De novo design of a synthetic riboswitch that regulates transcription termination

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ABSTRACT

Riboswitches are regulatory RNA elements typically located in the 5’-untranslated region of certain mRNAs and control gene expression at the level of transcription or translation. These elements consist of a sensor and an adjacent actuator domain. The sensor usually is an aptamer that specifically interacts with a ligand. The actuator contains an intrinsic terminator or a ribosomal binding site for transcriptional or translational regulation, respectively. Ligand binding leads to structural rearrangements of the riboswitch and to presentation or masking of these regulatory elements. Based on this modular organization, riboswitches are an ideal target for constructing synthetic regulatory systems for gene expression. While riboswitches for translational control have been designed successfully, attempts to construct elements regulating transcription have failed so far. Here, we present an in silico pipeline for the rational design of synthetic riboswitches that regulate gene expression at the transcriptional level. Using the well-characterized theophylline-aptamer as sensor, we designed the actuator part as RNA sequences that can fold into functional intrinsic terminator structures. In the biochemical characterization, several of the designed constructs show ligand-dependent control of gene expression in *E. coli*, demonstrating that it is possible to engineer riboswitches not only for translational but also for transcriptional regulation.
INTRODUCTION

Synthetic biology has become a rapidly developing branch of life sciences. At its heart is the rational design and construction of biosynthetic and regulatory networks, pathways and their constituent components. As RNA plays a central role in a diverse set of regulatory pathways, efforts in synthetic biology have produced novel RNA components capable of regulating gene expression in vivo {Isaacs 2006 #71}. Riboswitches are a particularly attractive type of gadget for synthetic biology applications {Wieland 2008 #106}{Topp 2010 #107}. These naturally occurring RNA elements determine the expression state of a gene in response to an external signal (in most cases a small molecule) through a conformational change, regulating either transcription or translation. The fundamental architecture of a riboswitch combines a sensor (aptamer domain) for the external stimulus and an actuator that influences gene expression. Both elements can in principle be freely combined. Depending on the response to the trigger molecule, gene expression is either enhanced (ON switch) or repressed (OFF switch). In transcription-regulating riboswitches, the actuator element contains a possible intrinsic transcription terminator structure consisting of a stem-loop element followed by a U-rich region. Depending on the bound or unbound state of the sensor, this terminator structure is either masked or formed, leading to activation or inhibition of gene expression, respectively.

While structural features of the conserved aptamer domains are well described, the organization of the regulatory platforms is in many cases unclear. In particular, transcription terminator elements can vary substantially, not only as isolated elements but also in the context of a specific riboswitch {Breaker 2011 #69}{Hoon 2005 #50}{Lesnik 2001 #51}. Hence, the design of transcription regulating riboswitches is a difficult task. This was demonstrated by Fowler et al., who intended to design transcriptional ON switches by fusing a theophylline aptamer to a B. subtilis
intrinsic terminator {Fowler 2008 #131}. Detailed analysis revealed, however, that the regulatory mechanism of the selected riboswitches was independent of the terminator structure and was obviously targeting translation. Accordingly, currently available riboswitches generated by rational and/or evolutionary approaches exclusively regulate translation {Hanson 2005 #2}{Ogawa 2011 #45}{Beisel 2009 #117}{Sinha 2010 #112}{Muranaka 2009 #35}{Wieland 2009 #113}.

We used the well characterized theophylline aptamer as a sensor element to construct synthetic riboswitches that regulate gene expression at the transcriptional level. This aptamer has been successfully incorporated in different regulatory mechanisms for gene expression in vivo, demonstrating its functionality within the cell. Besides representing a sensor platform in translational riboswitches {Desai 2004 #1}{Topp 2007 #34}{Suess 2004 #3}, it was used to modulate mRNA cleavage catalysed by yeast RNase III {Babiskin 2011 #141} as well as RNA splicing {Kim 2005 #142}{Thompson 2002 #143} and to control non-coding RNA activity {Qi 2012 #146}. Fusing this aptamer to a computer-designed spacer element followed by an aptamer-complementary sequence, which together form a transcriptional terminator structure, several synthetic riboswitches were generated. Biochemical analysis revealed that these elements are indeed able to efficiently regulate transcription of a reporter gene. Hence, while several in vivo screening approaches have failed so far, our data show that a strategy based on in silico design is successful in constructing functional transcriptional riboswitches in E. coli.

