Transcriptional Activators of Human Genes with Programmable DNA-Specificity

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Abstract

TAL (transcription activator-like) effectors are translocated by Xanthomonas bacteria into plant cells where they activate transcription of target genes. DNA target sequence recognition occurs in a unique mode involving a central domain of tandem repeats. Each repeat recognizes a single base pair in a contiguous DNA sequence and a pair of adjacent hypervariable amino acid residues per repeat specifies which base is bound. Rearranging the repeats allows the design of novel TAL proteins with predictable DNA-recognition specificities. TAL protein-based transcriptional activation in plant cells is mediated by a C-terminal activation domain (AD). Here, we created synthetic TAL proteins with designed repeat compositions using a novel modular cloning strategy termed “Golden TAL Technology”. Newly programmed TAL proteins were not only functional in plant cells, but also in human cells and activated targeted expression of exogenous as well as endogenous genes. Transcriptional activation in different human cell lines was markedly improved by replacing the TAL-AD with the VP16-AD of herpes simplex virus. The creation of TAL proteins with potentially any desired DNA-recognition specificity allows their versatile use in biotechnology.

Introduction

Transcription activator-like (TAL) effectors include key virulence factors of Xanthomonas that bind to promoter regions of plant genes and act as DNA sequence-specific transcriptional activators [1,2,3,4]. As a typical feature, TAL effectors contain a central domain of tandem repeats (1 to 33.5 repeats of typically 34 amino acids) [1]. First shown for the archetype TAL effector AvrBs3, this repeat domain is essential for DNA-binding [5,6] and represents a novel, modular type of DNA-binding domain [5]. One repeat corresponds to one DNA base pair, and the specificity of each repeat is encoded by two hypervariable amino acids (position 12 and 15) per repeat, also termed repeat-variable diresidue (RVD) [7,8]. The last repeat contains only the first 20 conserved residues including the RVDs and is referred to as a half repeat. Each repeat functions neighbor-independently, and the linear order of repeats defines the matching DNA-sequence. In addition, the target box is extended by a 5’ T [1,2,7,8]. Based on the repeat-specificity code, the target DNA specificities of several TAL effectors were correctly predicted [7,9,10]. While the number of repeats greatly varies in TAL effector family members, at least 10.5 repeats are required for maximal activity [7]. Accordingly, TAL effectors with different numbers of repeats (e.g. Hax2, 21.5 repeats and Hax3, 11.5 repeats) [11] show comparable transcriptional activation in reporter assays [7]. The modular architecture, a hallmark of the TAL protein repeat domain, enables simple rearrangements of desired repeat orders. Thus, TAL proteins with novel and clearly predictable DNA-recognition specificities can be constructed [7,12,13,14,15,16,17].

The possibility of generating proteins with programmed DNA-binding specificity is an exciting avenue to targeted genome editing and gene regulation. For these purposes, zinc finger (ZF) proteins that contain an array of ZFs targeting a given DNA sequence are already in use [18,19]. Specifically, ZF-nucleases representing fusions between ZF proteins and the nuclease domain of the restriction enzyme FokI were applied to induce insertions and deletions at specific sites in complex genomes [19,20]. Compared to ZF proteins, the DNA-binding specificities of TAL proteins are considerably easier to predict [1,19]. Thus, TAL-nucleases were recently generated that cut specific DNA sites [12,13,14,21,22].

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acidic transcriptional activation domains (ADs) [23,24,25]. So far, this notion was supported by infection experiments with Xanthomonas strains delivering TAL effectors where the C-terminus was deleted or substituted by the AD of the herpes simplex virus (HSV) transcription activator VP16. Monitored via elicited plant responses and in yeast reporter assays, TAL effector activity was inhibited without AD, while it was partly restored by the heterologous VP16-AD [23,24,25,26].

Construction of TAL proteins with ordered repeats is challenging due to the highly repetitive nature of the repeats. We developed a modular cloning strategy to easily assemble TAL proteins and tested the TAL-dependent modulation of gene expression in human cells. We show that native TAL proteins function as transcriptional activators in different human cell lines and that this activity can be significantly enhanced by substituting the endogenous AD by the VP16 AD. Importantly, specifically designed TAL effector derivatives were shown to induce the expression of an exogenous reporter gene as well as of target endogenous genes.

