Supplementary Material to:
Biological signal processing with a genetic toggle switch
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1 Derivation of a reduced model for the JK-latch with independent heterodimer binding sites

The reactions describing the genetic JK-latch, presented in Table S1 and S2 can be classified into three different categories: protein-protein reactions (including protein degradation), protein-DNA reactions and gene expression (including transcription and degradation of mRNA and translation into proteins). A full deterministic model can be derived by setting up a rate equation for the mean concentration of each biochemical species in the reactions. A first step towards a reduced model is to assume the occupation states of the promoters (that is the protein-DNA reactions) to be in equilibrium with the current transcription factor concentrations at all times. Since all TF’s act as repressors, we assume that the only promoter state contributing to mRNA production is the unoccupied one. Therefore the rate equation for mRNA of, for instance, gene A is,

\[
\frac{dm_A(t)}{dt} = \nu_{mA} P_A(t) - \lambda_m m_A(t),
\]

where \(\nu_{mA}\) is the maximal transcription rate, \(\lambda_m\) the mRNA degradation rate and \(P_A(t)\) denotes the promoter activity function, that is, the probability of the promoter to be unoccupied and free to bind RNA polymerase at a certain point in time. Hence, assuming all protein-DNA reactions to be in equilibrium, \(P_A(t)\) describes the equilibrium probability to find the promoter unoccupied as a function of current transcription factor concentration. This probability can also be calculated by thermodynamic models,
corresponding to [1,2]. Their specific form for the genetic JK-latch is defined in the boxes of Figs. 2 and 3.

As a further simplification, we assume all protein-protein reactions to rapidly reach equilibrium with the current total concentration of reacting proteins. For instance, for the concentrations of all protein species containing A and K this leads to the following set of equations\(^1\)

\[
A_{tot} = A + 2A_2 + KA, 
\]

\[
k_{A_2}^{on} A^2 - k_{A_2}^{off} A_2 - \lambda_p A_2 = 0, 
\]

\[
k_{KA}^{on} K A - k_{KA}^{off} K A - \lambda_p K A = 0, 
\]

where we can additionally set \(K = K_{tot} - KA\). After solving this system for the concentrations of homo- and heterodimers all promoter activity functions in Figs. 2 and 3 of the main text can then be expressed in terms of total concentrations, i.e., \(P_A(B_2, KA) = P_A(A_{tot}, B_{tot}, K_{tot}, J_{tot})\).

Altogether, we are left with two equations left to describe the total concentration of each gene product. For instance, the equations for gene product \(A\) are

\[
\frac{dA_{tot}(t)}{dt} = \nu_p A(t) - \lambda_p A_{tot}(t), 
\]

\[
\frac{dm_A(t)}{dt} = \nu_m P_A(t) - \lambda m_A(t). 
\]

As a last step, we set Eq. (6) to zero thereby assuming mRNA translation to be fast with respect to transcription (we will discuss this assumption in the next section). Solving the resulting equation for \(m_A\) and substituting it in Eq. (5) leads to

\[
\frac{dA_{tot}(t)}{dt} = \nu_A P_A(t) - \lambda_p A_{tot}(t), 
\]

with an effective maximal expression rate \(\nu_A = \nu_{m,A} \nu_{p,A} / \lambda_m\). Thus, each gene product in our models can be described by a single effective equation.

\(^1\)Note that we consider protein degradation to occur at the same rate for monomers and dimers (no cooperative stability), such that the total concentration of a gene product degrades with the same rate as monomers and dimers, see e.g. Ref. [3].
2 Derivation of a reduced model for the genetic J-K latch with mutually exclusive heterodimer binding sites

Making the same quasi-equilibrium assumptions for protein-protein and protein-DNA as in Section I, the reduced model of the genetic J-K latch with mutually exclusive binding sites for the heterodimers $KA$ and $JB$ reads

\[ \frac{dA_{\text{tot}}}{dt} = \alpha_A O_B(B_2) O_{KA}(KA, JB) - \lambda A_{\text{tot}}, \]

\[ \frac{dB_{\text{tot}}}{dt} = \alpha_B O_A(A_2) O_{JB}(KA, JB) - \lambda B_{\text{tot}}, \]

where only the operator occupancy functions of the heterodimers now also depend on the respective other heterodimer:

\[ O_B(B_2) = \left( 1 + \frac{B_2}{K_{B_2}} \right)^{-2}, \quad O_A(A_2) = \left( 1 + \frac{A_2}{K_{A_2}} \right)^{-2}, \quad (9a) \]

\[ O_{KA}(KA, JB) = \left( 1 + \frac{KA}{K_{KA}(1 + (JB/K_{JB}))} \right)^{-1}, \quad (9b) \]

\[ O_{JB}(KA, JB) = \left( 1 + \frac{JB}{K_{JB}(1 + (KA/K_{KA}))} \right)^{-1}. \quad (9c) \]