**Results**

We devised an in silico selection pipeline to rationally design transcriptional riboswitches that are composed of (1) the theophylline aptamer TCT8-4 that selectively binds its target at a $K_d$ of 0.32 µM and strongly discriminates against the
structurally closely related caffeine {Jenison 1994 #46}, (2) a spacer sequence, (3) a sequence complementary to the 3'-part of the aptamer and (4) a U-stretch (Fig. 1 A). The rational model behind such a construct is to terminate transcription of a host gene if the target molecule theophylline is absent. This is achieved by the formation of a terminator structure composed of the aptamer 3'-part and elements (2), (3) and (4). Ligand binding stabilizes the aptamer structure. Thus, the 3'-end of the aptamer sequence is unavailable for the formation of the terminator. As a consequence, the complete sequence of the host gene is transcribed and subsequently translated. Accordingly, a functional construct represents a transcriptional ON riboswitch.

The functional role of the spacer sequence is twofold. First, it has to ensure that the binding-competent aptamer structure is formed and has sufficient time during transcription to sense the target molecule. Second, it has to provide a functional part within the terminator structure. These requirements translate to specific constraints on both composition and length of a functional spacer sequence. The spacer must not form base pairs with the aptamer sequence during transcription in order to avoid potential inhibition of target molecule sensing. This implies a structural constraint. The design algorithm starts from randomly generated spacer sequences with a length between 6 and 20 nt and employs folding simulations on the concatenation of aptamer and spacer to remove candidates that violate the described constraints. The reverse complement of the 3'-part of the aptamer followed by an eight nucleotide long U-stretch is appended to the valid aptamer-spacer constructs, completing the possible intrinsic terminator structure. The co-transcriptional folding paths of these complete constructs are simulated using RNAfold predictions {Lorenz 2011 #135}.

Based on the above described design principles, twelve theophylline sensing riboswitches have been constructed. From these predictions, six riboswitch constructs were selected, based on the following criteria of their computationally
determined folding paths: (a) the unbound full length transcript disrupts the aptamer and folds into a stable terminator structure; (b) no stable secondary structure is formed downstream of the aptamer/theophylline complex; (c) the calculated energy difference between bound and unbound state should be close to the binding energy of -8.96 kcal/mol of the aptamer/theophylline complex {Gouda 2003 #132}. The designed constructs were assembled by overlap-extension PCR and cloned into plasmid pBAD2_bgaB between the ara promoter and the reporter gene for β-galactosidase from B. stearothermophilus {Klinkert 2012 #137} (Fig. 1B, 2A). Transcription was induced by arabinose in the presence or absence of 2 mM theophylline, and β-galactosidase activity of cell extracts was quantified using ONPG (ortho-nitrophenyl-β-D-galactopyranoside). Three of the constructs turned out to stimulate reporter gene expression by two-fold (RS1 and RS3) or three-fold (RS10) (Fig. 2B). Caffeine, as a control, could not activate the constructs. Fusing riboswitches RS2 and RS4 to the reporter gene resulted in a permanent active state of gene expression, independent of theophylline or caffeine. In contrast, in the RS8 construct, the reporter gene showed a rather low level of expression, resembling an inactive state under all tested conditions.

