Zinc-finger-based transcriptional repression of rhodopsin in a model of dominant retinitis pigmentosa

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Despite the recent success of gene-based complementation approaches for genetic recessive traits, the development of therapeutic strategies for gain-of-function mutations poses great challenges. General therapeutic principles to correct these genetic defects mostly rely on post-transcriptional gene regulation (RNA silencing). Engineered zinc-finger (ZF) protein-based repression of transcription may represent a novel approach for treating gain-of-function mutations, although proof-of-concept of this use is still lacking. Here, we generated a series of transcriptional repressors to silence human rhodopsin (hRHO), the gene most abundantly expressed in retinal photoreceptors. The strategy was designed to suppress both the mutated and the wild-type hRHO allele in a mutational-independent fashion, to overcome mutational heterogeneity of autosomal dominant retinitis pigmentosa due to hRHO mutations. Here we demonstrate that ZF proteins promote a robust transcriptional repression of hRHO in a transgenic mouse model of autosomal dominant retinitis pigmentosa. Furthermore, we show that specifically decreasing the mutated human RHO transcript in conjunction with unaltered expression of the endogenous murine Rho gene results in amelioration of disease progression, as demonstrated by significant improvements in retinal morphology and function. This zinc-finger-based mutation-independent approach paves the way towards a ‘repression–replacement’ strategy, which is expected to facilitate widespread applications in the development of novel therapeutics for a variety of disorders that are due to gain-of-function mutations.

INTRODUCTION

Proof-of-concept studies and the recent success of clinical trials are providing sound evidence for the use of gene-based complementation therapeutic strategies to treat genetic recessive traits (Aiuti et al, 2009; Bainbridge et al, 2008; Cartier et al, 2009; Hauswirth et al, 2008; Kaplitt et al, 2007; Maguire et al, 2009, 2008). Nevertheless, the design of effective therapeutic strategies for dominant genetic traits still poses difficulties. The common therapeutic approach to correct these genetic entities relies on the silencing of gene expression. The main target of genetic silencing strategies is the messenger RNA (mRNA) transcript, the function of which can be inhibited by antisense-
RNA-based, ribozyme-based and more recently by small-interfering (si)RNA-based and micro (mi)RNA-based approaches. In particular, RNA interference (RNAi) has great promise for treating dominant diseases in both mutation-dependent and -independent manners, through its efficiency of mRNA transcript cleavage (LaVail et al, 2000; Lewin et al, 1998; O’Reilly et al, 2007; Xia et al, 2004). A possible alternative to such RNA-targeting approaches is the modulation of gene expression at the transcriptional level, by using zinc-finger (ZF)-based artificial transcription factors (ZF-ATFs) that can be tailored to a desired DNA target sequence.

Artificial ZF proteins (ZFPs) are becoming a novel and powerful technological platform for both gene manipulation and development of therapeutics (Jamieson et al, 2003; Pearson, 2008; Segal & Barbas, 2001). ZFPs are composed of a DNA-binding domain (DBD) that is based on the Cys2His2 ZF scaffold fused with a transcriptional regulation domain. Their modular structure enables both the sequential assembling of multiple ZFs to generate DBDs with different target specificities and the use of various effector domains to engineer ATFs or nucleases. A single ZF domain consists of approximately 30 amino acids with a simple ββα fold that is stabilized by hydrophobic interactions and the chelation of a single zinc ion. Each ZF module primarily recognizes an overlapping 4-bp DNA sequence, where the last base pair is the first of the following target (the fourth base of each target is on the opposite strand). The binding takes place through key amino-acid residues, which can be exchanged to generate ZF modules with different sequence specificities. To obtain a DBD that is tailored to a unique target sequence in mammalian genomic DNA (genome size in humans, 3.0 x 10⁹ bp), theoretically, a sequence longer than 16 bp is needed, and this can be achieved by consecutive linking of at least six ZF modules.

To date, several functional ZF-ATFs have been generated to modulate target gene expression in vitro (Bartesvich & Juliano, 2000; Beerli et al, 1998; Liu et al, 2001; Zhang et al, 2000); however, none of the in vivo studies presented (Mattei et al, 2007; Rebar et al, 2002) have been aimed at silencing a disease gene via vector-mediated somatic-gene transfer.