However, in order to introduce a delay required for a successful toggle operation, the unbinding kinetics of the overlapping operator sites needs to be slow (see main text), such that the quasi-equilibrium assumption that lead to Eqs. (9a) and (9b) is not strictly valid. To describe the slow dynamics of the overlapping operator complex we set up the master equation for the occupancy states of the overlapping operators. There are three states to be accounted for: (i) the binding site for $KA$ is occupied, (ii) the binding site for $JB$ is occupied and (iii) both binding sites are unoccupied. We denote the probabilities of these states as $q_{KA}$, $q_{JB}$ and $q_0$, respectively. The probabilities equivalent to Eqs. (9b) and (9c) to find the operator in an unoccupied state are then given by $O_{KA} = 1 - q_{KA}$ and $O_{JB} = 1 - q_{JB}$. Taking $k_{on}$
to be the on rate of both operators and $k_{\text{off}}$ the respective off-rate, the master equations read:

\begin{align}
\dot{q}_{KA} &= k_{\text{on}} KA q_0 - k_{\text{off}} q_{KA} \tag{10a} \\
\dot{q}_{JB} &= k_{\text{on}} JB q_0 - k_{\text{off}} q_{JB} \tag{10b} \\
\dot{q}_0 &= k_{\text{off}} (q_{KA} + q_{JB}) - k_{\text{on}} (KA + JB) q_0 \tag{10c}
\end{align}

These equations are controlled by the external variables $KA$ and $JB$. For given values of these variables, the system has a unique fixed point, which we denote by $(q_{KA}^*, q_{JB}^*, q_0^*)$. In the fixed point, the right hand side of Eq. (10) becomes zero, thus we can rewrite Eqs. (10a) and (10b) as

\begin{align}
\dot{q}_{KA} &= k_{\text{on}} KA (q_0 - q_0^*) - k_{\text{off}} (q_{KA} - q_{KA}^*) \tag{11a} \\
\dot{q}_{JB} &= k_{\text{on}} JB (q_0 - q_0^*) - k_{\text{off}} (q_{JB} - q_{JB}^*) \tag{11b}
\end{align}

Furthermore, we want to investigate Eqs. (10) under a constant toggle signal. Therefore, it is reasonable to assume that both heterodimers $KA$ or $JB$ are abundant in the system at all times. Then, the effective on-rates $k_{\text{on}} KA$ and $k_{\text{on}} JB$ are much faster than the off-rate $k_{\text{off}}$ and thus the probability $q_0$ to find the overlapping operator sites unoccupied is approximately zero. Additionally, substituting $q_{KA} = 1 - O_{KA}$ and $q_{JB} = 1 - O_{JB}$, the approximated equations can be written as

\begin{align}
\dot{O}_{KA} &\approx -k_{\text{off}} (O_{KA} - O_{KA}^*) \tag{12a} \\
\dot{O}_{JB} &\approx -k_{\text{off}} (O_{JB} - O_{JB}^*) \tag{12b}
\end{align}

We can identify the fixed points $O_{KA}^*$ and $O_{JB}^*$ with the quasi-equilibrium operator occupancy functions defined in Eqs. (9b) and (9c), which are functions of the time dependent heterodimer concentration. These equations can then be solved by variation of the constant:

\begin{align}
O_{KA}(t) &= \int_{0}^{\infty} k_{\text{off}} e^{-k_{\text{off}}(t-\tau)} O_{KA}^*(\tau) \, d\tau = \int_{0}^{\infty} g_k(\tau) O_{KA}^*(t-\tau) \, d\tau \tag{13a}, \\
O_{JB}(t) &= \int_{0}^{\infty} k_{\text{off}} e^{-k_{\text{off}}(t-\tau)} O_{JB}^*(\tau) \, d\tau = \int_{0}^{\infty} g_k(\tau) O_{JB}^*(t-\tau) \, d\tau \tag{13b}
\end{align}

with "memory kernel" $g_k(\tau) = k_{\text{off}} e^{-k_{\text{off}} \tau}$. Instead of accounting for the initial conditions in an own term,
we formally assume that for all negative times the system is held fixed at the initial point and integrate infinitely long into the past. This auxiliary assumption also simplifies the initial condition problem for the delayed dynamical system: for a distributed delay, instead of a single initial value, strictly the entire history prior to $t = 0$ has to be defined. Since we are interested in the qualitative long term behavior of the solution, we assume this history to be a single point from which the system is released at $t = 0$. Putting this result back into Eq. (8) leads to the delayed differential equations with exponentially distributed delay introduced in Fig. 3 Model 2.

**Additional delays in the reduced model.** For the following stability analysis, we include the additional delay caused by the translation process. While this is done for the sake of completeness, this additional delay is about ten times smaller than the delay caused by the overlapping heterodimer operators and is therefore not considered in the discussion of the main text. The additional delay arises from the reaction chain between the beginning of mRNA transcription and completion of protein translation. In principle, these kinds of delay alone can already lead to oscillations [4, 5]. Consider the rate equation for an arbitrary protein concentration $A$, with regulated transcription, linear degradation and an explicit equation for its mRNA denoted by $m_A$:

\[
\begin{align*}
\frac{dA}{dt} &= \nu_p m_A - \lambda_p A, \\
\frac{dm_A}{dt} &= \nu_m P_A(t) - \lambda_m m_A.
\end{align*}
\]