Results

The TAL protein-AD can be functionally replaced by ADs from other organisms

To generate TAL proteins with a potential transcription factor activity in non-plant organisms, we deleted the ADs from the TAL effectors AvrBs3 and Hax3 and replaced them with the ADs of the HSV VP16 and the yeast GAL4 transcription activators, respectively (Figure S1A). All TAL proteins were designed such that they contained an N-terminal tag of green fluorescent protein (GFP). Following heterologous expression, all TAL proteins were confirmed to localize to plant nuclei indicating functionality of their nuclear localization signals (NLSs) and thus protein integrity (Figure S2). To determine the transcription factor activity of the TAL effector-derivatives in plant cells, a reporter assay was set up. Target DNA sequences specifically recognized by AvrBs3 and Hax3 (termed AvrBs3- and Hax3-box, respectively) were inserted upstream of a minimal promoter which has low background activity [27] into a reporter vector containing a promoterless uidA (β-glucuronidase, GUS) reporter gene [7] (Figures 1A and S1B). The reporter constructs were co-transfected into Nicotiana benthamiana leaf cells together with constructs that mediated the expression of the avrBs3- and hax3-derivatives, respectively (Figure 1A).

Quantification of GUS activity revealed that AvrBs3 and Hax3 only induced promoters that contained the AvrBs3- and Hax3-box, respectively. As expected, deletion of the ADs strongly reduced the transcription factor activity of both effectors (Figure 1B,C). Addition of the VP16-AD and GAL4-AD led to a slight increase of the AvrBs3- and Hax3-activities in planta; however, activity was still substantially lower than that of the original TAL proteins (Figure 1B,C). These data show that AvrBs3 and Hax3 still contain approx. 15% of their activity after deletion of the AD, and that the VP16-AD only partially complements the natural TAL protein-AD in planta.

TAL protein-directed transcriptional activation in human cells

Plant-infecting xanthomonads are not pathogenic to humans and there is no indication that the bacteria can translocate TAL effectors into cells of non-plant organisms. TAL proteins were nevertheless expected to specifically interact and to even function with matching DNA-sequences in non-plant organisms. This notion was fuelled by the observation that in planta as well as in vitro, the specificity of TAL protein-DNA interactions is solely dependent on the TAL protein repeat composition [5,6,10,28].

Moreover, there are no indications for the existence of kingdom specific eukaryotic host factors contributing to TAL protein functions.

To test if TAL proteins were able to direct gene expression in human cells, we established reporter constructs that contained the AvrBs3- or Hax3-box upstream of a minimal human cytomegalovirus (CMV) promoter and a luciferase (luc) reporter gene (Figures 2A and S1B). When either reporter plasmid was co-transfected into human HeLa293T cells together with a plasmid that expressed GFP via a fully active CMV promoter, only weak luciferase activity was detectable. This confirmed that the minimal CMV promoter only has a low basal activity (Figure 2B,C). In contrast, co-expression of native AvrBs3 or Hax3 resulted in a specific activation of promoters with matching target DNA boxes, respectively (Figure 2B,C). Interestingly, the

![Figure 1. The VP16 or GAL4 activation domains (ADs) can partially complement the endogenous ADs of AvrBs3 and Hax3 in planta. (A) Schematic drawing of reporter construct and co-transfection experiment. Target AvrBs3 and Hax3 boxes were inserted upstream of the minimal Bsp4 promoter into a GUS reporter vector [7]. GUS reporter and constructs driving expression of TAL genes were co-delivered via Agrobacterium into N. benthamiana leaf cells. (B) Specific induction of the AvrBs3 box by AvrBs3, an AD deletion mutant (ΔAD), the ΔAD mutant complemented with the VP16-AD (VP16), and the ΔAD mutant complemented with the GAL4-AD (GAL4). (C) Specific induction of the Hax3 box by Hax3 and AD-derivatives. (B, C) GFP and constitutively expressed GUS served as negative and positive controls, respectively. Samples were taken 2 dpi (error bars indicate SD; n = 3). GUS activity was measured with MUG (4-methylumbelliferyl-β-D-glucuronide) as substrate. doi:10.1371/journal.pone.0019509.g001]
transcriptional activation was low when we applied AvrBs3- and Hax3-derivatives with deleted ADs. Conversely, AvrBs3 and Hax3 that contained a VP16-AD caused a strong transcriptional activation of the luciferase reporter gene, i.e., transcription activation was 2–3 times higher than that induced by the original (in planta active) AvrBs3 and Hax3 and 50–80 times higher than transcription in the negative control (Figure 2B,C). In comparison, the GAL4-AD did not increase the activity of the δAD-TAL proteins significantly. Importantly, all expressed TAL proteins were found to localize to the nucleus of the human cells indicating correct nuclear import in the heterologous host (Figure S3A). Protein levels of TAL effector-derivatives expressed in human cells were comparable (Figure S3B,C). Our data show that TAL proteins are capable to specifically induce gene expression in human cells and that substitution of the endogenous AD by a VP16-AD strongly increases transcriptional activity.

