Control of Transcription Factor Activity and Osteoblast Differentiation in Mammalian Cells Using an Evolved Small-Molecule-Dependent Intein

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Abstract: Inteins are naturally occurring protein elements that catalyze their own excision from within a larger protein together with the ligation of the flanking “extein” sequences. Previously we reported the directed evolution of an intein-based molecular switch in which intein splicing in yeast cells was made dependent on the cell-permeable small molecule 4-hydroxytamoxifen (4-HT). Here we show that these evolved inteins are effective means of rendering protein function and biological signaling pathway activation dependent on 4-HT in mammalian cells. We have characterized the generality, speed, and dose dependence of ligand-induced protein splicing in murine NIH3T3 cells and in human HEK293 cells. Evolved inteins were used to control in mammalian cells the function of Gli1 and a truncated form of Gli3, two transcriptional mediators of the Hedgehog signaling pathway. Finally, we show that a complex biological process such as osteoblast differentiation can be made dependent on 4-HT using the evolved intein system. Our findings suggest that evolved small-molecule-dependent inteins may serve as a general means of achieving gene-specific, dose-dependent, post-translational, and small-molecule-induced control over protein activity in mammalian systems.

Introduction

The study of mammalian signaling pathways and regulatory networks is a persistent challenge due to their complexity and to inherent limitations of methods used to study them. Although methods that enable inducible and highly specific control over a protein of interest without perturbing the structure of the protein or its regulatory genetic elements would be ideal, many approaches to modulating protein function in bacteria and yeast which meet some of these goals cannot readily be applied to mammalian systems.

Strategies for small-molecule-based regulation of protein activity in mammalian cells that can be applied to arbitrary proteins of interest include transcriptional control by regulatable promoters and genetic modification by small-molecule-responsive Cre recombinase fusion proteins. While both methods have been used successfully to probe biological systems, each has significant limitations. Promoter-based systems place expression of the protein of interest under the control of a non-native promoter. While the structure of the resulting protein is not affected by this change, the ability of endogenous factors to interact with the promoter can be compromised, potentially disrupting regulatory networks that normally control target protein expression. In addition, promoter-based systems typically require the expression of heterologous transcriptional activators or repressors, some of which can interact with the endogenous cellular machinery, resulting in toxicity or immuno-negativity.

Recombine-based systems require concurrent expression of a foreign small-molecule-dependent recombinase. In addition, because protein expression induced by genetic recombination requires that each individual cell undergo a relatively slow recombination event, cell lines expressing a variety of small-molecule-dependent Cre recombinases typically achieve target protein activation in only ≤50% of cells after 48 h following treatment with the small molecule. More efficient transgenic cell lines have been developed that undergo higher levels of recombination, but their use requires longer periods of exposure or high doses of small molecule. An efficient small-molecule-dependent method of controlling the activity of any protein of

interest in mammalian cells without requiring changes to the target protein’s promoter or the concurrent expression of heterologous proteins may prove useful for probing protein function. In addition, control of proteins at the post-translational level, in contrast with the above approaches, may enhance the speed with which changes in activity can take place.

We recently described the directed evolution of an intein-based molecular switch that renders intein splicing and therefore target protein activity dependent on the presence of the cell-permeable small molecule 4-hydroxytamoxifen (4-HT) in yeast cells. Because the evolved intein-based switch is a 4-HT-dependent, self-excision protein element contained within a target protein, its use does not require promoter modification or any additional control elements.