As RS10 had the best theophylline-dependent activation ratio (Fig. 2B), it was chosen for further investigations. To optimize this riboswitch, we aimed for a reduction of the background expression level in the absence of theophylline. First, we replaced the tetraloop sequence of the original RS10 construct by GAAA which is frequently found in natural terminators {d'Aubenton Carafa 1990 #139}. Second, as a G-C base pair forms a stable basis for a tetraloop structure, we mutated the terminal A-U base pair in the terminator stem to G-C, leading to construct RS10loop (Fig. 3A). This modification led to an increased efficiency in termination, as the background activity of the riboswitch was reduced from 40 to about 15 Miller units (MU) (Fig. 3B).
However, theophylline-induced gene expression activity was also lowered compared to the original construct. Obviously, this optimization of the terminator element stabilized its structure, resulting in an inhibition of a ligand-induced refolding of the upstream aptamer element required for ongoing transcription. Furthermore, it was described that a stretch of eight U residues closely upstream of the ribosomal binding site (RBS) strongly enhances ribosome attachment and translation, while shorter U stretches have no such effect {Zhang 1992 #134}. As our riboswitches are located immediately upstream of the RBS, it is conceivable that the terminator U-stretch of eight residues (U₈) stimulates translation in a similar way, leading to an unspecific increase in background activity. Indeed, a construct bgaBU₈, consisting of the U₈ stretch only (Fig. 3A), showed a dramatic and constitutive increase in reporter gene activity up to 100 MU, compared to the original construct bgaB with 31 MU (Fig. 3B). To prove that this effect depends on the close proximity of the U₈ stretch and the ribosomal binding site, a 19 nt region from the original pBAD multiple cloning site was inserted upstream of the reporter gene without a riboswitch, leading to construct bgaBshift (Fig. 3A). In contrast to the U₈ insertion, this construct did not show a significant increase in gene activity compared to the original bgaB construct (Fig. 3B). Accordingly, we inserted the 19 nt spacer element downstream of RS10, leading to construct RS10shift (Fig. 3A). In the activity test, this riboswitch showed a strongly reduced background signal of approximately 10 MU, resulting in a 6.5-fold activation of gene expression upon theophylline addition (Fig. 3B). To demonstrate the generality of this observation, we inserted the same 19 nt spacer region between RS3 and the ribosomal binding site. Similar to RS10shift, the RS3shift construct also showed a 5.7-fold increased ON/OFF rate, due to a reduced background activity in the absence of the ligand (18 MU compared to 54 MU in RS3; Fig. 3A, B).
To further verify the functionality of the selected riboswitches, individual components were examined in the expression system. Deleting the 5'-part of the aptamer sequence led to isolated terminator structures T1 to T10 that showed efficient repression of the reporter gene activity (Fig. 4A, B). As a positive control, the terminator of the attenuator region of the pyrBI operon from *E. coli* was used {Turnbough 1983 #147}. This natural regulatory element showed a termination efficiency comparable to that of the synthetic terminators, indicating that all generated terminator elements are functional (Fig. 4B).

The observed terminator-dependent reduction of reporter gene activity is a clear indication that the synthetic riboswitches indeed control transcription. Hence, the integrity of the terminator structure should be essential for regulation, and a deletion of this element should lead to a constitutive active state. To test this, we removed the whole 3'-part of the terminator region of RS10, i.e. the complementary 3'-part of the hairpin and the U-stretch, resulting in RS10ΔT (Fig. 4A). This deletion led to a complete destruction of riboswitch function, and the construct exhibited permanent gene expression, independent of theophylline (Fig. 4B).

Because the essential control elements for translation, in particular the ribosomal binding site, are still present in RS10ΔT as well as the corresponding terminator construct T10, these findings are a clear indication that the RS10 riboswitch regulates at the transcriptional level. To confirm this observation, Northern blot analysis was performed for RS10 and the terminator-alone construct T10. Total RNA was prepared from corresponding *E. coli* TOP10 cells grown in the presence or absence of theophylline. Hybridization was performed using bgaB-specific oligonucleotide probes. To make sure that identical amounts of RNA were analysed, a probe for 23S rRNA was used as an internal control. While the intensity of this control signal was identical in each preparation, a full length mRNA of the RS10-
bgaB construct was detected only in the presence of theophylline (Fig. 4C). The T10 construct, lacking the aptamer domain, completely inhibited transcription. Taken together, this result is in agreement with the corresponding β-galactosidase activity tests and clearly shows that RS10 regulates at the transcriptional level.