Next, we analyzed whether the TAL protein-directed transcriptional activation was functional in different human cell lines. As also shown in Figure 2D, analogous experiments that were performed with TAL-VP16 derivatives and reporter constructs in Hek293T-Rex (human embryonic kidney) cells, Huh7 (hepatocarcinoma) cells, and HeLa (cervical cancer) cells led in each case to an evident and specific induction of reporter gene activity in matching TAL protein-target box combinations (Figure 2D). In sum, these results demonstrate that TAL effector-derivatives are functional as transcriptional activators in different human cells.

A modular cloning strategy to assemble TAL proteins with designed repeat architecture

To target defined DNA sequences and individual promoters with TAL proteins it is required to control their order of repeats. Unfortunately, the repetitive nature of TAL effector repeats makes it inherently complicated to use classical or PCR-based cloning techniques to generate TAL proteins with a given order of repeats. For that reason, we established a rapid and flexible cloning technique (Figure 3 and S4, see Text S1 for details).

Single TAL protein repeats were cloned as individual modules with flanking BsaI restriction sites. The type IIIs sites were designed to generate specific overhangs that determine the position of the repeat in a repeat array (Figure 3A) while preserving the amino acid sequence. For each repeat position, four key repeats (NI, HD, NN, NG) were cloned that specify the four DNA base pairs. Using the BsaI cut-ligation reaction, one to six repeats can be combined into an assembly vector that contained additional flanking BsaI sites which specify the position of the repeat assembly in the final repeat domain (Figure 3B). In addition, BsaI-flanked modules of the TAL protein N-terminal region and C-terminal region fused to VP16, and an N-terminal GFP module were designed. In a BsaI cut-ligation reaction, these modules can be specifically aligned into a vector backbone (Figure 3C). With a plasmid library of only 29 repeat modules and seven different assembly vectors, synthetic TAL proteins with 1 to 24 specific repeats can be generated in a matter of days. The modular technology also allows a simple integration of an expanded module toolbox with different TAL protein termini, tags, and vectors. Since this toolbox is essentially based on golden gate cloning [29,30], we termed this cloning technique “Golden TAL Technology”.

Designed TAL proteins direct the expression of endogenous human genes

Next, we considered it important to evaluate whether synthetic TAL proteins were also applicable to activate the transcription and thus the expression of different endogenous human genes.

Figure 2. AvrBs3 and Hax3 activate transcription in human cells. (A) Schematic drawing of reporter construct and co-transfection experiment. TAL boxes were cloned upstream of a minimal cytomegalovirus (CMV) promoter into a luciferase (luc) reporter vector. The λ operators are bound by a chimeric transactivator used as control. The luc reporter plasmids were co-transfected into human Hek293T-Rex cells together with expression plasmids where transcription of the TAL or GFP genes was driven by a fully active CMV promoter. (B) Specific induction of the AvrBs3 box by AvrBs3, an AD deletion mutant (δAD), the δAD mutant complemented with the VP16-AD (VP16), and the δAD mutant complemented with the GAL4-AD (GAL4). (C) Specific induction of the Hax3 box by Hax3 and AD-derivatives. (D) Comparison of induction levels of the Luc reporter plasmids using Hax3 with VP16-AD or via co-transfection of a plasmid expressing a chimeric transactivator for the λ operator and addition of the inducer coumermycin (light grey bars), respectively, in Hek293T-Rex cells. (E) Specific activity of AvrBs3 and Hax3 containing the VP16-AD, respectively, in human Huh7, and HeLa cells. (B-E) Measurements were done 1 dpi. Luc activity was plotted in arbitrary light units per μg protein. Dark grey bars: AvrBs3, white bars: Hax3. The activation is statistically significant (relative to GFP) with P≤0.005 for all values except AvrBs3-δAD, P≤0.01. doi:10.1371/journal.pone.0019509.g002
HuTAL expression plasmids resulted in a strong (22- to 34-fold) and specific induction of reporter activity only in matching TAL protein-target DNA box combinations (Figure 4A). Thus, we confirmed the specificity and functionality of the HuTALs.

Next, we transfected Hek293T-Rex cells with the CMV promoter-driven HuTAL genes to address if we could also affect the expression of endogenous human genes by the HuTALs. These experiments revealed that expression of the respective HuTAL in each case caused moderately increased levels of target endogenous human gene transcript levels (Figure 4B). While PUMA transcript levels were only slightly induced (1.5 x), the IFNα1 and IFNβ1 levels were 3.1-fold and 3.5-fold up-regulated, respectively (Figure 4B).

TAL protein-directed induction of endogenous gene expression was considerably lower than with our reporter constructs. This prompted us to determine if the promoter per se has an influence on TAL protein functionality. Hence, we replaced the minimal CMV promoter in the reporter plasmid by the native IFNβ1 promoter sequence. Co-transfection of the IFNβ1 reporter construct with the HuTAL targeting this promoter led to a moderate, 3.2-fold induction of reporter activity (Fig. 4C).