Other groups have also reported the use of inteins to modulate protein activity in eukaryotic cells. Umezawa and co-workers have shown that splicing of the split Synecocystis sp. DnaE intein fused to the N- and C-terminal halves of luciferase generates active luciferase in mammalian cells, albeit in an unregulated manner. Perrimon and co-workers developed a temperature-sensitive Saccharomyces cerevisiae VMA intein and used it to control the activity of the yeast Gal80 transcription factor in both yeast and Drosophila melanogaster. Muir and co-workers engineered a split VMA intein in which each half of the intein is fused to FKBP or FRB domains. Rapamycin induces dimerization of the FKBP and FRB domains, which unites the intein halves and enables splicing. Rapamycin-dependent splicing in this system has been shown to mediate the joining of two epitope tags in mammalian cells as well as the removal of an inhibitory peptide from a protein kinase A–split intein fusion using purified proteins in solution. Target proteins for which inhibitory peptides are not known or that are not stable when separated and expressed as two intein-fused halves, however, may prove difficult to manipulate using a split intein system. Regulated inteins have not yet been reported to control native protein function in mammalian cells.

Here we show that the evolved small-molecule-dependent intein confers 4-HT dependence on the activities of proteins in murine and human cells in a dose-dependent and fairly rapid manner. We demonstrate that native mammalian proteins controlled by the evolved intein are able to perform their endogenous biological functions in mammalian cells. Further, we show that the evolved intein can render a complex process such as osteoblast differentiation dependent on 4-HT. Our results validate the use of evolved small-molecule-dependent inteins as tools to introduce post-translational control elements into signaling pathways in mammalian cells.

Experimental Section

General. 4-Hydroxytamoxifen, Leptomycin B, N-terminal Sonic Hedgehog peptide (nShh), anti-3X Flag antibody, and anti-β-actin antibody were purchased from Sigma-Aldrich. Alexa Fluor 488 anti-mouse antibody was purchased from Invitrogen. Alexa Fluor 680 anti-rabbit and 800 anti-mouse antibodies were provided by Li-cor Biosciences. Restriction enzymes were purchased from New England Biolabs. Cell culture reagents were purchased from WVR and Invitrogen. Western blots were visualized and quantitated using an Odyssey imager (Li-cor Biosciences). Fluorescence activated cell sorting (FACS) analysis was performed using a MoFlo cell sorter (DakoCytomation). Sodium dodecyl sulfate—polyacrylamide gel electrophoresis and Western blotting were performed using standard protocols.

Plasmid Construction. To create a stably transfected cell line expressing a green fluorescent protein (GFP)–intein(3-2) fusion, the encoding gene was cloned into pcDNA3.1/Hygro (Invitrogen). The GFP-intein(3-2) gene was amplified by polymerase chain reaction (PCR) with Vent polymerase (New England Biolabs) from p414Gal1-GFP-intein(3-2) using oligonucleotides 5′-GCCGCCCTTTAAGAT- GGCAAGCCAAGGAGAAC-3′ and 5′-GCCGCCGCCGCGCTTATTTTAGCATCCTAC-3′. The resulting fragment was treated with AffIII and NorI (restriction sites underlined) and ligated into AffIII- and NorI-digested pcDNA3.1/Hygro.

C-terminally Flag-tagged Gli1-intein and Gli3T-intein expression vectors were constructed from p3XFlag-CMV-14-Gli1 and p3X-Flag-CMV-14-Gli3T. An Xhol restriction site was introduced into the Gli1 gene by silent mutation of the Ser 279 codon. An Xhol restriction site was removed from the Gli3T gene by silent mutation of the Leu 728 codon, while EcoRI and Xhol restriction sites were added by silent mutation of the Glu 512 and Ser 521 codons, respectively. All mutations was performed using standard protocols for QuiChange site mutagenesis (Stratagene). Gene fragments encoding intein(3-2) and intein(2-4) were amplified by PCR from p414Gal1-GFP-intein(3-2) and p414Gal1-GFP-intein(2-4). Fragments were amplified using oligonucleotides 5′-GCCGCCGATCTGCCTGGAGGGTACCCG-3′ and 5′-GCCGCCCTTCAGCAACTCCCTCAATGGCAGTGTGTGACGACAAC-3′ (for Gli1) or 5′-GCCGCCCTTCAGGAAACTCAAGACCGCCGGCACTTIGCACGACAAC-3′ (for Gli3T). Fragments were digested with EcoRI and Xhol and ligated into the appropriate EcoRI- and Xhol-treated vectors.