Discussion

We have successfully applied a combination of in silico, in vitro as well as in vivo experiments to generate theophylline-dependent riboswitches that activate transcription upon binding of the target molecule. This strategy is a promising alternative to riboswitch selection based on randomized RNA sequence libraries. The design of such libraries is crucial for the success of selection and requires features that differ depending on whether an ON or an OFF switch is to be designed {Muranaka 2009 #148}{Topp 2008 #37}. Furthermore, the screening approach with randomized RNA sequences was applied successfully only to riboswitches acting on translation {Desai 2004 #1}{Lynch 2007 #32}{Topp 2008 #4}{Nomura 2007 #109}{Fowler 2008 #131}. For such elements, the sole requirement is that the alternative structures lead to presentation or masking of the ribosomal binding site, which comprises a short sequence of four to six nucleotides {Steitz 1975 #149}. In transcription-controlling riboswitches, on the other hand, the alternative conformations have to promote or inhibit the formation of a larger and much more complex structure, i.e. an intrinsic terminator consisting of a hairpin element followed by a U-rich single-stranded region. Obviously, the screening of such a high sequence complexity with defined structural requirements is a very difficult task for a selection approach and corresponding efforts resulted again in riboswitches controlling translation, but not transcription as planned {Fowler 2008 #131}. Hence, our rational design seems to be more promising for the design of such synthetic regulators.
Interestingly, theophylline-dependent translational riboswitches and the transcription-controlling elements presented here show a similar efficiency in regulation (8-fold versus 6.5-fold) [Suess 2004 #3][Desai 2004 #1]. However, these constructs are transcribed from different promoters (B. subtilis XylA, E. coli IS10) that can also influence termination efficiency [Goliger 1989 #150]. Furthermore, it was shown that sequences downstream of the U-stretch can have an impact on termination efficiency [d'Aubenton Carafa 1990 #139][Lesnik 2001 #51], while additional U-rich elements enhance translation [Zhang 1992 #151]. Sequences immediately downstream of the start codon, described as downstream boxes, can also act as translational enhancers, as it was demonstrated for several E. coli as well as bacteriophage genes [Sprengart 1996 #152]. Besides these and other sequence-specific features, there are further properties that greatly influence the expression of a given gene. Secondary structure elements and position effects of the above mentioned regulatory regions within a translational unit play a crucial role for productivity of protein synthesis [Kozak 2005 #154][Makrides 1996 #153]. As these factors have a great impact on the efficiency of riboswitches in general, it is virtually impossible to compare the regulatory power of the elements. On the other hand, this plethora of features influencing bacterial gene expression allows a further optimization of the individual constructs, eventually leading to higher ON/OFF rates.

For transcription-controlling riboswitches, a kinetic control is discussed, where the riboswitch does not reach a thermodynamic equilibrium with its ligand, but is driven by the kinetics of RNA folding, ligand interaction and RNA synthesis [Wickiser 2005 #9][Wickiser 2005 #8][Rieder 2007 #18]. Such a competition between ligand binding and speed of transcription seems to be a common regulatory feature. Several transcriptional riboswitches force the polymerase to pause downstream of the terminator element, giving the sensor sufficient time for ligand binding and,
consequently, deciding whether to proceed or terminate transcription {Haller 2011 #25}. In our riboswitches, sophisticated structural features such as transcriptional pause sites {Wickiser 2005 #8} or the energetic contribution of ligand binding to the stability of the riboswitch conformation {Gouda 2003 #132} were not yet included in the design principles. The $K_d$ value used to estimate the binding energy between target molecule and aptamer depends on experimental conditions, e.g. salt concentration and temperature, which are uncertain for the theophylline-aptamer used here. Hence, the resulting $\Delta G$ value represents only a rough estimate and cannot be directly integrated into the pipeline at the moment. However, it should be possible to construct substantially more effective synthetic regulators of transcription by incorporating an explicit model of RNA polymerase and ligand binding behaviour into the rational design. The current design strategy is based on minimum free energy evaluations. This works as long as the unbound aptamer structure folds into the MFE conformation as it is the case for the used theophylline-aptamer. It needs to be replaced, however, if alternative structures are adapted preferentially. The usage of advanced co-transcriptional folding simulations and explicit predictions of alternative conformations will become necessary at that point.