Accordingly, we applied the Golden TAL Technology to generate artificial TAL proteins with repeats that matched DNA sequences in promoter regions of the human PUMA (p53 upregulated modulator of apoptosis), an interferon alpha (IFNα1), and an interferon beta (IFNβ1) gene, respectively (Figure S5A). PUMA is a pro-apoptotic member of the Bcl-2 protein family [31,32], whereas IFNα1 and IFNβ1 belong to type I interferons that are involved in the innate immune response in humans against viral infections [33,34]. To efficiently direct human gene expression, the designed “human TAL proteins” (HuTALs) contained a C-terminal VP16 activation domain. Protein levels of HuTALs expressed in human cells were comparable and the proteins localized to the nucleus of human cells (Figure S5B,C).

As an important pilot experiment, we first inserted the predicted target DNA boxes of the HuTALs into the reporter construct described above, i.e., upstream of the minimal CMV promoter and the promoterless luc gene (Figure 2A). Co-transfection of human Hek293T-Rex cells with the reporter constructs and
comparable to the level of endogenous gene induction. This suggests that the efficiency of TAL proteins also depends on the individual target promoter. Taken together, our experiments show that TAL protein-derivatives can be used to specifically induce expression of human endogenous genes and demonstrate the functionality of the TAL protein technology in other organisms besides plants.

Discussion

In this report we demonstrate that TAL protein derivatives can be used to specifically modulate the transcription of genes in human cells. Our findings confirm and extend the recent work of Miller et al. [14] and Zhang et al. [17]. In addition to those studies, we report here that native Xanthomonas TAL effectors (AvrBs3 and Hax3) exhibited considerable activity in the heterologous host cells. This is surprising considering that these are proteins from bacterial origin that were evolutionary shaped to function in plant cells. Two scenarios are conceivable to interpret this finding. On one hand, the mechanism that is applied by TAL proteins to activate the host’s transcription machinery may be based on the protein structure itself. Alternatively, TAL proteins may rely on elements that are conserved in the different hosts. Hence, it will be interesting to see if TAL proteins function ubiquitously in all eukaryotes or if there is a specific host determinant required for activity.

Deletion of the endogenous TAL protein activation domain (AD) strongly compromised transcriptional activation, but a measurable residual activity remained (Figs. 1 and 2). This finding supports two conclusions. First, the endogenous TAL-AD is clearly a major determinant of TAL protein function also in heterologous hosts, and, second, TAL proteins have a low but specific residual transcriptional activation activity, irrespective of the presence of the AD. This has not been demonstrated before, and suggests that this low activity is based on TAL-DNA interactions, which are not influenced by the AD [5], or possible AD-independent interactions with components of the host transcription machinery.

The substitution of the endogenous AD by the heterologous HSV VP16-AD strongly elevated the activity of AvrBs3 and Hax3 AD derivatives in human cells but not in plant cells (Figs. 1, 2 and 4). The human virus AD is adapted to function in human cells, and this function can be transferred to TAL proteins. Miller et al. [14] and Zhang et al. [17] applied artificial TAL proteins, where the VP16-AD was fused to full-length or C-terminally truncated TAL proteins. The TAL proteins designed by Miller et al. [14] and Zhang et al. [17] as well as our synthetic TAL proteins each activated gene expression in human cells indicating that the VP16-AD functions in two ways, as a substitute or as a supplement of the endogenous TAL-AD. In addition, we showed that TAL-VP16 fusions are active in different human cell lines (Fig. 2D). The TAL protein activity strongly varied between HEK cells showing highest and HeLa cells showing lowest transcriptional activation, respectively. Likely, this is in part due to different transfection efficiencies that were achieved with the different cell lines. Further studies will clarify if tissue-specific requirements exist or whether different TAL protein backbones (N- and C-termini) function differently well in human cells.

We designed the Golden TAL Technology as a toolbox for a flexible assembly of TAL proteins with designed repeats, different N-, C-termini, and tags. While our method is somewhat similar to the recently published method of Zhang et al. [17], it includes a number of aspects that we consider advantageous for the construction of designer TAL proteins. First, our procedure does not rely on PCR but instead uses plasmid sub-clones of single repeats which is user-friendly and prohibits PCR-based errors. Furthermore, Zhang et al. [17] altered the codon usage between TAL protein repeats to allow efficient PCR, while we preferred to keep the codon usage near-identical between repeats to reduce repeat-specific effects on translation. Second, our library consists of only 24 single-repeat plasmids, instead of the 48 PCR-library of Zhang et al. [17]. Third, our method allows construction of TAL proteins with a flexible number of repeats. At present TAL proteins with 1 to 24 repeats can be constructed based on assembly modules containing six or less individual repeats. Fourth, we designed the restriction overhangs such that they differ each at a minimum of two positions. We found this to be essential for an efficient error-free ligation. Finally, we employ the full power of Golden Gate cloning which is a fast and versatile method to assemble many fragments specifically in a one-step cut-ligation reaction [29,30]. To acknowledge this, we termed our cloning strategy Golden TAL Technology.