Genes encoding nonsplicing Cys–Ala mutants of Gli1-intein(3-2) and Gli3T-intein(3-2) were constructed by mutating the Cys-encoding codon at the C-terminus of the intein (TGC) to one encoding Ala (GCC). The genes encoding Flag-tagged Gli1, Gli1-intein(3-2) and Gli1-intein(2-4), were also cloned into pCAGGS for expression in 10T1/2 cells.

Cell Culture. HEK293 cells were cultured in Dulbecco’s modified Eagle medium (DMEM):F12 medium with 10% fetal bovine serum, while NIH3T3 and 10T1/2 cells were cultured in DMEM with 10% calf serum according to standard protocols. Transient transfections were performed using Lipofectamine2000 (Invitrogen) following the manufacturer’s protocol. To generate a stably transfected cell line expressing GFP-intein, HEK293 cells were plated in a 10 cm dish coated with Matrigel (BD Biosciences). Cells were transfected with pcDNA3.1-GFP-intein(3-2) using the ProFection calcium phosphate transfection system (Promega) following the manufacturer’s protocol. Seventy-two hours after transfection, cells were replated onto three 10 cm dishes in media containing 150 μg/ml G418 and 1 mg/ml Hygromycin B (EMD Biosciences). Resistant clones were replated individually and analyzed by Western blot using an anti-GFP antibody.

FACS Analysis. Stably transfected HEK293 cells expressing GFP-intein(3-2) were grown in six-well plates with the specified concentrations of 4-HT for 24 h. Cells were then trypanosized and resuspended in 500 μl of phosphate buffered saline with 1% fetal bovine serum and 75 U/ml DNase (New England Biolabs). At least 106 cells were analyzed for each sample.
Luciferase Assays. NIH3T3 cells were plated in a 24-well plate containing 5 × 10^4 cells per well. Eighteen hours after plating, cells were transiently transfected with a total of 800 ng of DNA, including 300 ng of ptcΔA136-GL3 luciferase reporter and 100 ng of pSV-β-galactosidase transfection control (Promega). The control transfection contained 100 ng of p3XFlag vector with no encoding gene. Transfections to express Gli1, Gli1-intein(3-2), Gli1-intein(2-4), and nonsplicing mutant Gli1-intein(3-2) contained 100 ng of the corresponding p3XFlag expression vector. Transfections to express Gli3T, Gli3T-intein(3-2), Gli3T-intein(2-4), and nonsplicing mutant Gli3T-intein(3-2) contained 10 ng of the corresponding p3XFlag expression vector, as well as 100 ng of p3XFlag-Gli1. PSK Bluescript (Stratagene) was added to each transfection mixture to bring the total DNA content to 800 ng. Forty-eight hours after transfection, the medium was changed to DMEM with 0.5% calf serum, containing 4-HT where appropriate. After an additional 48 h, cells were collected and assayed as previously described. Luciferase activity values were normalized to β-galactosidase activity values to correct for variations in numbers of transfected cells. Note that, although NIH3T3 cells contain Gli1- and Gli3-encoding genes, the level of CMV promoter-driven overexpression of transcribed Gli1 and Gli3T in this standard assay is much greater than endogenous Gli1 or Gli3 levels.

Immunohistochemistry. HEK293 cells were plated on Matrigel-coated glass coverslips in a six-well plate and transiently transfected with 4 μg of the specified p3XFlag expression vector. After growth in the absence or presence of 1 μM 4-HT for 48 h, cells were fixed with methanol and treated with anti-3XFlag antibody, Alexa 488 anti-mouse antibody, and 4',6-diamidino-2-phenylindole (DAPI) following standard protocols. For cells treated with Leptomycin B (LMB), 5 nM LMB was added 2 h prior to fixation.