We designed a two-step repression–replacement strategy (Cashman et al, 2005; Chadderton et al, 2009; Farrar et al, 2002; Gorbatyuk et al, 2005; Kiang et al, 2005; O’Reilly et al, 2007; Xia et al, 2004): (i) mutational-independent silencing of the human rhodopsin (hRHO) gene (transcriptional silencing targeted to both wild-type and mutated RHO alleles); and (ii) gene replacement of the endogenous RHO copies by adeno-associated virus (AAV)-vector-mediated photoreceptor gene transfer. Considering that autosomal dominant retinitis pigmentosa (adRP) is the most genetically heterogeneous inherited disease in humans, we designed this mutational-independent strategy to overcome the technical and economical magnitude of allele-specific targeted-designed therapeutics. Indeed, in the case of adRP, due to rhodopsin mutations, more than 150 allele-specific silencing molecules would be required to silence each specific mutation identified thus far (RHO gain-of-function mutations account for 25–50% of the total autosomal dominant adRP cases; Inglehearn et al, 1998; Sohocki et al, 2001).

Here, we set out to determine whether transcriptional repression by engineered ZFP technology represents a novel therapeutic gene-silencing paradigm for the treatment of gain-of-function mutations. To this end, we used a transgenic mouse model of adRP harbouring a Pro347Ser (P347S) mutation in hRHO.

We show that AAV-vector-mediated photoreceptor delivery of a selected humanized ZF artificial transcriptional repressor targeted to the mutated hRHO transgene in P347S adRP mice results in significant reduction of its expression, which in turn leads to improved retinal pathology and function, which validates the first limiting step of the repression–replacement strategy designed.

RESULTS

Design and generation of zinc-finger-based transcription factors to control rhodopsin gene expression

To control RHO expression, we designed ZF-ATFs targeted to the human rhodopsin promoter (hRHO-P). We first scanned the proximal hRHO-P to identify amenable target regions, applying the following criteria: (i) exclusivity of the 18-bp DNA sequences in the human genome; (ii) putative accessibility to the chromatin environment [in proximity to known endogenous transcription-factor binding sites; (Graslund et al, 2005)]; (iii) absence of annotated polymorphisms; and (iv) divergence between the human and murine rhodopsin promoter target sequences. We selected 10 different continuous or discontinuous 18-bp target sites within the proximal region of the hRHO-P (from −90 to −32, relative to the transcription start site; Fig 1a and Table 1) that fulfilled these four criteria. As individual ZFs typically bind 3 or 4 bp of DNA, we generated DBDs by assembling six-ZF arrays that were directed to the selected 18-bp DNA sequences, using the modular assembly method (http://www.zincfingertool.org; Beerli et al, 1998; Mandell & Barbas, 2006; Segal et al, 1999). Due to the lack of availability of ZFs for all possible triplets (Blancafort et al, 2003; Dreier et al, 2001, 2005, 2000; Segal et al, 1999) and to enhanced DNA-binding affinity (Kim & Pabo, 1998), we designed 10 ZF modules with two arms, each composed of three ZF units (arm 1 encoding fingers 1.2.3 and arm 2 encoding fingers 4.5.6) that were bridged together by linker peptides that allowed these ZF modules to span up to 10 bp of DNA gaps between their target-recognition sites (Fig 1b). For gaps longer than 3 bp, we used the structured linker reported by Moore (Moore et al, 2001), whereas for shorter gaps or continuous sequences, we used the flexible linker reported by Rebar (Rebar et al, 2002; Table 1). We next fused the ZF DBDs to a nuclear localization signal, to a C-terminal HA-tag, and to effector domains that included either the herpes simplex virus-based transcriptional activator VP64 domain (Seipel et al, 1992; ZF transcriptional activators; ZF-As) or the human-derived Krüppel-associated box (KRAB) repression domain (Margolin et al, 1994; ZF transcriptional repressors: ZF-Rs; Fig 1b).