Taking advantage of the Golden TAL Technology we were able to straightforwardly construct artificial TAL proteins with VP16-AD and repeats that match to promoters of various human target genes. These tailored TAL proteins (HuTALs) strongly induced reporter gene activation more than 30-fold and also endogenous human PUMA, IFNα1, and IFNβ1 mRNA levels. Endogenous gene induction was considerably weaker (1.5-fold to 3.5-fold), a phenomenon which was also observed with artificial ZF transcription factors [35,36]. The efficiency of TAL protein-dependent endogenous human gene induction in the two recent studies varied. While Miller et al. [14] activated endogenous NTF3 expression 20-fold, Zhang et al. [17] induced endogenous human SOX2 and KLF4 genes only 2.2 and 5.5 fold, respectively, or were not able to change expression levels of c-MYC and OCT4 at all. This observation makes sense considering that in the transient transfection experiments, multiple reporter gene copies are applied, whereas the endogenous genes are present at much lower copy-number. Moreover, the epigenetic environment of the endogenous genes (nucleosome packaging, transcriptional interference with other regulatory factors, etc.) may significantly affect the activity of HuTALs. This may also explain, why in the case of the PUMA gene no phenotype (accelerated cell death) could be observed. Hence, future optimization of the HuTAL system will be necessary to allow efficient regulation of endogenous human gene expression.

A growing collection of studies shows that the programmable DNA-binding specificity of TAL proteins renders these proteins similarly useful as the ZF technology [19] for different biotechnology applications. Our work and the recently published ones [14,17] are first crucial kick-off steps to use TAL proteins as highly potent and specific gene switches in human cells.

Materials and Methods

Plant growth and inoculations

Nicotiana benthamiana plants were cultivated in the greenhouse with day and night temperatures of 23°C and 19°C, respectively, 16 h light, and 40 to 60% humidity. Leaves of five- to seven-week-old plants were inoculated with an Agrobacterium suspension using a needleless syringe [7]. Inoculated plants were transferred to a Percival growth chamber (Percival Scientific) with day and night temperatures of 22°C and 18°C, respectively, and 16 h light.

Cell culture and transfection

HeK293T-Rex, HeLa and Huh7 cells were grown under standard conditions [37] in DMEM (Invitrogen) supplemented with 10% FCS (Invitrogen), 1% non-essential amino acids and 1%
Plasmids

For ADD constructs, the last 31 amino acids (aa) of AvrBs3 and Hax3 were deleted, respectively. 68 aa of VP16, or 113 aa of GAL4 were added as heterologous activation domains. Constructs for TAL expression in planta and in human cells were based on pVS300F [38] and pcDNA5 (Invitrogen), respectively. Reporter plasmids for plant cells were constructed as described [7]. Reporter plasmids for human cells were based on pF12A RM Flexi (Promega). The barnase gene of pF12A RM Flexi was replaced by a luciferase gene. To generate plasmids containing the recognition site of TALs, pF12A RM Flexi (Luc) was amplified with primers containing target boxes. The promoter and 5’ untranscribed region (675 bp) of the IFNβ promoter were amplified by PCR and inserted into the luciferase reporter plasmid. Further details on constructs and methods are given in the supplementary information and upon request.

GUS assays

GUS assays from plant samples were essentially done as described [7]. Briefly, Agrobacterium strains delivering TAL effector constructs and GUS reporter constructs were mixed in equal amounts, and inoculated into Nicotiana benthamiana leaves with an OD600 of 0.8. Two days post infiltration (dpi), two leaf discs (0.9 cm diameter) were sampled and quantitative GUS activity was determined using 4-methyl-umbelliferyl-β-D-glucuronide (MUG). Proteins were quantified by Bradford assays (BioRad). Details on constructs and methods are given in the supplementary information and upon request.