Alkaline Phosphatase Assays. 10T1/2 cells were plated in 96-well plates containing 10^4 cells per well. Twenty-two hours after plating, cells were transiently transfected with 200 ng of the appropriate pCAGGS expression vector or an empty pCAGGS vector as a control. Eight hours after transfection, fresh medium was added that contained 1 μM 4-HT or 50 nM 6λH where specified. Five days after transfection, cells were lysed by freeze--thaw and assayed using pNPF alkaline phosphate liquid substrate (Sigma). Reactions were incubated for 1 h at 37°C and measured by spectrophotometer.

Results

Evolved Intein Mediates Small-Molecule-Dependent Splicing of GFP in Mammalian Cells. Given the significant differences between yeast and mammalian cells, we first evaluated the ligand dependence and splicing ability of the previously yeast-evolved inteins10 in mammalian cells. We used the final 0.5×10^5 cells per well. To evaluate the speed of intein-mediated protein activation, we observed the time dependence of 4-HT responsiveness using GFP-intein(3-2). When cells stably expressing GFP-intein are treated with 4-HT, spliced protein can be detected by Western blot within 2 h and reaches its maximum concentration within 24 h (Figure 2C). In the absence of 4-HT, less than 0.5% of protein detected is spliced, while at 2 h (the earliest time point measured), 5% of protein is spliced. Western blot quantitation of three separate trials (representative shown in Figure 2C) indicates that, after 24 h, the amount of spliced protein remains relatively constant at 30–40% of total protein detected. These findings indicate that intein(3-2) splicing in the presence of 4-HT takes place on a time scale of hours.

Evolved Inteins Mediate Splicing of Mammalian Transcription Factors. To determine whether evolved inteins could control the activity of native proteins in living mammalian cells, we inserted evolved inteins into the murine transcription factor

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Gli1 and into a C-terminally truncated form of the transcription factor Gli3 (Gli3T). Gli1 and Gli3 are key mammalian transcriptional mediators of Hedgehog signaling and play key roles in developmental processes, including spinal cord patterning and limb development. Gli1 is a transcriptional activator, while Gli3T is an engineered version of the repressor form of the Gli3 transcription factor. Separate constructs were generated for both Gli1 and Gli3T containing either intein-(3-2) or intein-(2-4) preceding a zinc-coordinating cysteine in the highly conserved DNA binding domain present in both transcription factors (Cys 273 in Gli1, Cys 515 in Gli3T). In these constructs, protein splicing therefore generates Gli1 or Gli3T with primary structures that are identical to those of the native proteins.

When HEK293 cells were transiently transfected with each of the Gli1-intein or Gli3T-intein fusions, 4-HT-dependent splicing was observed via Western blot (Figure 3A,B). The amount of spliced protein generated in the presence of 4-HT was measured by FACS. The red spot represents the median fluorescence of untransformed cells. Data points and error bars reflect the mean and standard deviation, respectively, of three independent trials. (B) The fluorescence histograms of the cell populations described in (A) show that increasing concentrations of 4-HT induce increasing amounts of protein splicing throughout the cell populations. Individual histograms are normalized to a total cell count of $2 \times 10^5$. (C) HEK293 cells stably expressing GFP-intein-(3-2) were treated with 1 $\mu$M 4-HT for the indicated time periods and analyzed by Western blot using an anti-GFP antibody.

Figure 2. 4-HT-dependent splicing of evolved intein is dose-dependent and takes place within hours. (A) HEK293 cells stably expressing GFP-intein-(3-2) were grown for 24 h in 0–2.0 $\mu$M 4-HT. Fluorescence was measured by FACS. The red spot represents the median fluorescence of untransformed cells. Data points and error bars reflect the mean and standard deviation, respectively, of three independent trials. (B) The fluorescence histograms of the cell populations described in (A) show that increasing concentrations of 4-HT induce increasing amounts of protein splicing throughout the cell populations. Individual histograms are normalized to a total cell count of $2 \times 10^5$. (C) HEK293 cells stably expressing GFP-intein-(3-2) were treated with 1 $\mu$M 4-HT for the indicated time periods and analyzed by Western blot using an anti-GFP antibody.