Here, $P_A(t)$ denotes an arbitrary promoter activity function of gene $A$. Solving Eq. (15) and treating the initial conditions as above leads to

\[
m_A(t) = \frac{\nu_m}{\lambda_m} \int_0^\infty \lambda_m e^{-\lambda_m \tau} P_A(t - \tau) d\tau.
\]

Putting this back into Eq. 14 leads to

\[
\frac{dA}{dt} = \nu_A \int_0^\infty g_A(\tau) P_A(t - \tau) d\tau - \lambda_p A
\]

with an effective expression rate $\nu_A = \frac{\nu_p \nu_m}{\lambda_m}$ and a delay kernel $g_A(\tau) = \lambda_m e^{-\lambda_m \tau}$. Thus, taking the dynamics of mRNA into account, leads to an additional distributed delay acting on the entire promoter activity function. In a reduced model the protein concentration therefore responds to a change in promoter
activity on a timescale given by the mRNA degradation rate. This is generally valid for any gene controlled by regulated recruitment. There are, however, mechanisms of gene regulation, involving active degradation of mRNA [6, 7]. In that case, the dynamics of mRNA have to be accounted for by its full dynamic description.

3 Linear stability analysis on a system with distributed delays

Here we discuss linear stability analysis for nonlinear systems with multiple cascaded exponentially distributed delays, which closely follows the book of [8]. The results presented here are generally applicable to any system of that kind. The model system considered here is the simple J-K latch with overlapping heterodimer operators in Eqs. (8) and (9). As discussed in the last sections, it contains delays on the promoter activity functions $P_A$ and $P_B$ and, within those, on the operator occupation functions $O_{KA}$ and $O_{JB}$. The memory kernels $g_k(\tau)$ and $g_\lambda(\tau)$ of these delays are exponential distributions with rate parameters $k_{\text{off}}$ and $\lambda_m$, respectively.

To perform a linear stability analysis on our model system, it first is important to note that while delays change the stability of a fixed point, they do not alter their position with respect to the undelayed system. This is easy to see: consider a fixed point $FP = (A^*, B^*)$ of the undelayed system and the delayed function $D_{g_k} \circ O_{KA}(A^*)$, where $D_{g_k}$ abbreviates a distributed delay with memory kernel $g$. We assume that the delayed system has stayed in this fixed point for a very long time, such that the delayed function can be written as

$$D_{g_k} \circ O_{KA}(A^*) = \int_0^\infty g_k(\tau)O_{KA}(A^*(t-\tau)) \, d\tau$$

$$= O_{KA}(A^*) \int_0^\infty g_k(\tau) \, d\tau = O_{KA}(A^*),$$

for any normalized kernel $g_k(\tau)$. Therefore, a fixed point of the undelayed system is a fixed point of the delayed system as well.

To perform a linear stability analysis on the model system Eq. (8), we must take into account that the dynamic variables $A$ and $B$ occur within different delay integrals and hence need to be treated separately in the linearization around a fixed point $FP$. Therefore, we categorize $A$ and $B$ by the delays acting on
them. The model system is

\[\dot{A} = \int_0^\infty g_\lambda(\tau)O_B^2(B_{g_\lambda}(t-\tau))(\int_0^\infty g_k(\tau')O_{KA}(A_{g_\lambda,k}(t-\tau-\tau'), B_{g_\lambda,k}(t-\tau-\tau'))d\tau')d\tau - \lambda_p A,\]

\[\dot{B} = \int_0^\infty g_\lambda(\tau)O_A^2(A_{g_\lambda}(t-\tau))(\int_0^\infty g_k(\tau')O_{JB}(A_{g_\lambda,k}(t-\tau-\tau'), B_{g_\lambda,k}(t-\tau-\tau'))d\tau')d\tau - \lambda_p B,\]

which can more formally be written as

\[\dot{A} = f(A, B, g_\lambda, A_{g_\lambda,k}, B_{g_\lambda,k}), \]

\[\dot{B} = h(B, A, g_\lambda, A_{g_\lambda,k}, B_{g_\lambda,k}).\]

Here, variables with subscript \( g_\lambda \) occur in functions, which are delayed only by mRNA degradation, whereas variables with subscript \( g_{\lambda,k} \) are delayed by mRNA degradation and the off-rate of the overlapping operators. We now can linearize the system around a fixed point \( FP = (A^*, B^*) \) and obtain linear equations for small deviations \( \tilde{A} = A - A^* \) and \( \tilde{B} = B - B^* \):

\[\frac{d\tilde{A}}{dt} = \frac{\partial f}{\partial A}|_{FP} \tilde{A} + D_{g_\lambda} \circ \left( \frac{\partial f}{\partial B_{g_\lambda}}|_{FP} \tilde{B} + D_{g_k} \circ \left( \frac{\partial f}{\partial A_{g_\lambda,k}}|_{FP} \tilde{A} + \frac{\partial f}{\partial B_{g_\lambda,k}}|_{FP} \tilde{B} \right) \right),\]

\[\frac{d\tilde{B}}{dt} = \frac{\partial h}{\partial B}|_{FP} \tilde{B} + D_{g_\lambda} \circ \left( \frac{\partial h}{\partial A_{g_\lambda}}|_{FP} \tilde{A} + D_{g_k} \circ \left( \frac{\partial h}{\partial B_{g_\lambda,k}}|_{FP} \tilde{B} + \frac{\partial h}{\partial A_{g_\lambda,k}}|_{FP} \tilde{A} \right) \right),\]