In vitro screening of functional ZF-Rs

To select ZF-ATFs that are functional in hRHO-P transcriptional control, we first performed firefly luciferase reporter-gene
transactivation assays by co-transfecting HEK293 cells with both the reporter gene driven by the human rhodopsin proximal promoter (−330 to +32, relative to the transcription start site) and the individual ZF-A. Two out of the 10 constructs significantly transactivated luciferase expression to levels comparable to those obtained with a plasmid coding for the endogenous rhodopsin transcription activator CRX (cone-rod-homeobox-containing gene) (Chen et al., 1997; Rehemtulla et al., 1996). The shift was not observed using nuclear extract from cells expressing ZF-R6-shuffled (lane 2). Specific binding was abolished upon addition of 10× or 100× molar excess of cold probe to the mix (lanes 5 and 6), whereas no effects can be seen upon addition of 10× or 100× molar excess of an unspecific cold probe (lane 7 and 8). The supershift is obtained by adding an antibody that recognizes the HA-tag of the ZF-R6 in the mix (lane 9).

mediated transcription through triple transfection in HEK293 cells, which included the ZF-Rs, the CRX and the reporter plasmids. Notably, ZF-R2 and ZF-R6, which contain the same DBDs as ZF-A2 and ZF-A6, significantly reduced luciferase expression levels (81% and 64% repression relative to CRX transactivation, respectively; Fig 1d; n = 5, from three independent experiments; ZF-R2, p < 0.001; ZF-R6, p < 0.01), similar to the repression obtained with the CRX DBD fused to KRAB, which was used as the positive control (Chau et al., 2000).
ZF-R-mediated repression of human rhodopsin in retinal stem cells

To probe whether the two selected ZF-ATFs promote rhodopsin transcriptional repression in the chromosomal context, we took advantage of an in vitro system to differentiate retinal precursors (retinal stem cells; RSCs (Giordano et al., 2007)). Neurons derived from the murine adult ocular ciliary margin can differentiate in vitro into various retinal neuronal cell types, including photoreceptor precursors that express rhodopsin, the beta-subunit of phosphodiesterase-6 (PDE6b), and the cGMP-gated channel (Cnga) (Sanges et al., 2006). We explanted neurons from the adult P347S transgenic mouse. This mouse model of adRP harbours a human rhodopsin transcriptional unit (including 4 kb of the proximal rhodopsin promoter region) from a patient carrying a P347S mutation (Li et al., 1996), in addition to the endogenous wild-type murine rhodopsin alleles (mRho). Once differentiated into photoreceptor-like (rhodopsin positive; Rho+) cells, P347S neurons underwent apoptosis with kinetics similar to those observed in another severe retinal-degeneration model [the rd-1 mouse; (Sanges et al., 2006)]. Virtually all of the Rho+ cells were also TUNEL-positive 12 days after adding differentiating medium to the P347S retinal cell precursors (Figure S3 of Supporting Information). Thus, we expected that efficient and specific repression of the human mutated rhodopsin, combined with preservation of the endogenous murine rhodopsin expression, would result in rescue/survival of the photoreceptor precursors.

To this end, 7 days after adding the differentiating medium, we transduced P347S retinal precursor cells with retroviruses coding for the selected ZF-Rs (ZF-R2 and ZF-R6) or for ZF-R6-shuffled as a negative control. Twelve days after the induction of differentiation, ~70% of the cells expressed rhodopsin (i.e. were Rho+), 98% of which were also double positive (i.e. HA+, Rho+), confirming that retroviruses can transduce retinal cell precursors efficiently (Sanges et al., 2006). ZF-R2 and ZF-R6 transduction produced significant 77 and 92% protection from cell death, respectively, whereas virtually all of the ZF-R6-shuffled transduced cells underwent apoptosis (Fig 2a, b and d; n = 2, from two independent experiments; ZF-R2, \( p = 4.54 \times 10^{-11} \); ZF-R6, \( p = 1.18 \times 10^{-15} \)). Confocal images of triple-stained (HA+, Rho+ and TUNEL+) samples confirmed that only cells expressing ZF-R2 or ZF-R6 were protected from apoptosis (Fig 2c).

To determine whether protection from apoptosis was through specific repression of human rhodopsin transcription, we measured hRHO mRNA levels 4 days after transduction. Nonetheless, ZF-R2- and ZF-R6-transduced cells showed 51 and 88% reduction, respectively, of the hRHO transcript levels, compared to hRHO levels in cells treated with ZF-R6-shuffled (Fig 2e; n = 2, from two independent experiments; ZF-R2, \( p < 0.01 \); ZF-R6, \( p < 0.01 \)) after normalization with murine rhodopsin, which was used to discriminate photoreceptor precursor cells (Rho+ cells) from those which were not induced to express rhodopsin. These data show that ZFPs promote hRHO transcriptional repression in a chromosomal context and that this, in turn, can protect retinal progenitors of an adRP model from apoptosis.