As transcriptional riboswitches require an overlapping region between sensor (theophylline aptamer) and actuator (terminator), it is not possible to simply fuse a naturally occurring terminator to the aptamer. Natural intrinsic terminators consist of a GC-rich hairpin followed by a U-rich region that readily dissociates from the DNA-template {d’Aubenton Carafa 1990 #139}{Martin 1980 #155}. As the hairpin element does not show a high conservation at the sequence level, this part of the terminator can be easily manipulated without destroying its regulatory function {Wilson 1995 #156}. As a consequence, it is possible to design artificial terminators simply by adjusting the 5'-half of the hairpin element to the sequence required for proper base
pairing with the 3'-part of the aptamer. This is demonstrated by the fact that all
designed terminator structures significantly shut down gene expression and
represent therefore functional elements, comparable to natural terminators.
Obviously, the simple descriptors used for our in silico construction are sufficient to
predict efficient termination of the reporter system.
As shown in figure 2, not every generated terminator led to a functional riboswitch.
Nevertheless, the calculated free energy values of the individual terminators are in
good agreement with functionality of the constructs (Fig. 5). The terminator structure
of RS8 showed the highest stability (-29.00 kcal/mol), suggesting that it is formed
even in the presence of theophylline, and the bound aptamer structure cannot
compete with the more stable terminator hairpin. Correspondingly, RS8 exhibits a
constitutive low gene expression level. In contrast, terminators of RS2 and RS4, with
constitutive gene expression, are the least stable structures with -19.70 kcal/mol and
-20.60 kcal/mol, respectively. Here, the binding-competent structure of the aptamer is
probably more stable and remains intact even in the absence of theophylline,
interfering with terminator formation. The stabilities of terminators in functional
riboswitches RS1, RS3 and RS10 were between -21.00 and -25.80 kcal/mol, allowing
a theophylline-dependent switch from the unbound to the bound form. In agreement
with this correlation are the regulatory features observed for RS10loop, where the
optimization of the terminator hairpin leads to an increased stability. Due to this more
robust terminator structure (delta G?), RS10loop has a reduced background activity.
This is correlated with a lowered ON rate, as the ligand-bound aptamer structure now
has to compete with a more stable terminator element. Hence, RS10loop shows an
intermediate phenotype located between RS8 (most stable terminator with deltaG)
and T3 (less stable terminator with deltaG, allowing efficient structural
rearrangements upon theophylline binding). Although the adjusted loop sequences
are found in most of the naturally occurring intrinsic terminators {d'Aubenton Carafa 1990 #139}, one cannot transfer common features of stand-alone terminators into transcriptional riboswitches. Obviously, the requirements for structural rearrangements between ON and OFF states do not allow optimal terminator structures but favor a compromise between efficient termination and the possibility of riboswitch refolding upon ligand binding.

While these values show a good correlation with the regulatory effects of the synthetic riboswitches, they do not represent a general and irrevocable parameter for predicting a functional construct. The mutually exclusive RNA structures representing ON and OFF states depend on the relative stabilities of aptamer and terminator structures. Hence, the free energy values of these elements should be rather similar, allowing an easy switch between the different structures. Without the energy contribution of the bound ligand, the formation of the terminator hairpin should be the preferred structure, while binding of the ligand should lead to a stabilized aptamer, inhibiting terminator formation. As these energy values are not known for most of the available aptamers, it is not possible to predict a priori which of the designed terminators are suited for a certain aptamer domain to form a functional riboswitch. Here, biochemical analyses are required to verify a ligand-dependent regulation of transcription.

