Antiviral Interferon-Beta Signaling Induced by Designed Transcription Activator-Like Effectors (TALE)

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Abstract

Here we show that designed transcription activator-like effectors (TALEs) that bind to defined areas of the interferon beta promoter are capable to induce IFN-beta expression and signaling in human cells. Importantly, TALE-mediated IFN-beta signaling occurs independently of pathogen pattern recognition but effectively prohibits viral RNA replication as demonstrated with a hepatitis C virus replicon. TALEs were thus indicated to be valuable tools in various applications addressing, for example, virus-host interactions.

Introduction

TALEs were originally characterized as virulence factors of plant pathogenic bacteria that reprogram gene transcription of the host cells. TALEs contain a DNA binding domain that is composed of similar tandem repeats of typically 34 amino acids. For transcription activation, each repeat binds one base pair of the target DNA, and a repeat-variable di-residue (RVD) specifies the bound base [1–4]. Thus, ‘designer TALEs’ containing a defined order of repeats and a suitable transcription activation domain can be constructed and applied to induce the transcription of human genes [5–7].

An attractive target for transcription activation is the cytokine IFN-beta, which is well characterized regarding its antiviral activity and also used during treatment of multiple sclerosis [8,9]. The IFN-beta promoter mainly consists of an enhancer that is flanked by two nucleosomes, one masking the transcriptional TATA-box (Fig. 1A).

The transcription of the IFN-beta gene is stringently regulated. During cell activation, which may be provoked by interactions of cellular pattern recognition
receptors (PRR) with pathogen associated molecular pattern (PAMP), signaling cascades induce the assembly of transcription and nucleosome remodeling factors (the so-called ‘enhanceosome’) at the enhancer’s positive regulatory domains (PRD; Fig. 1A). This, in turn, enables the association of the TATA-binding protein and Pol II-mediated transcription to initiate [10–13]. Accordingly, in non-activated cells (i.e., in the absence of the enhanceosome), the IFN-beta promoter was considered to be accessible for the binding and transcription-inducing activity of TALEs.

In this report we show that TALEs directed to bind to certain sites of the IFN-beta promoter induce an effective antiviral signaling cascade also in the absence of an external stimulus.

**Methods**

**Cloning of TALE genes**

Using a repeat library and the Golden TALE technology [5] six single-repeat modules were ligated into assembly vectors as a six-repeat array by cut-ligation using BpiI. The procedure was essentially performed as described by Geissler et al.
To assemble the complete TALE-coding sequence, three six-repeat arrays were ligated with modules encoding a green fluorescent protein (GFP) tag, the N- and C-terminus, and the herpes simplex virus VP16 transcription activation domain (C-terminal 68 aa), respectively, into a pcDNA3 (Invitrogen) derivative using BsaI. The amino acid sequence of the applied TALEs is given in Table S1 in File S1.

Construction of reporter plasmid

The plasmid was based on pF12A RM Flexi (Promega) where the barnase gene was replaced by a luciferase gene. To generate plasmids containing the TALE recognition sites, pF12A RM (Luc) was amplified with primers containing target boxes. The promoter and 5′ untranslated region (675 bp) of the IFN-beta1 promoter were amplified by PCR and inserted into the luciferase reporter plasmid [5].

In vitro transcription

The replicon-encoding plasmid pSGR-JFH1 was kindly provided by Dr. Wakita (Tokyo Metropolitan Institute for Neuroscience) and modified as described in [14]. The plasmid was digested with XbaI and transcribed by run-off in vitro transcription (standard protocol) with T7 RNA-polymerase (Stratagene) using the protocol of Geissler et al. 2012 [14].

Cell culturing and transfection conditions

Huh7 cells [14] were cultured in DMEM (Invitrogen) supplemented with 10% FCS (PAN-Biotech), 1% penicillin/streptomycin (Invitrogen), 0.1% d-Biotin and 0.1% hypoxanthin (Sigma) [15]. Transfection of plasmids was performed with 70% confluent cells, 20 μg plasmid DNA/10 ml growth medium using Turbofect (Fermentas) and the manufacturer’s instructions. The replicon RNA was transfected using 300 ng (ca. 100 fmol) and the Bio-Rad Gene Pulser II (1 pulse without controller at 0.2 kV and 950 μF).