**Delivery of ZF-Rs to murine photoreceptors decreases adRP disease progression**

The ZF-R6 construct, which provided robust hRHO transcriptional repression and apoptosis protection, was selected for in vivo experiments. We generated an AAV2/8 vector, as this is currently considered to be most efficient AAV delivery vehicle for photoreceptor gene transfer (Allocca et al., 2007; Lebherz et al., 2008; Natkunarajah et al., 2008; Stieger et al., 2008). This AAV2/8 vector contained the ZF-R6 construct under the photoreceptor-specific rhodopsin kinase (RHOK) promoter element (Allocca et al., 2007) (AAV2/8-RHOK-ZF-R6). In homozygous P347S mice, over-expression of the mutated rhodopsin leads to the fast progression of retinal degeneration. Indeed, by P21, only one row of photoreceptor nuclei was detectable by histology (Figure S4 of Supporting Information). Thus, to decrease human P347S rhodopsin expression levels, we crossed P347S mice with a wild-type C57BL/6 background. The resulting progeny (P347S+/−) showed milder retinal degeneration (Figure S4 of Supporting Information). At P70, P347S+/− retinae still retained 5–6 rows of photoreceptor nuclei and residual functionality, thus allowing evaluation of disease progression over time by both histology and electroretinography.
ERG) analysis. We first assessed transcriptional repression by subretinal co-injection in P4 P347S+/−/− of 1 × 10^9 genome copies (GC) of AAV2/8-RHOK-ZF-R6 and a vector coding for the enhanced green fluorescent protein (EGFP; AAV2/1-CMV-EGFP), to identify the transduced retinal area. The contralateral eye received a mix of AAV2/1-CMV-EGFP and AAV2/8-RHOK-EGFP vectors as control. Western blotting showed expression of the ZF-R6 protein in the retina 12 days post-injection (P18, before photoreceptor degeneration; Fig 3a). Quantitative real-time polymerase chain reaction (PCR) analysis demonstrated that the EGFP-positive portion of the retinae treated with AAV2/8-RHOK-ZF-R6 showed significant, ~26%, reduction in the hRHO transcript levels relative to the endogenous mRho, compared to the contralateral EGFP-treated retinae (Fig 3b; n = 9, from two independent littermates; p < 0.01). In contrast, the expression profiles of three photoreceptor-specific genes were unchanged (Fig 3b; n = 4).

We next evaluated the ability of AAV2/8-RHOK-ZF-R6 to impact on P347S photoreceptor disease progression. As we noted that there was variability in the retinal function assessed by recording the electrophysiological response of the retina to increasing light intensities (ERG) among different littermates,
Figure 3. AAV-mediated photoreceptor ZFP gene transfer reduces the severity of adRP retinal dystrophy.

A. Representative Western blot with an anti-HA antibody of the lysates from P347S retinae (P18) injected with AAV2/8-RHOK-ZF-R6 at P4. Lane 1: HEK293 cells transfected with a plasmid encoding for ZF-R6; lane 2: P347S retinal extracts transduced with AAV2/1-CMV-EGFP and AAV2/8-RHOK-EGFP; lane 3: P347S retinal extracts transduced with AAV2/8-RHOK-ZF-R6 and AAV2/1-CMV-EGFP. Actin expression was used for normalization (anti-actin antibody).

B. Histogram from quantitative RT-PCR analysis on P18 retinae transduced at P4 with AAV2/8-RHOK-ZF-R6 and AAV2/1-CMV-EGFP. Specific down-regulation, by 26%, of human rhodopsin transgene expression was measured in the treated eyes (white bars) compared to the control-injected eyes (black bars).

Photoreceptor specific gene expression levels (murine rhodopsin, peripherin, PDE6β and Cnga) remained unaltered in treated and control retinae (values normalized versus endogenous GAPDH transcript levels). ∗∗p < 0.05, as statistically significance difference between ZF-R6 and EGFP treated eyes of the same animals (t-test).