where \( D_{g_\lambda} \) denotes a delay with kernel \( \lambda_m e^{-\lambda_m \tau} \) and \( D_{g_k} \) a delay with kernel \( k_{off} e^{-k_{off} \tau} \). Since the position of the fixed point does not change for delayed variables, all partial derivatives are evaluated in
the same point \( FP \). Taking into account that the functionals \( D \) are linear, the system can be written as

\[
\begin{align*}
\frac{d\tilde{A}}{dt} &= \left. \frac{\partial f}{\partial A} \right|_{FP} \tilde{A} + \left. \frac{\partial f}{\partial B} \right|_{FP} D_{g_A} \circ D_{g_k} \circ \tilde{A} + \left. \frac{\partial f}{\partial A} \right|_{FP} D_{g_A} \circ D_{g_k} \circ \tilde{B}, \\
\frac{d\tilde{B}}{dt} &= \left. \frac{\partial h}{\partial B} \right|_{FP} \tilde{B} + \left. \frac{\partial h}{\partial A} \right|_{FP} D_{g_A} \circ D_{g_k} \circ \tilde{B} + \left. \frac{\partial h}{\partial A} \right|_{FP} D_{g_A} \circ D_{g_k} \circ \tilde{A}.
\end{align*}
\] (25, 26)

Herein, the delayed variables take the form

\[
\begin{align*}
D_{g_A} \circ \tilde{A} &= \int_0^\infty \lambda e^{-\lambda \tau} \tilde{A}(t-\tau) d\tau, \\
D_{g_A} \circ D_{g_k} \circ \tilde{A} &= \int_0^\infty \lambda_m e^{-\lambda m \tau} \left( \int_0^{\infty} e^{-k_{\text{off}} \tau'} \tilde{A}(t-\tau-\tau') \right) d\tau' d\tau.
\end{align*}
\] (27, 28)

We make the usual ansatz for a linear system: \( \dot{A} = c_A e^{zt} \) and \( \dot{B} = c_B e^{zt} \) with \( z \in \mathbb{C} \) to be determined. Upon this ansatz, the delayed variables become

\[
\begin{align*}
D_{g_A} \circ \tilde{A} &= c_A \int_0^\infty \lambda_m e^{-\lambda m \tau} e^{z(t-\tau)} = c_A e^{zt} \int_0^\infty \lambda_m e^{-\lambda m \tau} e^{z \tau} = \tilde{A} \times L(g_A(\tau); z), \\
D_{g_A} \circ D_{g_k} \circ \tilde{A} &= \tilde{A} \int_0^\infty \lambda_m e^{-\lambda m \tau} e^{-z \tau} \times \int_0^\infty k_{\text{off}} e^{-k_{\text{off}} \tau'} e^{-z \tau'} d\tau' d\tau = \tilde{A} \frac{\lambda_m}{\lambda_m + z}.
\end{align*}
\] (29, 30)

Variables with two delays can be evaluated in a similar way:

\[
\begin{align*}
D_{g_A} \circ D_{g_k} \circ \tilde{A} &= c_A \int_0^\infty \lambda_m e^{-\lambda m \tau} \left( \int_0^\infty k_{\text{off}} e^{-k_{\text{off}} \tau'} e^{z(t-\tau-\tau')} d\tau' \right) d\tau \\
&= \tilde{A} \int_0^\infty \lambda_m e^{-\lambda m \tau} e^{-z \tau} d\tau \times \int_0^\infty k_{\text{off}} e^{-k_{\text{off}} \tau'} e^{-z \tau'} d\tau' \\
&= \tilde{A} \frac{\lambda_m}{\lambda_m + z} \frac{k_{\text{off}}}{k_{\text{off}} + z}.
\end{align*}
\] (31)
Putting these results back into Eqs. 25 and 26 yields a linear equation for the coefficients $c_A$ and $c_B$:

$$
\begin{pmatrix}
  c_A z \\
  c_B z 
\end{pmatrix}
= 
\begin{pmatrix}
  \frac{\partial f}{\partial A} + \frac{\partial f}{\partial \lambda_{\lambda,k}} \frac{\lambda_m}{\lambda_m + z} k_{\text{off}} + \frac{\partial f}{\partial B} + \frac{\partial f}{\partial g\lambda,k} \frac{\lambda_m}{\lambda_m + z} k_{\text{off}} + z \\
  \frac{\partial h}{\partial A} + \frac{\partial h}{\partial \lambda_{\lambda,k}} \frac{\lambda_m}{\lambda_m + z} k_{\text{off}} + \frac{\partial h}{\partial B} + \frac{\partial h}{\partial g\lambda,k} \frac{\lambda_m}{\lambda_m + z} k_{\text{off}} + z 
\end{pmatrix} 
\begin{pmatrix}
  c_A \\
  c_B 
\end{pmatrix}
.$$  

(32)

Note that all partial derivatives have to be evaluated at the fixed point under consideration. Subtracting the left hand side from the diagonal elements of the right hand side leads to the familiar form an eigenvalue problem. Here, in contrast to a common linear stability analysis, additional polynomial terms are incorporated, representing the delays. In order for the system to have a solution, its determinant has to be zero. Thereby we obtain a characteristic equation for the exponents $z$.