qRT-PCR, western blotting and luciferase assay

Human cells were harvested 24 h after transfection with plasmids. For RNA extraction, a standard protocol was used as described [37]. RNA was heated for 5 min to 70°C and reverse transcription of DNaseI (Roche), EcoRI, and BglII (Fermentas) treated RNA (2 μg) was performed by RevertAid H Minus Reverse Transcriptase (Fermentas) for 1 h at 42°C, followed by heat inactivation for 10 min at 70°C as described in the manufacturer’s manual. Quantitative PCRs (for primers see Table S1) were performed with the QuantiTect SYBR Green PCR Kit (Qiagen) according to the manufacturer’s manual using a LightCycler 2.0 instrument (Roche). PCR conditions were: 95°C 15 min, (95°C, 10 sec; 60°C, 25 sec; 72°C 25 sec) x 45 cycles. The qRT-PCR data were normalized to GAPDH as internal control and analyzed by the ddCt method [39]. For western blotting total protein extracts were prepared with Luciferase Cell Culture Lysis Reagent (Promega) and analyzed by western blotting for GFP (α-GFP, Invitrogen) and Vinculin (hVinc-1, Sigma-Aldrich). Luciferase assays were prepared as described in the Luciferase Assay System (Promega). The data points were generated from three biological replicates (separate transfections).

Golden TAL Technology

For details on individual modules, vectors, and cloning strategy see Text S1 and Figure S4. Briefly, single repeats were PCR-amplified with flanking BpiI sites (for primers see Table S1) and subcloned by cut-ligation using Smal and T4-DNA ligase into a pUC57-derivative (Amp) with mutated BsaI site. The assembly vectors [Kan] contain a lacZ gene flanked by BpiI sites which is replaced in cloning step 1 by the inserted repeats and used for blue-white selection of correct assembly products. BsaI-flanked modules are assembled in cloning step 2 into a target expression vector. Assembly reactions were set up as cut-ligation with 50-100 ng of each plasmid module, 1 μl BstI (NEB), 2 μl ATP (10 mM), 2 μl restriction buffer no. 4 (NEB), 1 μl T4-DNA ligase (5 u/μl) in a 20 μl reaction for 1 h at 37°C followed by 20 min. inactivation of enzymes at 70°C.

Supporting Information

Text S1 TAL gene expression constructs, reporter constructs, the Golden TAL Technology toolbox, and promoter regions of human genes chosen as targets for TAL protein-directed expression. (DOC)

Figure S1 TAL effector-derivatives and reporter constructs. (A) Functional domains of TAL effector-derivatives used. The endogenous C-terminal transcriptional activation domain (AD: C-terminal 31 aa) was deleted or replaced by the ADs from herpes simplex virus VP16 (C-terminal 68 aa) or yeast GAL4 (C-terminal 113 aa). NLS: nuclear localization signals; red: repeat domain of AvrBs3 and derivatives; yellow: repeat domain of Hax3 and derivatives. All constructs carried an N-terminal green fluorescent protein (GFP) tag. (B) Reporter constructs used for transient expression in plant and human cells, respectively. For experiments in plants, TAL protein target DNA boxes were inserted in front of a minimal Bs4 promoter which has low basal activity upstream of a promoterless uidA (GUS) reporter gene as described [4]. For studies with human cells, TAL protein target DNA boxes were placed upstream of the minimal CMV promoter and a promoterless luciferase (Luc) gene into a pF12A RM Flexi (Promega) reporter vector. To study activity of the endogenous interferon β promoter, the minimal CMV promoter was replaced by part of the INFβ promoter. (TIF)

Figure S2 GFP-TAL protein fusions localized to the plant nucleus. GFP-TAL protein fusions or GFP alone were expressed transiently in leaf cells of Nicotiana benthamiana via Agrobacterium-mediated delivery. Two days post infiltration, N. benthamiana leaf epidermis cells were stained with DAPI (4′,6-diamidino-2-phenylindole) and analyzed by confocal laser scanning microscopy. Green: GFP fluorescence; blue: DAPI fluorescence; red: chlorophyll autofluorescence. Scale bars indicate 50 μm. (TIF)

Figure S3 GFP-TAL protein fusions are expressed and localize to the nucleus in human cells. (A) GFP-TAL protein fusions or GFP alone were expressed after transfection of human Hek293T-Rex cells. One day post transfection, cells were stained with DAPI (4′,6-diamidino-2-phenylindole) and analyzed by fluorescence microscopy. Green: GFP fluorescence; blue: DAPI fluorescence. (B) Western-blot analysis of whole cell extracts of transfected Hek293T-Rex cells. (C) TAL protein-derivatives expressed in HeLa cells show degradation patterns. (Br-C) GFP and GFP-TAL protein fusions were detected using α-GFP antibody. An α-Vinculin antibody was used as constitutive loading control. (TIF)

Figure S4 Golden TAL Technology toolbox. Single repeat modules were subcloned with flanking BpiI sites that specify their position in the repeat array. A library of four repeat types (NI = A, HD = C, NN = G, NG = T) was constructed for each of the six repeat positions. Stop repeats (1s to 5s) can be used to terminate the repeat assembly. In cloning step 1, up to six repeats are inserted into an assembly vector to generate a repeat assembly. Different assembly vectors are used to position the repeat assembly within the final TAL protein repeat domain. In cloning step 2, individual repeat assemblies, Hax3 N- and
C-termini, and an N-terminal GFP-tag are ligated into an expression vector to generate the complete TAL gene.