Western blot

The western-blot analysis was performed at 24 h p.t. of the TALE-expressing plasmids. Ca. 2 × 10⁶ cells were centrifuged at 1000 × g, washed with phosphate buffer saline and lysed by 1x lysis buffer (Promega). The protein amount was determined by a standard Bradford assay (BioRad) and 20 μg of total cell protein separated per lane by SDS-PAGE. Following the transfer to nitrocellulose (Millipore), the reaction was carried out using standard conditions and the following antibodies: anti-GFP (Invitrogen A-6455) 1:2000, anti-GAPDH (Santa Cruz sc-47724) 1:15000; secondary antibodies (each at 1:5000 dilution) anti-rabbit (Licor Cw 800 926-32213), anti-mouse (Licor Cw 800 926-32212).
RT-PCR analysis

For qRT-PCR analysis, RNA was isolated from ca. $2 \times 10^6$ cells with Trizol at the indicated time points and the qRT-PCR performed using Revert Aid reverse transcriptase (Thermo) and the PCR-MasterMix qRT (Roboklon GmbH, Germany). For reverse transcription of total mRNA, an oligodT$_{19}$ primer was applied; for reverse transcription of the HCV replicon JFH, we applied the primer JFH reverse (ACA TGA TCT GCA GAG AGA CCA G). The RT conditions were 1 h at 42°C (amounts of applied RNA 500 ng); the PCR conditions were 94°C, 15 sec; 60°C, 25 sec; 72°C 25 sec $\times$ 40 cycles. The HCV RNA levels were normalized to GAPDH RNA as internal control. For further details, see Geissler et al. [14]. All applied DNA oligonucleotides (purchased from Eurofins, Germany) are summarized in Table S2 in File S1.

Data evaluation and statistics

Data evaluation and statistics were done as described previously [14].

Ethics statement

Not applicable.

Results and Discussion

Following earlier work, which revealed the general option to induce IFN-beta expression in human cells by a TALE [5], this study aimed at investigating if an entire antiviral IFN-beta signaling cascade may be navigated by designed effectors. For this purpose, we generated a set of six TALE-expressing constructs applying the ‘Golden TALE technology’ [5]. The corresponding effectors were designed to bind to different regions of the IFN-beta promoter (see Fig. 1A and Table 1).

After transient transfection of the expression plasmids, all six effectors were comparably expressed in Huh7 cells (human hepatoma cells). This was demonstrated by western-blot that detected the TALEs via a fused green fluorescent protein (GFP) reporter (Fig. 1B). In a subsequent experiment, we expressed the individual TALEs in Huh7 cells and measured the amount of IFN-beta mRNA by qRT-PCR (Fig. 1C). This data revealed that with promoter-associating TALEs, the transcription of the IFN-beta gene was up to 4-fold increased in comparison to experiments with a non-related TALE. Interestingly, most effective were TALEs1 and 6 that were binding to the sites of the promoter that were indicated to be covered by nucleosomes (Fig. 1A and C). This suggests that the effectors, besides attracting the Pol II transcription machinery, may also facilitate nucleosomal remodeling.