C. Electrophysiological responses of retina recorded by ERG analysis, on P347S+/− mice injected subretinally with AAV2/8-RHOK-ZF-R6 and AAV2/1-CMV-EGFP or AAV2/1-CMV-EGFP and AAV2/8-RHOK-EGFP at P30. The amplitudes represent retinal responses evoked by increasing light intensities under scotopic (dim light) and photopic (bright light) conditions. B-wave amplitudes before treatment (base-line: P30; open triangles, n = 34 eyes, 17 animals) and after treatment (P90; filled circles, n = 16 eyes, AAV2/8-RHOK-ZF-R6 treated animals; open circles, n = 15 eyes, AAV2/8-RHOK-EGFP treated animals). Sixty days after vector delivery, the electrophysiologic responses of the retina were preserved in ZF-R6–treated eyes (black circles) compared to control contralateral EGFP–injected eyes (open circles). ∗∗p < 0.01; ∗∗∗p < 0.001, as statistically significance differences between EGFP control eyes (P90) and base-line responses (P30).

D. Representative indirect ophthalmoscopy of retinal fundus. Fluorescent EGFP expression indicates the extension of retinal exposure after AAV vector administration (P90 P347S retina co-injected with AAV2/8-RHOK-ZF-R6 and AAV2/8-RHOK-EGFP).

E. Representative immunofluorescence histological analysis of AAV2/8-RHOK-ZF-R6 treated retina showing the EGFP–positive retinal portion transduced (right; dotted square, selected field at higher magnification, 20 ×); co-injection of AAV2/8-RHOK-ZF–R6 and AAV2/8-RHOK-EGFP). This treatment resulted in partial preservation of photoreceptor cell nuclei, as compared to EGFP–negative area (left; dotted square as before).
and despite similar P347S rhodopsin levels, we decided to contextually measure base-line ERG parameters and treat the animals subretinally. We performed subretinal injections of 1 × 10^8 GC of AAV2/8-RHOK-ZF-R6 in three independent 1-month-old P347S/− littermates (P30; for a total of 17 animals, 34 eyes: 17 RHOK-ZF-R6 and 17 RHOK-EGFP eyes), with the paired eye of the same animals receiving the same vector dose of AAV2/8-RHOK-EGFP. At 60 days following this subretinal vector administration (P90), the eyes that received AAV2/8-RHOK-ZF-R6 showed preservation of ERG responses, as compared to the control EGFP-treated eyes of the same animals, in which the degeneration led to a ~40% reduction in the b-wave maximum amplitude (Table 1 of Supporting Information, Fig 3c). In addition, histological analysis of mice at P90–100 demonstrated that the ZF-R6-transduced portion of the retina (typically 30–50% of the retina, Fig 3e), was associated with partial protection of the photoreceptor degeneration in seven eyes out of the nine analysed (Fig 3d and e).

Finally, to evaluate potential toxicity on retinal function of a healthy mouse retina due to ZF-R6 treatment, we injected AAV2/8-RHOK-ZF-R6 subretinally in a cohort of 10 C57BL/6 animals. Two months after vector delivery, no histological abnormalities or reductions in either the a-waves or b-waves were seen for the treated eyes, as compared to untreated parental eyes, thus supporting the safety of this approach (Figure S5 of Supporting Information).

**DISCUSSION**

In the present study, we have demonstrated that photoreceptor delivery of an AAV2/8 vector containing artificial ZF transcriptional repressors (ZF-R6) targeted to the hRHO promoter can repress hRHO transgene over-expression, and this, in turn, was associated with improvement in photoreceptor disease in the P347S mouse model of adRP. Our findings extend the possible uses of artificial ZF-based DBPs as novel therapeutics for the treatment of dominant diseases.