$$
\begin{vmatrix}
  \frac{\partial f}{\partial A} + \frac{\partial f}{\partial \lambda_{\lambda,k}} \frac{\lambda_m}{\lambda_m + z} k_{\text{off}} + \frac{\partial f}{\partial B} + \frac{\partial f}{\partial g\lambda,k} \frac{\lambda_m}{\lambda_m + z} k_{\text{off}} + z \\
  \frac{\partial h}{\partial A} + \frac{\partial h}{\partial \lambda_{\lambda,k}} \frac{\lambda_m}{\lambda_m + z} k_{\text{off}} + \frac{\partial h}{\partial B} + \frac{\partial h}{\partial g\lambda,k} \frac{\lambda_m}{\lambda_m + z} k_{\text{off}} + z 
\end{vmatrix} = 0.
$$  

(33)

Evaluating the determinant, we obtain a polynomial of 6th order in $z$. In order for the considered fixed point to be stable, each solution of Eq. 33 must have a negative real part. The timescales of the system considered here are given by $\tau_{\lambda} = 1/\lambda_m$ and $\tau_{k} = 1/k_{\text{off}}$, which are the mean values of the memory kernel. These characteristic times are usually referred to as the mean delay [8]. Written in terms of the mean delay $\tau$, the Laplace transform of an exponential distribution becomes $1/(1 + \tau z)$. Thus, if the delay is small, its contributing terms in the characteristic equation are close to one and therefore do not change the stability of a fixed point. In the following we will employ the derived method of linear stability analysis to investigate the genetic JK-latch with overlapping heterodimer operators.

### 4 Analysis of deterministic and stochastic oscillations in the genetic JK-latch

To analyze the stability properties of the genetic JK-latch with overlapping heterodimer operators for a certain choice of parameters and concentration of input proteins $J_{\text{tot}}$ and $K_{\text{tot}}$, we applied the following procedure using Mathematica (Wolfram Research, Inc.): (i) calculate the concentrations of all homo- and heterodimers using Eqs. (2)-(4). For the stability analysis we manipulate the concentration of input proteins $J_{\text{tot}}$ and $K_{\text{tot}}$ directly, instead of controlling their transcription rate. (ii) Find all fixed points by
setting Eq. (8) to zero. (iii) For each fixed point solve the characteristic equation defined in Eq. (31). If the real parts of all solutions of the last step are negative, the fixed point is stable otherwise it is unstable. (iv) If necessary, characterize the system’s qualitative behavior (e.g. oscillatory) by solving its equations numerically. Employing this scheme for different parameter values we identify two important bifurcations: the system undergoes a Hopf bifurcation [9, 10] if the concentration of input proteins $J_{\text{tot}}$ and $K_{\text{tot}}$ is increased (see Fig. S2), then is oscillatory for a certain concentration range until it becomes stable again. Additionally, the system exhibits another Hopf bifurcation if the mean delay time is increased (i.e. the parameter $k_{\text{off}}$ is decreased), while a constant toggle signal is applied. This is illustrated in Fig. S3A: at a critical operator dwell time $\tau_{\text{crit}} \approx 32$ min the system becomes oscillatory with a period that increases with the dwell time.

In the form we set up the reduced model for the genetic JK-latch with overlapping heterodimer operators, we couple the dynamics of protein concentrations with the intrinsically fluctuating switching of operator states. Therefore, although the deterministic model provides valuable insights into the working principles of the circuit, the quantitative period of the stochastic system is not correctly reproduced. This is shown in Fig. 3B, where it is apparent that the average of the stochastic period is much shorter than the period of the deterministic model. Also the stochastic system displays oscillations in a regime of short heterodimer dwell times, in which the deterministic system does not oscillate at all. In fact, it is known that a small number of reactant molecules together with negative feedback and time delay in gene expression can lead to delay-induced instabilities, such that a system turns oscillatory even when its deterministic counterpart is not [4].

If the delay (dwell time) is long, the strength of the toggle signal is chosen appropriately and the signal duration is well timed, then high probabilities of a correct toggle response can be achieved – as can be seen in Fig. S4A and B.

5 Genetic master-slave latch

We extended the reaction system of the JK-latch to a master-slave latch by adding additional regulation of the signal genes $J$ and $K$ – see Table S4. This was done by firstly including homodimerization of signal proteins. $J_2$ and $K_2$ bind two operators on the promoter of the respective other gene to repress it. Additionally, gene $J$ is repressed by heterodimers $KA$ by binding to a third operator in its promoter
region, while gene $K$ is repressed by $JB$ in the same way. Notably, the restrictive promoter layout of the overlapping heterodimer operators on genes $A$ and $B$ with very slow unbinding rates is no longer necessary in the master-slave latch and is changed to independent operators with normal unbinding rates.

