Efficient and exclusive induction of Tet repressor by the oligopeptide Tip results from co-variation of their interaction site

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Abstract

Protein–protein interactions are an important element of signal transfer within and between organisms. They are mainly mediated by short oligopeptide motifs and represent a widely used alternative to small, organic molecules for conveying information. The transcription factor TetR, a regulator of tetracycline resistance in Gram-negative bacteria, is naturally induced by tetracycline or its derivatives. The oligopeptide Tip (Transcription inducing peptide) fused to either N- or C-terminus of Thioredoxin A (TrxA) has been isolated as an artificial inducer for TetR in Escherichia coli. This inducing property can be exploited to monitor the in vivo expression of a protein of interest by fusing Tip to its C-terminus. We improve the induction efficiency of Tip by adding an aromatic amino acid before residue 1 of Tip in C-terminal fusions to TrxA. The induction efficiency of that modified TrxA-Tip fusion is further enhanced when the effector-binding pocket of TetR is enlarged by the N82A and F86A mutations. The double mutant is also insensitive to induction by tetracyclines. Thus, Tip is an exclusive inducer of this TetR variant, representing the first example of fully converting a small molecular weight effector-dependent transcription factor into one depending solely on protein–protein recognition.

Introduction

Peptides are involved in many essential processes in signal transduction and cell-to-cell communication. Several prominent examples are the peptide hormones that bind to cell surface receptors triggering a signal cascade within the cell (1), peptides involved in innate immunity (2), or the quorum sensing peptides found in Gram-positive bacteria (3). Since methods like phage display or mRNA display facilitate the isolation of peptides that recognize protein targets, the number of synthetic peptides with novel biological functions is steadily and rapidly growing. These include enzyme and toxin inhibitors (4,5), peptides with antiviral activity (6,7) or with the ability to specifically detect tissues with histological changes associated with diseases (8,9). Peptides selected in this manner mostly exert their effect by binding to a specific target site.

Recently, a 16-mer peptide termed Tip (Transcription inducing peptide; Figure 1A) was isolated via phage display. Tip is not only able to bind specifically to its target, the bacterial transcription factor Tet repressor (TetR), but also triggers a conformational change in vivo normally induced by tetracyclines, the natural effectors of TetR (10). TetR is widely used to control regulation of gene expression in prokaryotes and, as a fusion protein with an activation domain, in eukaryotes (11), because it combines high specificity for its cognate DNA sequence (tetO) with extremely sensitive induction by tetracyclines (tc), especially the potent analogues doxycycline (dox) and anhydrotetracycline (atc) (12,13). Non-tc inducers of TetR, like Tip, may lead to alternative inducers when used as a scaffold for peptidomimetics to generate novel, small-molecule effectors. The recently published crystal structure of the TetR–Tip complex provides a molecular basis for such efforts (14). Another application of Tip is to fuse it to a protein of interest and use it to analyse target protein expression in vivo by monitoring a TetR-controlled reporter (15). Although a wide variety of tagsystems suitable for diverse applications exist, Tip is the only protein tag that allows quantitative in vivo analysis of protein expression. Mandatory for this application is a high efficiency of induction to ensure that low levels of Tip fusion proteins can be detected. We demonstrate here that addition of an aromatic residue between C-terminally fused Tip and TrxA along with mutations...
in the effector-binding pocket of TetR lead to a strong increase of reporter induction. Furthermore, the introduced TetR mutations render it insensitive to induction by tetracyclines, thereby creating exclusive specificity for Tip.

**MATERIALS AND METHODS**

**Materials and general methods**

Chemicals were obtained from Merck (Darmstadt), Sigma (Munich) or Roth (Karlsruhe) and were of the highest purity available. Media, buffers and solutions were prepared with Millipore water or deionized water and autoclaved. Heat labile substances were dissolved and purified with Millipore water or deionized water and autoclaved. Heat labile substances were dissolved and filtered with a sterile filter (0.2 μm). Enzymes for DNA restriction and modification were obtained from New England Biolabs (Frankfurt/M.), Roche Diagnostics (Mannheim), Invitrogen (Karlsruhe) or Fermentas (St. Leon-Rot). Sequencing was carried out according to the protocol provided by PE Applied Biosystems (Weiterstadt) for cycle sequencing.