A different technological platform based on RNAi was first used in vivo to silence a dominant allele of ataxin-1 for the treatment of spinocerebellar ataxia type 1 in a mutational-independent manner; this approach was then applied to other mouse models of human disease, including the P347S adRP mouse (O’Reilly et al, 2007; Xia et al, 2004). Thus, based on the data presented here, transcriptional (artificial ZFP technology) or post-transcriptional (RNAi) silencing can be considered as alternatives to one another; however, in the light of the different molecular mechanisms underlying these two silencing strategies (promoter binding and transcription inhibition of ZF vs. enzyme-like degradation of mRNA by RNAi), a side-by-side comparison will be necessary to determine which of these two systems is the safest and the most efficacious. Moreover, as well as the off-target silencing events that will potentially set the hurdles for the development of a safe ZFP-based or RNAi-based system, studies have shown that high levels of siRNAs can cause cellular toxicity through various mechanisms (Boudreau et al, 2009; Grimm et al, 2006). Also, an attractive and therapeutically valuable feature of ZFPs is the possibility to control their expression levels through the use of inducible promoter systems, which have not yet been optimized for the control of RNAi-based repression systems. Indeed, a key safety and efficacy element for the success of silencing strategies relies on the fine-tuning control of the expression levels of the repressor system.

The transcriptional, mutational-independent strategy used in the present study is aimed at overcoming the heterogeneity of mutations in the RHO gene. Indeed, the design of mutation-specific silencing molecules is technically, and economically, particularly challenging, considering that: (i) retinitis pigmentosa is widely thought to be the most genetically heterogeneous disorder in man (Wright et al, 2010); and (ii) over 150 dominant mutations in rhodopsin are known to date (OMIN database 180380). As has been previously suggested (e.g. RNAi-mediated suppression and replacement of human rhodopsin in vivo; (O’Reilly et al, 2007)), we propose here to apply a general strategy that is based on the repression of both the mutated and the wild-type alleles, and the simultaneous replacement of the endogenous gene copies with the exogenous vector-delivered gene [repression–replacement strategy; (O’Reilly et al, 2007)]. In the present study, the ZF-Rs acted as transcriptional hRHO-specific silencers, whereas the endogenous copies of murine Rho alleles functioned as a replacement gene. Thus, one future development should be the use of an amenable mouse model (ideally harbouring both a mutated and a normal hRHO gene on a mRho-knock-out background) to determine whether down-regulation of both alleles together with simultaneous replacement with a hRHO transgene (under the transcriptional control of a promoter insensitive to the ZF-R) will result in phenotypic rescue. Although this needs to be tested, the feasibility of this proposed approach is based on the following considerations: (i) we have demonstrated the ability to down-regulate the levels of transcription somatically (via AAV-vector delivery), which represents the major limiting step in the strategy; (ii) the therapeutic levels of transcriptional silencing result in phenotype amelioration (26%), and thereby the levels of RHO necessary for replacement appear feasible to achieve, considering the proof-of-concept obtained with the gene-replacement approach in different animal models of retinal degeneration [recently also in Rho−/− mice; (Chadderton et al, 2009)], as well as clinical trials (Bainbridge et al, 2008; Hauswirth et al, 2008; Maguire et al, 2008); and (iii) there is the possibility to incorporate two genes into the same vector (both the silencing and the replacement constructs in our case), as this has been demonstrated in AAV vectors by us and others (Allocca et al, 2007; Nguyen et al, 2007). Indeed, a vector that incorporates both the artificial ZF-based repressor and the replacement gene will ensure their simultaneous and balanced silencing and repression action in the same transduced photoreceptors [e.g. with bidirectional promoters allowing the coordinated expression of the two transgenes; (Amendola et al, 2005)]. Moreover, it has been shown that both high and low rhodopsin expression levels can be deleterious to rod function, which suggests that tight regulation of rhodopsin expression levels is required for the successful application of any retinal repression–replacement strategy.
Finally, the versatility of ZFP design theoretically allows to switch the repressor with an activator domain (maintaining the same DBD) to also treat dominant diseases that are due to haploinsufficiency.

The results from the present study demonstrate that the novel and generally applicable mutational-independent strategy based on artificial ZFPs allows transcriptional control of an endogenous causative gene and has therapeutic relevance for the treatment of diseases due to gain-of-function mutations.