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**Figure S5** HuTALs designed to target expression of human genes. (A) HuTALs with 17.5 to 19.5 repeats matching a DNA sequence (box) in the promoter region of target genes are assembled as N-terminal GFP fusions using the “Golden TAL technology”. (B) GFP-TAL protein fusions were expressed after transfection of human Hek293T-Rex cells. One day post-transfection, cells were stained with DAPI (4′,6-diamidino-2-phenylindole) and analyzed by fluorescence microscopy. Green: GFP fluorescence; blue: DAPI fluorescence. (C) Western-blot analysis of whole cell extracts of Hek293T-Rex cells transfected with HuTAL expression constructs.

**(TIF)**

**References**

1. Boch J, Bonas U (2010) Xanthomonas AvrBs3 family-type III effectors: discovery and function. Annu Rev Phytopathol 48: 419–436.

2. Bogdanove AJ, Schornack S, Lahaye T (2010) TAL effectors: finding plant genes for disease resistance. Curr Opin Microbiol 13: 394–401.

3. White FF, Yang B (2009) Host and pathogen factors controlling the rice Xanthomonas 13-33guc peg interaction. Plant Physiol 150: 1677–1686.

4. Scholze H, Boch J (2011) TAL effectors are remote controls for gene activation. Curr Opin Microbiol 14: 292–300.

5. Kay S, Hahn S, Marois E, Hause G, Bonas U (2007) A bacterial effector acts as a plant transcription factor and induces a cell size regulator. Science 318: 648–651.

6. Römer P, Hahn S, Jordan T, Strauß T, Bonas U, et al. (2007) Plant-pathogen recognition mediated by promoter activation of the pepper Bp resistance gene. Science 318: 645–648.

7. Boch J, Scholze H, Schornack S, Landgraf A, Hahn S, et al. (2009) Breaking the code of DNA binding specificity of TAL-type III effectors. Science 326: 1509–1513.

8. Moscou MJ, Bogdanove AJ (2009) A simple cipher governs DNA recognition by TAL effectors. Science 326: 1501.

9. Antony G, Zhou J, Huang S, Li T, Liu B, et al. (2010) Rice sa13 recessive resistance to bacterial blight is defeated by induction of the disease susceptibility gene Os-In3. Plant Cell 22: 3864–3876.

10. Römer P, Recht S, Strauß T, Elaeesser J, Schornack S, et al. (2010) Promoter elements of rice susceptibility genes are bound and activated by specific TAL effectors from the bacterial blight pathogen, Xanthomonas oryzae pv. oryzae. New Phytol 187: 1048–1057.

11. Kay S, Boch J, Bonas U (2005) Characterization of AvrBs3like effectors from a B. savannaea pathogen reveals virulence and avirulence activities and a protein with a novel repeat architecture. Mol Plant-Microbe Interact 18: 838–848.

12. Christian M, Cermak T, Doyle EL, Schmidt C, Zhang F, et al. (2010) TAL nucleases with novel DNA binding specificity create double-strand breaks. Proc Natl Acad Sci USA 93: 1757–1761.

13. Mahfouz MM, Li L, Shamimuzzaman M, Wibowo A, Fang X, et al. (2011) De novo-engineered transcription activator-like effector (TALE) hybrid nuclease with novel DNA binding specificity creates double-strand breaks. Proc Natl Acad Sci USA 108: 4165–4170.

14. Zhu W, Yang B, Chittoo JM, Johnson LB, White FF (1998) AvrXa10 contains an acidic transcriptional activation domain in the functionally conserved C terminus. Mol Plant-Microbe Interact 11: 824–832.

15. Zhu W, Yang B, Willis N, Johnson LB, White FF (1999) The C terminus of AvrXa10 can be replaced by the transcriptional activation domain of VP16 from the herpes simplex virus. Plant Cell 11: 1665–1674.

16. Wysoczka J, Herr W (2003) The herpes simplex virus VP16-induced complex: the makings of a regulatory switch. Trends Biochem Sci 28: 969–973.

17. Schornack S, Peter K, Bonas U, Lahaye T (2005) Expression levels of avrBs3-like genes affect recognition specificity in tomato B.粥 but not in pepper B.粥-mediated perception. Mol Plant-Microbe Interact 18: 1215–1223.

18. Kay S, Hahn S, Marois E, Wiedwald R, Bonas U (2009) Detailed analysis of the DNA recognition motifs of the Xanthomonas type III effectors AvrB3 and AvrB3rep16. Plant J 59: 859–871.