**Bacterial strains, plasmids and phage**

All bacterial strains are derived from *Escherichia coli* K12. Strains DH5α (*hsdR17 (RK- mK+)*, recA1, endA1, gyrA96, thi, relA1, supE44, φ80lacZΔM15, Δ(lacZYA-argF)U169) (16) and RB791 ([IN[rndD-rndE]], lacFΔL8) (17) were used for general cloning procedures. Strain DH5α (λtet50) (Tn10 tetA-lacZ transcriptional fusion) (10,18) served as host strain for β-galactosidase assays. The plasmids pWH1200 (19), pWH527- and pWH1413-derivatives for TetR variant expression (10), pWH806 (20), pWH2100-, pWH2200- and pWH2300-derivatives for C- and N-terminal TrxA/SbmC-XTip variant expression (10) were used in the *in vivo* studies. Class B TetR (21) was used in all experiments.

**Insertion of Met in C-terminal TrxA-Tip fusion**

Insertion of Met into the C-terminal TrxA-Tip fusion was done by standard 2-step PCR using pWH2101 as template, the mutagenic primer 5′-GTCGGGTGGAGC TATGTGACTTGGAAATG-3′ to introduce the Met codon (underlined) and the flanking oligonucleotides 5′-TGACAATTAATCATCGGCTCG-3′ and 5′-AAGG AATGGTGACATGCCCTGC-3′. The product obtained was restricted with HindIII and PstI and cloned into likewise-digested pWH2101 (10). The resulting plasmid was named pWH2102.
Randomization of Met at position –1 of TrxA-MTip

Randomization of the Met at position –1 in TrxA(C)-MTip was performed by combined chain reaction (22). The phosphorylated oligonucleotide 5'-GTCGGGTGGAGCT(NNK)TGGATGGAATG-3' introducing the randomized codon (NNK: N = A, C, G or T; K = G or T) and the flanking oligonucleotides 5'-TGGACATTAATCATCGGCTCG-3' and 5'-AAGGAAATGGTGCATGCTGC-3' were used in a standard PCR reaction with pWH2102 as template. The resulting DNA was restricted with HindIII and PstI and cloned into likewise-digested pWH2102. Candidates were identified by sequencing and named pWH2102-X with X denoting the amino acid replacing the M.

Construction of the C-terminal SbmC(C)-Tip fusion

The construction of a plasmid-encoded SbmC(C)-Tip fusion was done by amplification of the fragment encoding the fusion from genomic DNA from E. coli BW25113(λtet50)sbmC(C)-tip, a strain containing the linker and Tip-encoding sequence as used in the TrxA fusion fused to the 3' end of the sbmC gene at its position in the E. coli genome (unpublished data) with the primers 5'-ATGATGCTCTAGAAACTACGAGATTAAGCAGGAAAGAAAAGC-3' introducing a restriction site for XbaI (underlined) and 5'-GTAGTTGCTAGCTTACGAAACC TTCACCACATGAGGC-3' introducing a restriction site for SpI (underlined). The product was restricted with XbaI and SpI and cloned into likewise-digested pWH610 (23). The resulting plasmid was named pWH2301.

Insertion of Met in the C-terminal SbmC(C)-Tip fusion

Insertion of Met (underlined) into the C-terminal SbmC-Tip fusion protein was done genetically by standard 2-step PCR using pWH2301 as template, a mutagenic primer 5'-CTCGGTTGGAGCTATGTGGACATT GGAATG-3' and the flanking primers 5'-ATGATGCTCTAGAAACTACGAGATTAAGCAGGAAAGAAAAGC-3' and 5'-GTAGTTGCTAGCTTACGAAACC TTCACCACATGAGGC-3', introducing restriction sites (underlined) for XbaI and SpI, respectively. The product obtained was cleaved with both enzymes and cloned into likewise-digested pWH610 (10). The resulting plasmid was named pWH2301-Met.

Construction of tetR-N82A-F86A

The F86A mutation was introduced into tetR-N82A encoded by pWH527-N82A via standard 2-step PCR using a mutagenic primer 5'-CTGAATAACGCTAAGATGATGATGCTTT-3' and the two flanking primers 5'-CTCGAGATCATTGTTACCG-3' and 5'-CGCGTACTGCCCCTTGG-3'. The resulting double mutant was cloned via restriction with NcoI and XbaI into pWH527 resulting in low-level constitutive expression of the TetR mutant (10).