Figure 3. 4-HT-dependent splicing of evolved inteins generates active Gli1 and Gli3T in mammalian cells. (AB) HEK293 cells were transiently transfected with Flag-tagged Gli1- and Gli3T-intein expression vectors, grown for 24 h in the absence or presence of 1 $\mu$M 4-HT, and analyzed by Western blot using an anti-Flag antibody. (C) NIH3T3 cells were transiently transfected with the indicated expression vectors, as well as a luciferase reporter vector and a $\beta$-galactosidase transfection control vector. The control expression vector contains no gene, Gli1-(2-4) and Gli1-(3-2) refer to the Gli1-intein constructs, and Gli1-(3-2)mut refers to the Cys$\rightarrow$Ala mutant of Gli1-intein(3-2). The luciferase activity resulting from the control vector in the absence of 4-HT is defined as 1. (D) NIH3T3 cells were transiently transfected with the indicated expression vectors, the luciferase reporter vector, and a $\beta$-galactosidase control vector. Gli3T-(2-4) and Gli3T-(3-2) refer to the Gli3T-intein constructs, and Gli3T-(3-2)mut refers to the Cys$\rightarrow$Ala mutant of Gli3T-intein(3-2). Luciferase activity induced by Gli1 is defined as 100. For all luciferase assays, luciferase activity was normalized to $\beta$-galactosidase activity. Bar heights and error bars reflect the mean and standard deviation, respectively, of two or three independent trials.
was approximately 50–60% of the total Gli1 or Gli3T protein detected, but in the absence of 4-HT was only 0–10% of total Gli1 or Gli3T protein produced. These results demonstrate that insertion of evolved intein(3-2) or intein(2-4) into Gli1 and Gli3T confers 4-HT-dependent control over each protein’s primary structure.

4-HT-Dependent Splicing Generates Active Gli1 and Gli3T in Mammalian Cells. To analyze the activity of the Gli1- and Gli3T-intein fusions, we used an established luciferase-based reporter assay.17 Murine NIH3T3 cells were co-transfected with either Gli1-intein or Gli3T-intein expression vectors and a reporter construct containing the consensus Gli binding sequence from the promoter of patched, an endogenous target of Gli regulation,26 upstream of the firefly luciferase gene. Without 4-HT treatment, cells transfected with Gli1-intein(2-4) exhibited no significant increase in luciferase activity over cells transfected with a control expression vector that contains no gene (Figure 3C). Cells transfected with Gli1-intein(3-2) exhibited a 2-fold increase in luciferase production over the control in the absence of 4-HT. In the presence of 1 μM 4-HT, Gli1-intein(2-4) induced approximately 25% as much luciferase production as wild-type Gli1, while Gli1-intein(3-2) induced approximately 35% as much. As controls, no luciferase production was observed in response to 4-HT in cells not expressing Gli1-intein protein (Figure 3C), and 4-HT treatment of cells transfected with wild-type Gli1 had no effect on luciferase production (data not shown). These results demonstrate that expression of Gli1-intein(2-4) or Gli1-intein(3-2) results in 4-HT-dependent transcriptional activation of the luciferase reporter.

To measure the activity of the intein-controlled Gli3T repressor, Gli3T-intein expression vectors were co-transfected with the reporter and a 10-fold excess of wild-type Gli1 expression vector (Figure 3D). Repression by native Gli3T decreases Gli1-induced luciferase production by 80% (down to 20% of its unrepressed level). In the presence of 4-HT, Gli3T-intein(2-4) and Gli3T-intein(3-2) decrease luciferase production by 50% and 70%, respectively, of its unrepressed level. Therefore, expression of Gli3T-intein(2-4) or Gli3T-intein(3-2) leads to 4-HT-dependent repression of luciferase production. These results collectively show that the evolved inteins, upon small-molecule-triggered splicing, generate Gli1 and Gli3T proteins that retain their native activities.