With this additional regulation, signal genes $J$ and $K$ become bistable when their maximal transcription rate is increased externally (by the toggle signal) and thereby form the master toggle switch. Then, the master toggle switch settles into a state that is determined by the current concentrations of heterodimers and therefore by the current state of the slave latch. Dependent on the operator strengths for homodimers and for heterodimers two kinds of erroneous responses are possible upon a toggle signal: If homodimers bind to strongly with respect to heterodimers, then the master switch is prone to stochastic fluctuations at the onset of the toggle signal and biased towards that signal protein which is first abundant enough to form homodimers. If, on the other hand, heterodimer binding is too strong, random single bursts of the repressed signal gene can be sufficient to switch the state of the master latch. Fig. S4C shows the toggle probability as a function of those key parameters for a toggle signal duration of 60 min. It can be seen that the master-slave latch is robust in that sense, that it responds correctly for a broad range of those parameters.
JK-latch with overlapping heterodimer operators

**Promoter and operator states**

- $P_A | \cdot \rangle + B_2 \xrightarrow{k_{on}} P_A | B_2 \rangle$
- $P_A | \cdot \rangle + B_2 \xrightarrow{k_{off}} P_A | \cdot \rangle + B_2$
- $P_A | B_2 \rangle + B_2 \xrightarrow{k_{on}} P_A | B_2 | B_2 \rangle$
- $P_A | B_2 \rangle + B_2 \xrightarrow{k_{off}} P_A | B_2 | B_2 \rangle$
- $P_B | \cdot \rangle + A_2 \xrightarrow{k_{on}} P_B | A_2 \rangle$
- $P_B | \cdot \rangle + A_2 \xrightarrow{k_{off}} P_B | \cdot \rangle + A_2$
- $P_B | A_2 \rangle + A_2 \xrightarrow{k_{on}} P_B | A_2 | A_2 \rangle$
- $P_B | A_2 \rangle + A_2 \xrightarrow{k_{off}} P_B | A_2 | A_2 \rangle$
- $O | \cdot \rangle + KA \xrightarrow{k_{on}} O | KA \rangle$
- $O | \cdot \rangle + KA \xrightarrow{k_{off}} O | \cdot \rangle + KA$
- $O | \cdot \rangle + JB \xrightarrow{k_{on}} O | JB \rangle$
- $O | \cdot \rangle + JB \xrightarrow{k_{off}} O | \cdot \rangle + JB$
Dimerization

\[ 2A \xrightarrow{k_{{}^{on}}A} A_2 \]
\[ 2B \xrightarrow{k_{{}^{on}}B} B_2 \]
\[ A_2 \xrightarrow{k_{{}^{off}}A} 2A \]
\[ B_2 \xrightarrow{k_{{}^{off}}B} 2B \]

Turnover

\[ m_A \xrightarrow{\lambda_{m_A}} A \]
\[ m_B \xrightarrow{\lambda_{m_B}} B \]
\[ m_J \xrightarrow{\lambda_{m_J}} J \]
\[ m_K \xrightarrow{\lambda_{m_K}} K \]
\[ A \xrightarrow{\lambda_p} A \]
\[ B \xrightarrow{\lambda_p} B \]

Transcription and translation

\[ P_J \xrightarrow{\nu_{m_J}} m_J \]
\[ P_K \xrightarrow{\nu_{m_K}} m_K \]
\[ P_A \xrightarrow{\nu_m} P_A \]
\[ P_B \xrightarrow{\nu_m} P_B \]
\[ P_A \xrightarrow{\nu_{m_J} + O} m_A \]
\[ P_B \xrightarrow{\nu_{m_K} + O} m_B \]

Table S1. All reactions involved in the genetic JK-latch with overlapping heterodimer binding sites.

A graphical representation of the reaction network is shown in Fig. S1. Proteins and their dimers are denoted by capital letters; transcripts of a gene \( X \) are denoted by \( m_X \). A gene \( X \) is represented by its promoter \( P_X \), which can be occupied by transcription factors. Each occupation state of a promoter is represented by an own species of reactants for which an empty operator is indicated by \( \cdot \) and an occupied operator by the name of the respective transcription factor. In this notation the different binding sites are separated by the symbol \( | \). To reduce the number of occupation state combinations the operator complex for heterodimers is separated from the other promoter states and denoted by \( O \). To make transcription nevertheless conditional on heterodimers, we include the respective species for an empty binding site as reactant and product into the corresponding transcription reaction. This ensures that transcription requires an empty heterodimer operator to proceed but does not change its concentration. All corresponding parameter values are listed in Table S3.

Additional reactions in J-K latch without overlapping heterodimer operators

\[ O \xrightarrow{\nu_{m_J} + O} P_A \]
\[ O \xrightarrow{\nu_{m_K} + O} P_B \]