β-Galactosidase (β-Gal) activity measurements

Repression and inducibility of TetR was assayed in E. coli DH5α(λtet50) (10). The strain was transformed with pWH527-derivatives expressing TetR variants at a low level or pWH1413-derivatives expressing TetR at a higher level and with plasmids from the pWH2100/2200/2300 series encoding the different C- and N-terminal Tip fusions to TrxA or SbmC. Overnight cultures and log-phase cultures were grown at 28°C in LB medium supplemented with 100 µg/ml ampicillin and either 60 µg/ml kanamycin for pWH527-derivatives or 25 µg/ml chloramphenicol for pWH1413-derivatives. For this purpose, stationary phase cultures were diluted 1:100 in fresh medium and expression of the fusion proteins induced using 60 µM IPTG, if not indicated otherwise. The cells were then grown to an OD600 of ~0.4 and their β-Gal activities determined as described by Miller (24). Three independent clones were assayed for each combination of constructs and experiments repeated at least twice. The values obtained were normalized to the maximal β-Gal activity in the absence of TetR which was set to 100% and typically varied from 6500 to 7500 Miller units in the individual experiments.

Western blot analysis of the TrxA-XTip fusion variants and the TetR variants

Escherichia coli DH5α(λtet50) was transformed with the derivatives listed in the respective β-Gal assays and grown under the same conditions as stated there. Cells were harvested at an OD600 of 0.4. The crude lysates were prepared by sonication and centrifugation. Ten microgram of crude lysate of each construct was loaded either on a 14% (TrxA-XTip fusions) or a 10% SDS-PAA gel (TetR variants) and electrophoresed. Proteins were transferred (120 mA, overnight at 4°C) to a PVDF membrane (BIORAD) in a Mini V8.10 blotting apparatus (Gibco-BRL) using 10 mM NaHCO3, 3 mM Na2CO3 and 20% (v/v) methanol as transfer buffer. After blocking with 0.2% I-Block™ (TROPIX) in phosphate-buffered saline (75 mM Na-phosphate; 68 mM NaCl; pH 7.8) with 0.1% Tween 20, membranes were treated either with a monoclonal anti-TrxA antibody (TrxA-XTip fusions), Anti-Thio™ antibody, Invitrogen) or a polyclonal TetR-specific antibody (SA1851; lab stock). Signals were detected with anti-mouse IgG (TrxA-XTip fusions) or anti-rabbit IgG (TetR variants) conjugated to horseradish peroxidase (Amersham Pharmacia) and the ECL+ kit (Amersham Pharmacia) following the manufacturer’s instructions.

RESULTS

Induction of TetR by C-terminal TrxA-XTip variants

Figure 1A shows the relevant features of the E. coli screening strain used in this study to score the efficiency of TetR induction by TrxA fusions with Tip. This strain has been described in detail (10). It contains one plasmid encoding TetR or its variants and LacI. TetR represses a chromosomally located lacZ, while LacI controls expression of the respective TrxA-Tip fusion protein. The latter is encoded on a second, compatible plasmid giving rise to IPTG-inducible β-Gal expression mediated by TrxA-Tip. For our studies, Tip was fused C- or N-terminally to
TrxA, and it had been observed that the latter is a much better inducer of TetR than the former (10) (see Figure 1B and C). A potential reason might be a sequence difference in Tip resulting from the failure of cleaving the initiator amino acid M from the N-terminus in E. coli when the following residue is W (see Figure 1B) (25,26). This M residue is missing in the C-terminal fusion. To examine this possibility we inserted an M residue into the C-terminal TrxA(C)-Tip fusion at the corresponding position termed −1 (see Figure 1B). This insertion leads to a nearly 2-fold increase in β-Gal activity, but still reaches only half of the β-Gal activity obtained with the N-terminal TrxA(C)-Tip fusion (see Figure 1C). The steady-state protein levels of all three TrxA fusion proteins are similar as concluded from Western blots (see Figure 1D) confirming that the induction levels are intrinsic properties of the fusions.