MATERIALS AND METHODS

Rational design of zinc-finger-based artificial transcription factors

DBDs targeted to 10 different target sites in the hRHO promoter (Table 1) were obtained using the web-based Zinc Finger Tools (http://www.zincfingertools.org; Mandell & Barbas, 2006). These were composed of two arms, each of which recognized a 9-bp half-target site, and they were fused together through a linker sequence, to bridge the gap sequence. We used two different linkers according to the lengths of the gaps: for gaps longer than 3 bp (target sites from 1 to 4), we used a complex structured linker, as reported by Moore (Moore et al., 2001); CRSSVEASCVTLVALLPATSAPTQVSG, while for longer gaps (target sites from 5 to 10), we used a flexible linker, as described by Rebar (Rebar et al., 2002); QNKKGCGDGKKQKH. We optimized the corresponding DNA sequences to facilitate the subsequent cloning steps, and purchased these as transgenes in the pLS-standard vector (TOP Gene Technologies, Quebec, Canada). After the in vitro selection steps, we purchased an additional DBD with the fingers of DBD number six shuffled (from 1.2.3–4.5.6 to 5.1.6–3.4.2) to be used as a negative control for a total of 11 DBDs. All of the DBDs were flanked by a Cai and an Xbal site at the 5’ and 3’ ends, respectively.

DBD number 1 was purchased as a complete protein fused at the N-terminal with the repressor domain KRAB (Krueppel-associated box, DAKSLTAWSTLRVTKDFVDFRTREEWLDTAQQILYRNVMLENYKFVQ) and the nuclear localization signal (PKKKRKV) from the SV40 large T antigen, and fused at the C-terminal with the HA-tag (YPYDVPDYAS). Also, DBD number 2 was purchased as a complete protein, but fused at the C-terminal with the nuclear localization signal, the activator domain VP64 (tetrameric repeat of herpes simplex VP16 minimal-activation domain, GRADALDDFDLDMLGSDALDDFDLDMLGSDALDDFDLDMLGSDALDDFDLDMLGSDALDDFDLDMLGSDALDDFDLDMLGSDALDDFDLDML), and the HA-tag. In the two proteins, the DBDs can be exchanged through a double digestion with the restriction endonucleases Cai and Xbal.

The 11 different transcription activators (ZF-As) and repressors (ZF-Rs) were inserted in the pAAV2.1-RK-EGFP NotI and HindIII sites, downstream of the RHOK promoter (Table 1), by removing the EGFP coding sequence for transfection experiments. For AAV preparations, ZF-R6 was inserted in the pAAV2.1-RK-EGFP NotI and HindIII sites, downstream of the RHOK promoter, by removing the EGFP coding sequence. For retrovirus preparations used in RSC experiments, ZF-R6 was inserted in the pAAV2.1-RK-EGFP NotI and HindIII sites, downstream of the cytomegalovirus (CMV) promoter, by removing the EGFP coding sequence for transfection experiments. For AAV preparations used in RSC experiments, ZF-R6 was inserted in the pAAV2.1-CMV-EGFP NotI and HindIII sites, downstream of the CMV promoter (Table 1). For retrovirus preparations used in RSC experiments, ZF-R6-shuffled was cloned in the lusn plasmid EcoRi and HindIII sites downstream of the CMV promoter.

Luciferase reporter plasmids were generated from the pGL3-Basic vector (Promega, Madison, WI, USA), by cloning the proximal upstream regions of the human or murine rhodopsin gene (see Materials and Methods Section of Supporting Information).

Screening of the functional ZF-ATFs in human cells by transient transfection

HEK293 cells were plated in six-well plates at a density of 400,000 cells/well. Twenty-four hours later, the cells were co-transfected using FUGENE 6 (Roche, Basel, Switzerland), following the manufacturer instructions. Transfection cocktails included 200 ng reporter plasmid (or pGL3Basic as a negative control), 300 ng ZF-ATF, 300 ng of a plasmid encoding for murine CRX (alone, as a positive control in transactivation assays, or in combination with the ZF-Rs, for repression assays) and 10 ng Renilla luciferase reporter plasmid control (Promega), to normalize for transfection efficiency. The amount of DNA was kept constant by adding pAAV2.1-TBG-EGFP, to a total of 1 µg. Cells were harvested 48 h after transfection in 1× PLB Lysis Buffer (Promega). Firefly and Renilla luciferase activities were measured in a GloMax™ luminometer (Promega), using the Dual Luciferase Reporter Assay System (Promega).

Electromobility shift assay

Electromobility shift assays were performed using Lightshift Chemiluminescent Electromobility Shift Assay kits (Pierce), following the manufacturer instructions. The detailed procedure is available in the Materials and Methods Section of Supporting Information.