Taken together, our in silico design to construct functional synthetic riboswitches acting at the level of transcription is a first promising approach to generate and optimize sophisticated regulatory RNA control elements. The implication of further parameters like transcriptional pause sites and ligand binding behavior will lead to a more efficient regulatory power, comparable to that of the natural counterparts. Hence, these synthetic elements have the potential to develop into useful and versatile building blocks in synthetic biology.
Material and Methods

Riboswitch design

The rational design algorithm starts from the known sequence and secondary structure of the aptamer and constructs the sequence of the remaining part of the riboswitch, i.e. the spacer and the terminator. It proceeds by proposing a large set of candidates that are then evaluated with respect to their secondary structure and properties of their folding paths. At the proposal stage it incorporates some knowledge of sequence and structure of terminator elements for the given biological system in which the construct is supposed to function, here *E. coli*. In particular, we used the fact that the terminator hairpin must have a minimal size and that it is followed by a U-stretch. First, a spacer database containing randomly generated sequences with lengths between 6 and 20 nucleotides is created. The size of this database is user defined. For each position *k* of the 3’-part of the aptamer we form the sequence $\overline{X}_k$ that is complementary to the subsequence from *k* to the end of the aptamer. For the theophylline aptamer, values of *k* ranging from position 21 to 32 were used. The sequences of the aptamer, a spacer from the database, a complementary sequence $\overline{X}_k$, and the U-stretch are concatenated to form a complete riboswitch candidate.

Each of these constructs is then evaluated by folding simulations. In the current implementation, folding paths are represented as a sequence of secondary structures computed for sub-sequences starting at the 5’-end. We used individual transcription steps of 5 to 10 nucleotides to simulate co-transcriptional folding with varying elongation speed. Secondary structures are computed by RNAfold, a
component of the Vienna RNA Package (Lorenz 2011), with parameter settings -d2 --noPS --noLP. If one of the transcription intermediates forms base pairs between aptamer and spacer, it is likely that this will interfere with the ligand-binding properties; hence such a candidate is rejected. On the other hand, if the full length transcript does not contain a single hairpin structure composed of the aptamer 3’-part, the spacer and the sequence $\overline{X}_k$ complementary to the 3’-part of the aptamer, the construct cannot form a functional terminator and is also rejected. For each construct that passes these filters, additional features are calculated that are used as further selection criteria. The energy difference between the MFE (minimum free energy) fold of the full length construct and a structure where the aptamer region is constrained to form the ligand-binding competent fold is calculated. This parameter provides a rough measure for the stabilizing effect required to prevent terminator formation when the ligand is bound. Furthermore, a z-score of the structure downstream of the constrained aptamer fold is estimated as follows: The stability (MFE) of the candidate sequence is compared to the mean $\mu$ and standard deviation $\sigma$ of the folding energies computed for a set of 1000 di-nucleotide shuffled sequences. A positive value of the z-score ($z = (\text{MFE} - \mu)/\sigma$) indicates a structure less stable than expected by chance. This is used to estimate whether stable local structures downstream of the constrained aptamer fold might interfere with transcription.

The six experimentally tested synthetic riboswitch constructs are summarized in Figure 2A. Secondary structures and free energy values of individual sequences were calculated using RNAfold 2.0.7.

Given the dissociation constant $K_d$, the binding energy is given by: $\Delta G = RT \ln K_d$. With $T = 298$ K, $R = 1.98717$ cal/K and the experimentally determined value $K_d =$
0.32 μM {Jenison 1994 #46} for the theophylline aptamer, we obtain ΔG = -8.86 kcal/mol for the stabilization of the aptamer structure.

**Chemicals**

Oligonucleotides were purchased from biomers.net and dNTP were obtained from Jena Biosciences. LB medium was purchased from Applichem, theophylline and caffeine were obtained from Sigma-Aldrich. Ampicillin and Arabinose were purchased from Carl Roth.

**Plasmid construction**

Riboswitch constructs were generated by overlap extension PCR and fused to bgaB reporter in pBAD2_bgaB {Klinkert 2012 #137} by QuickChange site directed mutagenesis (Stratagene). Plasmid constructs were isolated using GeneJET™ Plasmid Miniprep Kit (Fermentas). Correctness of individual assemblies was verified by sequencing.