We next asked if M is the optimal amino acid at position −1 for induction of TetR by randomizing this residue. Western blot analyses were carried out with the resulting 19 TrxA(C)-XTip variants to make sure their expression levels are the same. Only TrxA(C)-PTip showed a reduced steady-state level while all other variants were present in similar amounts as TrxA(C)-MTip and TrxA(C)-Tip (data not shown). The TrxA(C)-XTip variants were then scored for their induction efficiencies in the E. coli screening strain (see Figure 1A). The results are summarized in Table 1. All activities of the TrxA(C)-XTip fusions are presented relative to 100% β-Gal activity defined by a strain lacking TetR. TrxA(C)-XTip variants with P, charged residues like D, E, K or R or with aliphatic, hydrophobic residues like V, L or I display either no or only marginal induction of TetR. All other amino acids at this position yield fusion proteins that induce TetR. They are either less active (Y), as active (S, T, C, F) or more active (G, A, N, Q, H, W) than the one with M.

### Table 1. Induction of TetR by all 20 possible variants of TrxA(C)-XTip

| TetR-inducing protein | β-Gal activity (%) |
|-----------------------|--------------------|
|                       | Repression          | Induction          |
| TrxA(C)-Tip           | 2.3 ± 0.1           | 24 ± 2             |
| TrxA(C)-MTip          | 2.6 ± 0.1           | 37 ± 3             |
| TrxA(C)-XTip          | 3.5 ± 0.4           | 57 ± 5             |
| Alanine               | 3.7 ± 0.2           | 63 ± 4             |
| Serine                | 2.7 ± 0.1           | 41 ± 3             |
| Threonine             | 2.4 ± 0.2           | 34 ± 3             |
| Cysteine              | 2.4 ± 0.2           | 36 ± 3             |
| Valine                | 2.2 ± 0.1           | 6.9 ± 0.5          |
| Isoleucine            | 2.6 ± 0.2           | 8.7 ± 1.2          |
| Leucine               | 2.1 ± 0.1           | 6.3 ± 0.6          |
| Aspartic Acid         | 2.2 ± 0.1           | 2.0 ± 0.1          |
| Asparagine            | 3.1 ± 0.1           | 46 ± 4             |
| Glutamic Acid         | 1.9 ± 0.2           | 2.3 ± 0.2          |
| Glutamine             | 3.8 ± 0.1           | 64 ± 3             |
| Lysine                | 2.1 ± 0.1           | 2.2 ± 0.1          |
| Arginine              | 2.0 ± 0.1           | 2.2 ± 0.1          |
| Histidine             | 3.4 ± 0.1           | 57 ± 4             |
| Phenylalanine         | 3.3 ± 0.2           | 38 ± 1             |
| Tyrosine              | 2.3 ± 0.1           | 21 ± 1             |
| Tryptophan            | 2.7 ± 0.1           | 55 ± 1             |
| Proline               | 2.0 ± 0.1           | 1.5 ± 0.4          |

### Induction of TetR-N82A, -F86A and -N82A-F86A by Tip variants

X-ray crystallography of TetR complexed with a 16-mer Tip oligopeptide acetylated at the N-terminus revealed details of the location of Tip inside the tc-binding pocket. The Cα atom of the acetyl moiety is in close proximity to residues N82 and F86 of TetR (14). Figure 2 shows an excerpt of the structure highlighting this proximity. We assumed that a large residue at this position, as found in several active C-terminal TrxA(C)-XTip variants, might lead to sterical hindrance. Therefore, we replaced the bulky N82 and F86 residues with a smaller A residue and scored the induction properties of these mutants with TrxA(C)-MTip. The results are shown in Figure 3. Even in the absence of IPTG, in which only basal expression of TrxA(C)-MTip from the leaky tac promoter occurs, we observed induction of TetR-N82A and TetR-F86A. However, TrxA(N)-Tip is still the more efficient inducer under these conditions. At the higher, IPTG-induced, expression level of TrxA(C)-MTip, induction of TetR-N82A reaches the same level as with TrxA(N)-Tip and wt TetR. A slightly smaller maximal induction by TrxA(C)-MTip is observed for TetR-F86A. As expected, altering the size of the binding pocket in TetR-N82A and TetR-F86A does not lead to increased induction by TrxA(C)-Tip lacking the additional M residue.