Table S2. With independent heterodimer operators, heterodimers can bind simultaneously to their binding sites. All other reactions are the same as in the JK-latch with overlapping heterodimer operators, listed in Table S1.
| Parameter | Value | Description and References |
|-----------|-------|-----------------------------|
| **Transcription** | | |
| \( \nu_m^A, \nu_m^B \) | 5 min\(^{-1} \) | strong promoter [11] |
| \( \nu_m^J, \nu_m^K \) | (0.01 − 1) min\(^{-1} \) | inducible promoter [11]; induction can be achieved e.g. by upstream binding activators [12] or via small non-coding RNAs [13] |
| **Translation** | | |
| \( \nu_p^A, \nu_p^B, \nu_p^J, \nu_p^K \) | 0.23 min\(^{-1} \) | corresponds to a burst factor of \( b = 10 \) [14] |
| **Degradation** | | |
| \( \lambda_m \) | 0.23 min\(^{-1} \) | active degradation by RNase [15]; half-life of 3 min |
| \( \lambda_p \) | 0.138 min\(^{-1} \) | active proteolysis by SsrA-tags [16]; half life of 5 min |
| **Dimerization** | | |
| \( k^{on}_A, k^{on}_B, k^{on}_{KA}, k^{on}_{JB} \) | 0.2 nM\(^{-1} \text{min}^{-1} \) | assumed to be diffusion limited [17] |
| \( k^{off}_A, k^{off}_B, k^{off}_J, k^{off}_K \) | 2 min\(^{-1} \) | typical value for dimerization |
| **Protein-DNA binding** | | |
| \( k^{on}_{O_A2}, k^{on}_{O_B2} \) | 0.2 nM\(^{-1} \text{min}^{-1} \) | assumed to be diffusion limited [17] |
| \( k^{off}_{O_A2}, k^{off}_{O_B2} \) | 2 min\(^{-1} \) | modified operator sequence assumed; parameter in the range of \( \text{tet-O}_1 \) binding strength [18] |
| \( k^{on}_{O_{KA}}, k^{on}_{O_{JB}} \) | 0.2 nM\(^{-1} \text{min}^{-1} \) | assumed to be diffusion limited [17] |
| \( k^{off}_{O_{KA}}, k^{off}_{O_{JB}} \) | 0.0126 min\(^{-1} \) | very slow off-rate assumed [19,20] |

Table S3. Values and references for all parameters used in the reactions of the genetic JK-latch.
**Additional reactions in master-slave latch**

**Promoter and operator states**

- $P_J|· + K_2 \xrightarrow{s_{\text{on}}^{K_2}} P_J|K_2|·$
- $P_J|· + K_2 \xrightarrow{s_{\text{off}}^{K_2}} P_J|K_2|·$
- $P_J|K_2| + K_2 \xrightarrow{s_{\text{on}}^{K_2}} P_J|K_2|K_2$
- $P_J|K_2| + K_2 \xrightarrow{s_{\text{off}}^{K_2}} P_J|K_2|K_2$
- $P_K|· + J_2 \xrightarrow{s_{\text{on}}^{J_2}} P_K|J_2|$
- $P_K|· + J_2 \xrightarrow{s_{\text{off}}^{J_2}} P_K|J_2|$
- $P_K|J_2| + J_2 \xrightarrow{s_{\text{on}}^{J_2}} P_K|J_2|J_2$
- $P_K|J_2| + J_2 \xrightarrow{s_{\text{off}}^{J_2}} P_K|J_2|J_2$
- $O_J|· + KA \xrightarrow{s_{\text{on}}^{KA}} O_J|KA$
- $O_K|· + JB \xrightarrow{s_{\text{on}}^{JB}} O_K|JB$

**Dimerization**

- $2J \xrightarrow{s_{\text{on}}^{J}} J_2$
- $2K \xrightarrow{s_{\text{on}}^{K}} K_2$

**Turnover**

- $J_2 \xrightarrow{\lambda_p} ∅$
- $K_2 \xrightarrow{\lambda_p} ∅$

**Transcription**

- $P_J|· + O_J|· \xrightarrow{\nu_{m_J}} P_J|· + O_J|· + m_J$
- $P_K|· + O_K|· \xrightarrow{\nu_{m_K}} P_K|· + O_K|· + m_K$

**Table S4.** Additional reactions involved in regulation of genes J and K. All other reactions are the same as in the genetic JK-latch with independent heterodimer operators. Values of new parameters are listed in Table S5.
| Parameter                      | Value               | Description and References                                                                 |
|-------------------------------|---------------------|---------------------------------------------------------------------------------------------|
| **Transcription**             |                     |                                                                                             |
| $\nu_{mJ}, \nu_{mK}$          | $(0.05 - 5) \text{ min}^{-1}$ | strong inducible promoter [11]; induction can be achieved e.g. by upstream binding activators [12] or via small non-coding RNAs [13] |
| **Dimerization**              |                     |                                                                                             |
| $s^{on}, s^{on}_K$            | $0.2 \text{ nM}^{-1}\text{min}^{-1}$ | assumed to be diffusion limited [17]                                                          |
| $s^{off}, s^{off}_K$          | $2 \text{ min}^{-1}$ | equal to dimerization of $A$ and $B$                                                          |
| **Protein-DNA binding**       |                     |                                                                                             |
| $s^{on}_{O_{J2}}, s^{on}_{O_{K2}}$ | $0.2 \text{ nM}^{-1}\text{min}^{-1}$ | diffusion limited [17]                                                                          |
| $s^{off}, s^{off}_{O_{K2}}$   | $2 \text{ min}^{-1}$ | optimized for highest toggle probability                                                       |
| $s^{on}_{O_{KA}}, s^{on}_{O_{JB}}$ | $0.2 \text{ nM}^{-1}\text{min}^{-1}$ | binding to the operator on the promoter                                                        |
| $s^{off}_{O_{KA}}, s^{off}_{O_{JB}}$ | $0.35 \text{ min}^{-1}$ | optimized for highest toggle probability                                                       |
| **Changes in slave circuit**  |                     |                                                                                             |
| $k^{off}_{O_{KA}}, k^{off}_{O_{JB}}$ | $2 \text{ min}^{-1}$ | use same binding strength as homodimers                                                       |