Secreted IFNs function by binding to the IFN receptor (IFNAR) of neighboring cells and by activating the canonical JAK/STAT pathway. This leads to the formation of interferon-stimulated gene factor complexes (ISGF3) that drive the transcription of interferon-stimulated genes (ISGs). ISGs encode antiviral proteins.
like OAS (2'-5'-oligoadenylate synthetase), MX (GTPase), ADAR (adenosine deaminase) and signaling proteins as the interferon regulatory factor IRF7 [16–18]. To understand next if IFN-beta expression correlated with ISG expression, we performed qRT-PCR that measured the mRNA levels of IFN-beta side-by-side with those of OAS1 and 2, MX1, IRF7 and ADAR1. This was done with two cell types, Huh7 and Huh7.5. Huh7.5 differ from Huh7 such that RIG-I (retinoic acid-inducible gene I), an important intracellular PRR [19] is defective in these cells [20, 21]. Thus, with the pathogen hepatitis C virus (HCV) it is well understood that PAMP as the tri-phosphate at the viral RNA’s 5’-end and an HCV-specific RNA motif in the 3’ untranslated region (3’UTR) of the viral genome are recognized by RIG-I and that RIG-I-induced signaling cascades lead to the activation of transcription factors that regulate IFN gene expression [22–25]. To compare HCV- and TALE-induced IFN-beta signalling, both cell types were transfected either with an HCV subgenomic RNA replicon [26] (Fig. 2A) or with the TALE6-expressing plasmid. Interestingly, as shown in Fig. 2C, TALE6 induced the expression of IFN-beta and ISGs in both cell types, while replicating HCV RNA did so only in Huh7 (Fig. 2B). Controls performed with a non-replicating HCV RNA or a non-specific TALE showed neither an induction of IFN-beta nor of ISGs (Fig. 2B and C). This data highlights that TALE6 triggered the entire IFN-beta signaling cascade. Most importantly, with the experiment applying Huh7.5 cells, we confirmed that TALE-induced IFN-beta signaling occurred independently of PAMP signaling.

In a final experiment, we addressed the question if the TALE6-mediated induction of IFN-beta and ISGs was capable to interfere with viral RNA replication. For this, Huh7 and Huh7.5 cells were transfected with the TALE6-encoding plasmid and, 24 h later, with the HCV replicon. As shown in Fig. 2D, prior expression of TALE6 effectively inhibited viral RNA replication while expression of a non-related TALE did not. A considerable inhibitory effect on viral replication was also observed in cells when we expressed TALE6 in Huh7 cells where HCV replication was already established (Fig. 2E).

The activation of IFN-beta gene expression via PRR-PAMP signaling is a well-studied model of how transcriptional output is regulated in the cell. Here we demonstrate that TALEs, by operating directly on the transcriptional level, bypass

| TALE | target sequence |
|------|----------------|
| control | T CGG TCT GGC TTG ACA TGA |
| 1 | T TAA ATA AAG AGT TTT AGA |
| 2 | T AGA AAC TAC TAA AAT GTA |
| 3 | T AGG AAA ACT GAA AGG GAG |
| 4 | T GGG AAA TTC CTC TGA AT |
| 5 | T CTC ATA TAA ATA GGC CAT |
| 6 | T CTC ATA TAA ATA GGC CAT |

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Table 1. TALE binding sites within the IFN-beta promoter (see also Fig. 1A).
PAMP/RIG-I-mediated signaling. Moreover, TALE-triggered IFN-beta signaling effectively prohibits HCV replication in hepatoma cells. Thus, in comparison to the IFN-beta response that is induced by the HCV replicon in Huh-7 cells (Fig. 2B), the TALE6-stimulated IFN-beta and ISG expression (Fig. 2C) leads to a clearly detectable decrease in the level of viral RNA (Fig. 2D). We explained this by the multiple ways of how viral factors, which are also encoded by the applied replicon may inhibit IFN signaling [27]. For example, the HCV NS3/4A protease
inhibits HCV PAMP-RIG-I signaling by proteolytic degradation of a RIG-I signaling adapter [28].

Taken together, our data recommend IFN-beta inducing TALEs as potentially valuable tools for future vaccination or treatment applications. Specifically designed TALEs may also be helpful to unravel the function of yet insufficiently characterized host factors participating in (antiviral) cell signaling or viral replication.

Supporting Information

File S1. Supporting tables. Table S1, Organization and amino acid sequences of applied TALEs. Table S2, DNA oligonucleotides applied for qRT-PCR.
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Author Contributions

Conceived and designed the experiments: IR RG SF SEB. Performed the experiments: IR NF RG SF AP. Analyzed the data: IR NF RG SF SEB. Contributed to the writing of the manuscript: SEB.

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