The induction properties of TrxA-Tip could depend in part on the TrxA portion of the fusion protein. To address this question, we constructed Tip and MTip fusions to the C-terminus of the E. coli protein SbmC (27) and replaced...
the TrxA-Tip fusion in our screening strain by them. SbmC is a small, soluble protein of 157 residues with solvent-exposed and flexible N- and C-termini (28). It is part of the SOS regulon and involved in reducing DNA damage (29). Induction of TetR by SbmC(C)-MTip is slightly more efficient than induction by SbmC(C)-Tip. In contrast, TetR-N82A is induced to a 5-fold higher level of β-Gal activity by SbmC(C)-MTip compared to SbmC(C)-Tip as shown in Figure 4. This result establishes an only marginal influence of the carrier protein on Tip activity. Hence, the improved induction of the TetR variants by the elongated Tip results solely from their effects on Tip–TetR interaction.

We wondered if the induction efficiency profiles of the TrxA(C)-XTip variants depend on the shape of the inducer-binding pocket. In addition to the TetR-N82A and -F86A mutants, we constructed the double mutant TetR-N82A-F86A and introduced that also into the screening strain. The Western blot shown as insert in Figure 5 indicates that TetR and TetR-F86A are expressed to about the same level, while TetR-N82A and TetR-N82A-F86A are present in roughly half the amount of the former variants. The induction activities were scored without IPTG at the basal expression levels of the TrxA(A(C)-XTip variants to increase the sensitivity and are shown in Figure 5. TrxA(C)-Tip shows about the same induction for TetR-N82A and TetR-F86A (~2% β-Gal activity), but the double mutant appears slightly de-repressed under these conditions (~4%). No other TrxA(C)-XTip variant induces TetR-N82A to a significantly higher degree than TrxA-MTip, and only TrxA(C)-FTip and TrxA(C)-WTip reach about the same induction efficiency (see the white columns in Figure 5). In contrast, TetR-F86A is very efficiently induced by TrxA-XTip variants with an aromatic residue. While TrxA-MTip shows only ~6% β-Gal activity, TrxA(C)-HTip and TrxA(A(C)-YTip lead to ~35% β-Gal activity, TrxA(C)-WTip gives rise to 53% and TrxA-FTip even to 60% β-Gal activity. The latter variant is hence the most efficient inducer of TetR-F86A. Similar to TetR-F86A, we observed the highest induction activities for TetR-N82A-F86A by TrxA-XTip derivatives with aromatic residues, albeit in a slightly changed order: TrxA-YTip exhibits the same induction efficiency for both TetR variants, while TrxA-HTip, TrxA-FTip and TrxA-WTip are better inducers for the double than for the single TetR mutant. Taken together, it appears that TrxA(C)-FTip and TrxA(C)-WTip are the best inducers for all three TetR variants, but their efficiencies increase from TetR-N82A via TetR-F86A to TetR-N82A-F86A. It is also noteworthy to mention that these induction profiles do not correlate generally to the differences in expression levels established for these three TetR variants.

The TetR residues N82 and F86 contact tc as derived from the crystal structure of the TetR-te complex (30). Hence, we tested whether the three TetR mutants in which one or both of these residues are replaced by A are inducible by tc, atc or dox, three widely used inducers of TetR. The results are shown in Figure 6. TetR-F86A is not induced by te, but atc and dox are very efficient inducers. In contrast, TetR-N82A and TetR-N82A-F86A are not induced by any of these compounds. Since these two are best induced by Tip we have created functionally novel TetR mutants, which will not respond to the most commonly used inducers atc and dox anymore, but instead are highly sensitive to Tip.
DISCUSSION

An additional Met preceding Tip increases its induction efficiency

TrxA(N)-Tip exhibits a much higher activity as inducer of TetR than TrxA(C)-Tip (10). We show here that part but not all of this activity difference is due to the fact that an additional M residue is present when Tip is N-terminal, owing to the fact that removal of the initiator M residue in E. coli is very inefficient before a W residue (26). The crystal structure of the TetR–Tip complex (14) reveals that residues W1 to N4 of Tip are located in the tc-binding pocket, thereby piercing the TetR core. We assume that an N-terminally fused Tip may enter the inducer-binding pocket more easily because the interacting residues can enter in a head first orientation. This is not possible for a C-terminal fusion of Tip, and hence, may contribute to the lower induction efficiency.