Table S5. Additional parameters used in the genetic master-slave latch. Instead of the symbol $k$ for rates in the slave circuit, the symbol $s$ is used for rates of all additional processes (dimerization and protein-DNA binding of signal proteins) in the master circuit. Since a delay in heterodimer binding is no longer needed, the assumption of very slow unbinding kinetics of heterodimers from promoters of genes $A$ and $B$ has been released.
Figure S1. Schematic of the reaction network of the JK-latch with overlapping heterodimer operators. All reactions contained in the model are depicted together with their rates. On- and off-rates ($k_{on}$ and $k_{off}$) are combined to dissociation constants $K = k_{off}/k_{on}$. 
Figure S2. (A) Bifurcation diagram of the genetic JK-latch with overlapping heterodimer operators. The relative position of the system’s fixed points is shown with the total concentrations of input proteins $J_{\text{tot}}$ and $K_{\text{tot}}$ as synchronously tuned bifurcation parameter. (B) Value of the critical root calculated in the linear stability analysis, presented in Section 3, for the upmost fixed point (ON state) in Fig. (A). The root is critical in that sense that it is the only solution of the characteristic equation, which becomes positive for certain values of $J_{\text{tot}}/K_{\text{tot}}$. Starting from a low concentration of input proteins the system initially has two stable (solid line) and one unstable (dotted line) fixed point. The upper stable fixed point represents the ON state, whereas the lower one represents the OFF state. As the concentration of input proteins is increased, the two stable fixed points lose their stability at a critical concentration $c_1$, after which delay induced oscillations commence (orange lines). At input protein concentration $c_2$ the three fixed points collapse to one. That, however, does not alter the system’s oscillatory behavior. At concentration $c_3$ the system’s single fixed point becomes stable again. This is due to depletion of homodimers by increasingly forming heterodimers, which eventually abolishes the switch-like behavior of genes $A$ and $B$. 

Figure S3. (A) Frequency of the race condition as a function of the average dwell time $\tau$ of a heterodimer on the overlapping operator complex (inset shows the period of the same). At a critical dwell time $\tau_{\text{crit}} \approx 32$ min the system undergoes a Hopf bifurcation and thereafter oscillates with a period that is approximately linear in $\tau$. (B) Average period of the stochastic race condition as a function of the mean dwell time $\tau$. In comparison to (A) the stochastic system oscillates even for small dwell times (although with very low amplitude) and has a shorter period than the deterministic model. The latter is due to the fact that the deterministic model couples the dynamics of protein concentrations with probabilities of operator switching, which has a distorting effect on the oscillatory dynamics.
Figure S4. Toggle probability $p_{\text{toggle}}$ as a function of key parameters. Contours indicate equal probability to toggle successfully into the complementary state. Each data point (a grid of 30 by 30 per plot) is estimated by testing the final state of 5000 simulation runs of the respective full stochastic model. (A) Dependence of $p_{\text{toggle}}$ on the duration $T$ and the strength of the toggle signal in the JK-latch. Here the strength of the toggle signal is tuned by a concerted variation of the transcription rates of genes $J$ and $K$, i.e., $\nu_J = \nu_K = \nu$. (B) Dependence of $p_{\text{toggle}}$ on the duration $T$ of the toggle signal and the off-rate $k_{\text{off}}$ for unbinding the overlapping operator sites. As expected, $p_{\text{toggle}}$ increases as the delay (dwell time of heterodimers on the binding site) is increased. (C) Dependence of $p_{\text{toggle}}$ in the master-slave latch on the additional rates $s_{J2}^{\text{off}}$, $s_{K2}^{\text{off}}$ for unbinding the homodimer operators and $s_{JB}^{\text{off}}$, $s_{KA}^{\text{off}}$ for unbinding the heterodimer operators in the additional toggle switch (master latch). The master-slave latch is robust in that sense that $p_{\text{toggle}}$ is high for a broad range of parameters.
Dynamics of the deterministic model of the master-slave latch. (A) Phase diagram of the toggle switch as a function of the effective maximal transcription rates $\tilde{\nu}_A$ and $\tilde{\nu}_B$. The parameters are chosen such that the system is in the bistable regime in the absence of input signals (point O) and the circuit is set to the ON state initially. The curves indicate dynamic changes of $\tilde{\nu}_A$ and $\tilde{\nu}_B$, incurred by applying the toggle signal (simultaneous expression of both input genes J and K) for 200 min (green solid curve) and then releasing it (red curve). In contrast to the JK-latch, even if the toggle signal is applied a long time, the system enters the correct monostable regime (A low, B high), switches to the OFF state and returns to the bistable region without approaching the other monostable regime. In particular, the master-slave latch does not oscillate, even under a continuous toggle signal. (B) The same trajectories in the $([A_{\text{tot}}],[B_{\text{tot}}])$-plane.
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