**Quantitative activity test for β-galactosidase**

Enzymatic activity of the expressed β-galactosidase was determined at 40°C {Waldminghaus 2007 #49}. All activity tests were performed in three independent experiments.

**Northern Blot analysis**

*E. coli* Top10 cells were grown in the presence or absence of 2 mM theophylline until OD\(_{600}\) of 0.5-0.6 in 25 ml LB medium supplied with ampicillin and arabinose. Total RNA was extracted according to J. Lawrence (pers. communication; http://rothlab.ucdavis.edu/protocols/D50.html).
Northern Blot was performed using the NorthernMax Formaldehyd-Based System for Northern blots (Ambion) according to manufacturer’s instructions. 5 µg total RNA were separated on 1 % agarose formaldehyde gel and transferred to BrightStar™-Plus Membrane (Ambion).

DNA oligonucleotides (reverse complementary to target sequences) were radioactively labeled using γ-³²P-ATP (Hartmann Analytic). The primer sequences used are:

bgaB: 5’-GTTCGATCTTGCCTCCAACTG-3’
23S rRNA: 5’-ACGACGGACGTACACCCCG-3’

Acknowledgements

The authors thank H. Betat, B. Klinkert and P. Kerpedjiev for valuable discussions and G. Domin for cloning RS8. Thanks to M. Mießler and S. Bonin for expert technical support. This work was supported by the Deutsche Forschungsgemeinschaft and the Christiane-Nüsslein-Vollhard-Stiftung.

References

Figure Legends

Fig. 1. Riboswitch constructs. (A) Design strategy for theophylline-dependent riboswitches controlling transcription. The sequence of the TCT8-4 theophylline aptamer (red) was fused to a short spacer region (cyan) followed by a sequence complementary to the 3’-part of the aptamer (blue) and a U-stretch (black). The ribosomal binding site (black box) and the open reading frame of the reporter gene bgaB are located immediately downstream of this construct. Upon theophylline
binding, terminator structure formation should be inhibited and transcription should proceed, resulting in expression of the reporter gene. (B) The final riboswitch constructs are transcribed from the arabinose promoter of plasmid pBAD. The transcription start point is indicated by the angled arrow. The ribosomal binding site (AGGA) of the reporter gene for β-galactosidase is located immediately downstream of the riboswitch insert. The start codon of the reporter gene is underlined.

**Fig. 2. The six experimentally tested riboswitch constructs.** (A) The theophylline aptamer represents the sensor (red), followed by a variable spacer sequence (cyan) and the 3'-part of the terminator (blue), forming a hairpin structure with a subsequence of the sensor domain. The terminator element is completed by a stretch of eight U residues (black). Below each construct, dot bracket notations of the secondary structures with terminator or antiterminator formation are indicated. The overlapping situation of sensor (theophylline aptamer, red box) and actuator element (terminator element, grey box) is required for a mutual exclusion of the two alternative structures. For each construct, calculated energy values of complete riboswitch elements (RS) as well as isolated terminators (T) are indicated. (B) Activity tests of the reporter enzyme β-galactosidase. Activities are indicated in Miller units (MU). All synthetic riboswitches were tested in the absence (black bars) and presence (grey bars) of 2mM theophylline. As a control, the structurally related caffeine was offered (white bars).