Random mutation of position –1 in TrxA(C)-XTip yields more efficient inducers of TetR

The M residue in position –1 of Tip is not the only, nor the optimal residue for increased induction of TetR. Indeed, randomizing the residue at this position in TrxA(C)-XTip revealed that several other amino acids also lead to an increase in TetR induction. The better induction seen with small residues like G, A or S could simply be interpreted as an extension of the linker sequence separating Tip from the TrxA scaffold, thereby increasing the flexibility of Tip. This simple explanation does not hold true for the increased induction seen...
when aromatic residues occupy position $-1$, as TrxA(C)-FTip is as active as TrxA(C)-MTip, and TrxA(C)-WTip and TrxA(C)-HTip are even slightly more active. Furthermore, TrxA(C)-DTip, TrxA(C)-ETip, TrxA(C)-KTip or TrxA(C)-RTip carrying charged residues exhibit no induction of TetR, while the uncharged, but similarly sized residues TrxA(C)-NTip and TrxA(C)-QTip are very good inducers of TetR. According to the crystal structure of the Tip–TetR complex, the N-terminus of Tip is located in a predominantly hydrophobic environment formed by residues L60, F67, F86, L90 and L142 of the tc-binding pocket. Placing a charged residue in the hydrophobic core of a protein should be very unfavourable (31,32). Hence, we conclude that the residue at position $-1$ must exert its function via contacts with the inducer-binding pocket of TetR.

There is a clear distinction among the hydrophobic amino acids between aromatic and aliphatic residues. C-terminal TrxA(C)-XTip variants with V, L or I show a decrease in activity compared to M, while variants with H, F, Y or W induce TetR more efficiently. Comparison of the contacts in the TetR inducer-binding pocket to tc and the first five residues of Tip (Figure 7) shows that most of the key interactions for tc (H64, N82, F86, H100, P105) are not formed with Tip (14). In particular, the space occupied by the A-ring of tc appears to be empty in the Tip–TetR complex. It is thus conceivable that the aromatic residues mimic the A-ring structure of tc best, followed by the flexible M residue with the easily polarizable sulphur, while the large hydrophobic residues fit least well.

TetR mutants with altered binding pocket are more efficiently induced by TrxA-XTip

Figures 2 and 7 display the proximity of the N-terminus of Tip to the residues N82 and F86 in the inducer-binding pocket of TetR. The replacement of these bulky residues by the much smaller A in single and double exchange mutants of TetR results in a dramatic increase of the induction activities of several TrxA(C)-XTip variants. These gain of function mutants reinforce the hypothesis that the residue at position $-1$ in TrxA(C)-XTip is located in the inducer-binding pocket facing the altered residues of the TetR variants. This conclusion is highlighted by the observation that TrxA(C)-XTip variants show different activity profiles for induction of the TetR-N82A, -F86A and -N82A-F86A mutants. We assume that this reflects the slightly different interactions of the various residues with the mutated tc-binding pockets.

The increase in in vivo induction efficiency of the TrxA(C)-XTip TetR variant pairs is indeed dramatic because nearly complete induction is obtained without having to relieve the LacI-mediated repression of TrxA(C)-XTip expression. Previous studies of the expression levels of TrxA(C)-Tip have revealed that the fusion protein is barely detectable in Western blots in the absence of IPTG (10). Since we also demonstrate that the induction efficiencies are not confined to TrxA fusions, we have created much more sensitive tools for monitoring the expression levels of proteins in general as compared to the one used previously (15). Furthermore, two of the TetR mutants are not inducible by tc, atc or dox anymore,
allowing their use in conjunction with tc inducible expression systems (11). Taken together, the data presented here suggest that proteins carrying a modified Tip as a C-terminal protein tag in combination with a reporter system under control of a TetR mutant allows highly sensitive detection of their expression levels. Inserting such a tag at the C-terminus of a protein is of great advantage over the N-terminus as one can expect less interference with translation initiation (33).

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Conflict of interest statement. None declared.

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