AAV vector production and purification

AAV vectors were produced by the TIGEM AAV Vector Core using pAAV2.1-RK-ZF-R6 and pAAV2.1-CMV-EGFP. Recombinant AAV8 viruses were produced by triple transfection of HEK293 cells followed by CsCl2 purification of the vectors, as previously described (Auricchio et al., 2001). For each viral preparation, physical titers (GC/ml) were determined by both PCR quantification using TaqMan (Perkin-Elmer, Life and Analytical Sciences, Inc.) and by dot-blot analysis.

RNA preparation and measurement of rhodopsin transcript levels by real-time PCR

RNAs from tissues and cells were isolated using TRIzol reagent (Invitrogen), according to the manufacturer protocol. cDNA was amplified from 500 ng isolated RNA using SuperScript™ III First-Strand Synthesis (Invitrogen), as indicated in the manufacturer instructions. Transcript levels of human and murine rhodopsin were measured by real-time PCR using the ABI Prism 7000 Sequence Detection System (Applied Biosystems, Foster City, CA, USA) and the following primers: hRho_forward 5’...CCGGCTTGCGGATCACT...3’ and hRho_reverse 5’...TTGTCGCCCTCCGTTGAG...3’, mRho_forward 5’...TTTTGAAACAGCATTCCGGA...3’ and mRho_reverse 5’...TCCGATCTCCACGATTGACT...3’. The PCs with cDNA were carried out in a total volume of 25 µl, using 12.5 µl SYBR Green Master Mix (Applied Biosystems) and 400 nM primers under the following conditions: preheating, 50°C for 2 min, and 95°C for 10 min; cycling, 40 cycles of 95°C for 15 s and 60°C for 1 min. All of the reactions were standardized against murine GAPDH using the following primers: mGAPDH_forward 5’...CTTCACCACGATCGAGGC...3’ and
The paper explained

PROBLEM:
Autosomal dominant forms of Retinitis Pigmentosa (adRP) are incurable progressive inherited retinal degenerations, which inexorably lead to blindness. The major hurdles to design effective therapeutic strategies for these diseases are: the genetic heterogeneity (a number of distinct mutations affect numerous individual gene), and the tools to turn off mutated genes are inefficient and difficult to be delivered specifically and efficiently to diseased cells.

RESULTS:
We generated artificial zinc-finger transcriptional repressors (ZFRs) capable to bind DNA regulatory regions controlling rhodopsin gene (Rho) expression, one of the most frequently genes mutated in adRP. To overcome genetic heterogeneity we designed ZFRs that silence RHO in a mutational-independent fashion. We showed that adeno-associated (AAV) viral vector-mediated delivery of ZFR to photoreceptors of a mouse model of adRP that carries a dominant RHO mutation, result in robust down-regulation of RHO and that this in turn results in improvement of disease progression.

IMPACT:
Our results are the first to demonstrate that ZF technology can be exploited to induce transcriptional gene silencing of an endogenous gene via viral vector-mediated delivery in vivo. This proof-of-concept study holds promise for the development of novel therapeutics for adRP-associated with any rhodopsin mutation and a variety of other inherited disorders.
Author contributions

The overall study was conceived and designed by CM and EMS, with important contributions from DS, CB, VM, AA and GM. CM, DS, EM, CB and EMS: performed the experiments. CM, DS, EM, UD, AA, GM and EMS: analysed the data. AA and VM: contributed reagents or analysis tools. EMS wrote the paper with substantial contribution from CM.

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Supporting information is available at EMBO Molecular Medicine online.

The authors declare that they have no conflict of interest.

For more information

Zinc Finger technology:

The Zinc Finger Tool
http://www.scripps.edu/mr/barbas/zfdesign/zfdesignhome.php
The Zinc Finger Consortium
http://www.zincfingers.org/default2.htm
Retinitis Pigmentosa:

Online Mendelian Inheritance in Man (OMIM)
http://www.ncbi.nlm.nih.gov/omim/268000
http://www.ncbi.nlm.nih.gov/omim/180380
Retinal Information Network database
http://www.sph.uth.tmc.edu/Retnet/
Retinal International
http://www.retina-international.org/
European Vision Research
http://www.vision-research.eu/
Foundation Fighting Blindness
http://www.blindness.org/
The Vision of Children Foundation
http://www.visionofchildren.org/
Author’s website:
http://www.tigem.it/researchers/enrico-maria-surace

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