**Fig. 3. Optimized RS10 and RS3 variants and control constructs.** (A) In RS10loop, the tetraloop sequence of the terminator hairpin was adjusted to the frequently found sequence GAAA, including a closing G-C base pair. In RS10shift, a spacer sequence of 19 nt was inserted between the U₈ stretch of the terminator and
the ribosomal binding site (black box). The same 19 nt spacer was inserted in RS3shift at the identical position. In control bgaB, the upstream untranslated region of the reporter gene did not carry any riboswitch or terminator inserts. Furthermore, the U₈ stretch alone was placed immediately upstream of the ribosomal binding site and the 19 nt spacer sequence region was inserted into the original bgaB control construct, leading to bgaBU₈ and bgaBshift, respectively. (B) β-galactosidase activity test of the constructs described in (A). Activities are indicated in Miller units (MU). The tetraloop adaptation (RS10loop) in the terminator element led to a reduction of the background activity, but lowered also the expression of the reporter gene. Increasing the distance between the riboswitch U₈ stretch and the ribosomal binding site of the reporter gene in RS10shift increased the ON/OFF rate substantially, compared to the original RS10 construct. A similar effect was observed for RS3shift. In the controls, the U₈ stretch immediately upstream of the ribosomal binding site led to a rather high gene expression (bgaBU₈), while increasing the distance between these sequences reduced the β-galactosidase activity (bgaBshift).

Fig. 4. Functional analysis of synthetic terminator elements. (A) The isolated terminators of the tested riboswitch constructs were cloned upstream of the reporter gene. In a control construct, the 3'-complementary sequence of terminator 10 and the U₈ stretch was deleted, leading to RS10ΔT. As shown in Fig. 3, construct bgaBU₈ represents the positive control. (B) β-galactosidase activity of terminator constructs. While the positive control bgaBU₈ as well as the terminator deletion RS10ΔT show a strong enzyme activity, all terminators reduce the β-galactosidase activity at an efficiency comparable to that of the natural terminator from the pyrBI operon in E. coli (pyrBI). The terminator elements of the functional riboswitches 1, 3 and 10 show the strongest activity reduction. (C) Northern blot analysis of RS10. Upper panel: While
the terminator-alone construct T10 showed no transcription of the reporter gene, RS10 allowed bgaB gene transcription exclusively in the presence (+) of theophylline. C, control expression of bgaB using pBAD2-bgaB without riboswitch inserts. Lower panel: 23S rRNA was used as an internal standard for normalizing the presented lanes.

**Fig. 5. Terminator stability and riboswitch function.** Among all tested terminators, T8 has the lowest free energy value and forms a stable structure that obviously cannot be disrupted upon ligand binding. As a consequence, RS8 is locked in an OFF state. In contrast, T2 and T4 form a much weaker hairpin, shifting the equilibrium towards the aptamer structure that interferes with terminator formation. The functional riboswitches RS1, RS3 and RS10 show intermediate stabilities of their terminator elements that allow the desired ligand-dependent rearrangement of the constructs, switching between ON and OFF states. The stabilized terminator hairpin of RS10loop is located between the terminators of RS8 and the functional RS3. Although this stabilized hairpin reduces the background transcription rate considerably, it obviously impedes the structural rearrangement induced by ligand binding and allows only a moderate level of gene expression.
Fig. 1
### Table

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<td>-20.6</td>
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<tr>
<td>RS5</td>
<td>-16.7</td>
<td>-25.8</td>
</tr>
<tr>
<td>RS6</td>
<td>-17.3</td>
<td></td>
</tr>
<tr>
<td>RS7</td>
<td>-25.4</td>
<td>-29.0</td>
</tr>
<tr>
<td>RS8</td>
<td>-22.2</td>
<td></td>
</tr>
<tr>
<td>RS9</td>
<td>-28.3</td>
<td>-21.9</td>
</tr>
<tr>
<td>RS10</td>
<td>-15.1</td>
<td></td>
</tr>
</tbody>
</table>

### Diagram

**Fig. 2**
Fig. 3
Fig. 4

A

B

C

Fig. 4
Fig. 5

<table>
<thead>
<tr>
<th>Terminator</th>
<th>T8</th>
<th>T3</th>
<th>T10</th>
<th>T1</th>
<th>T4</th>
<th>T2</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\Delta G$ (kcal/mol)</td>
<td>-29.0</td>
<td>-25.8</td>
<td>-21.9</td>
<td>-21.0</td>
<td>-20.6</td>
<td>-19.7</td>
</tr>
</tbody>
</table>

Terminator Stability