19. Engler C, Gruetzmacher K, Kandria R, Marillonnet S (2009) Golden gate shuffling: a one-pot DNA shuffling method based on type IIS restriction enzymes. PLoS ONE 4: e5355.

20. Engler C, Kandria R, Marillonnet S (2008) A one pot, one step, precision cloning method with high throughput capability. PLoS ONE 3: e647.

21. Nakano K, Yoshida KH (2003) PUMA, a novel proapoptotic gene, is induced by p53. Mol Cell 7: 683–694.

22. Yu J, Zhang L, Hwang PM, Kinzler KW, Vogelstein B (2001) PUMA induces cell cycle arrest of mammalian cells. Mol Cell 7: 673–682.

23. Prinz M, Kalenka U (2010) New lessons about old molecules: how type I interferons shape Th1/Th17-mediated autoimmunity in the CNS. Trends Mol Med 16: 379–386.

24. Vilec J (2006) Fifty years of interferon research: aiming at a moving target. Immunity 25: 343–348.

25. Park KS, Seol W, Yang HY, Lee SJ, Kim SK, et al. (2005) Identification and use of zinc finger transcription factors that increase production of recombinant proteins in yeast and mammalian cells. Biotechnol Prog 21: 664–670.

26. Reik A, Zhou Y, Collingwood TN, Warle, I, Bartsevich V, et al. (2007) Enhanced protein production by engineered zinc finger proteins. Biotechnol Bioeng 97: 1180–1189.

27. Isken O, Baroth M, Grassmann CW, Weinlich S, Ostareck DH, et al. (2007) Nuclear factors are involved in hepatitis C virus RNA replication. RNA 13: 1675–1692.

28. Van den Ackerveken G, Marois E, Bonas U (1996) Recognition of the bacterial avirulence protein AvrBs3 occurs inside the host plant cell. Cell 87: 1675–1692.

29. Mahfouz MM, Li L, Shamimuzzaman M, Wibowo A, Fang X, et al. (2011) De novo-engineered transcription activator-like effector (TALE) hybrid nuclease with novel DNA binding specificity creates double-strand breaks. Proc Natl Acad Sci USA 108: 4165–4170.

30. Schornack S, Peter K, Bonas U, Lahaye T (2005) Expression levels of avrBs3-like genes affect recognition specificity in tomato B.粥 but not in pepper B.粥-mediated perception. Mol Plant-Microbe Interact 18: 1215–1223.

31. Kay S, Hahn S, Marois E, Wiedwald R, Bonas U (2009) Detailed analysis of the DNA recognition motifs of the Xanthomonas type III effectors AvrB3 and AvrB3rep16. Plant J 59: 859–871.

32. Engler C, Gruetzmacher K, Kandria R, Marillonnet S (2009) Golden gate shuffling: a one-pot DNA shuffling method based on type IIS restriction enzymes. PLoS ONE 4: e5355.

33. Engler C, Kandria R, Marillonnet S (2008) A one pot, one step, precision cloning method with high throughput capability. PLoS ONE 3: e647.

34. Nakano K, Yoshida KH (2003) PUMA, a novel proapoptotic gene, is induced by p53. Mol Cell 7: 683–694.

35. Yu J, Zhang L, Hwang PM, Kinzler KW, Vogelstein B (2001) PUMA induces cell cycle arrest of mammalian cells. Mol Cell 7: 673–682.

36. Prinz M, Kalenka U (2010) New lessons about old molecules: how type I interferons shape Th1/Th17-mediated autoimmunity in the CNS. Trends Mol Med 16: 379–386.

37. Vilec J (2006) Fifty years of interferon research: aiming at a moving target. Immunity 25: 343–348.

38. Park KS, Seol W, Yang HY, Lee SJ, Kim SK, et al. (2005) Identification and use of zinc finger transcription factors that increase production of recombinant proteins in yeast and mammalian cells. Biotechnol Prog 21: 664–670.

39. Reik A, Zhou Y, Collingwood TN, Warle, I, Bartsevich V, et al. (2007) Enhanced protein production by engineered zinc finger proteins. Biotechnol Bioeng 97: 1180–1189.

40. Isken O, Baroth M, Grassmann CW, Weinlich S, Ostareck DH, et al. (2007) Nuclear factors are involved in hepatitis C virus RNA replication. RNA 13: 1675–1692.

41. Van den Ackerveken G, Marois E, Bonas U (1996) Recognition of the bacterial avirulence protein AvrBs3 occurs inside the host plant cell. Cell 87: 1675–1692.

42. Livak KJ, Schmittgen TD (2001) Analysis of relative gene expression data using real-time quantitative PCR and the 2(-Delta Delta C(T)) Method. Methods 25: 402–408.