To determine whether protein splicing is necessary for the 4-HT-dependent activity of the Gli1- and Gli3T-intein fusions, we made mutants of Gli1- and Gli3T-intein(3-2) in which the nucleophilic cysteine at the C-terminus of the intein is mutated to alanine. An analogous mutation in the Pyrococcus GB-D Pol intein has been shown to completely eliminate splicing.27 The nonsplicing mutant Gli1-intein(3-2)mut induces no luciferase production in response to 4-HT (Figure 3C), and the nonsplicing Gli3T-intein(3-2)mut does not repress luciferase production significantly either in the absence or in the presence of 4-HT (Figure 3D). These results indicate that the small-molecule-dependent transcription factor activity of the Gli1- and Gli3T-inteins is contingent upon protein splicing. Furthermore, any background activity of the Gli1 and Gli3T-inteins in the absence of 4-HT is most likely due to low levels of protein splicing that occur in the absence of 4-HT, and not to any residual activity of unspliced protein.

The high sensitivity and quantitative nature of the luciferase assay enables a more detailed comparison of the activities of intein(3-2) and intein(2-4). Consistent with their properties in yeast,10 intein(3-2) generates greater quantities of spliced protein in the presence of 4-HT than intein(2-4), although the activity of either intein-containing transcription factor in the presence of 4-HT is lower than the activity of the wild-type transcription factor (Figure 3C,D). Intein(2-4) leads to a smaller amount of background splicing in the absence of 4-HT than does intein(3-2) in mammalian cells. This background splicing causes a 2-fold increase in luciferase production over the control in cells expressing Gli1-intein(3-2), but no observable increase in cells expressing Gli1-intein(2-4). The background splicing of Gli3T-intein(2-4) and Gli3T-intein(3-2) results in repression of Gli1-induced luciferase production by 20% and 25%, respectively. This background repression is surprising, given that no spliced protein is observed in the absence of 4-HT by Western blot. Gli3T may be a sufficiently potent transcriptional repressor that very low concentrations of active Gli3T may produce significant measurable effects in the luciferase reporter assay.

Spliced Gli1-Intein and Gli3T-Intein Localize Identically to Their Native Counterparts. Next we examined the subcellular localization of Gli1 and Gli3T containing the evolved intein. HEK293 cells were transiently transfected with Flag-tagged Gli1, Gli3T, Gli1-intein(3-2), or Gli3T-intein(3-2) and then fixed and stained with anti-Flag antibody (Figure 4). Normally the repressor form of Gli3 is localized to the nucleus,28 while Gli1 undergoes nuclear export as part of its regulatory mechanism and can therefore appear in both the nucleus and the cytoplasm.29 Consistent with these observations, Gli1 expressed in transfected cells appears broadly distributed throughout the nucleus and cytoplasm (Figure 4A), but addition of the small molecule LMB, a potent inhibitor of nuclear export, leads to accumulation of Gli1 in the nucleus (Figure 4B). Gli3T expressed in transfected cells is nuclearly localized (Figure 4C).

In cells transfected with Gli1-intein(3-2) or Gli3T-intein(3-2), protein produced in the absence of 4-HT is cytoplasmically sequestered, in contrast with the localization of native Gli1 and Gli3T (Figure 4A,C). We attribute cytoplasmatic sequestration in the absence of 4-HT to the interaction of Hsp90 with the mutated estrogen receptor ligand-binding domain that exists within the evolved intein,30 even though in yeast cells we demonstrated that this interaction is not solely responsible for the 4-HT dependence of protein splicing.10 When cells transfected with Gli1-intein(3-2) were grown in the presence of 4-HT, protein appeared broadly distributed (Figure 4A). However, when Gli1-intein(3-2) transfected cells were grown in the presence of 4-HT and treated with LMB, nuclear localization was observed (Figure 4B). When cells transfected with Gli3T-intein(3-2) were grown in the presence of 4-HT, the resulting protein localized to the nucleus (Figure 4C). The ability of Gli1- and Gli3T-intein(3-2) to localize correctly within the cell in the

presence of 4-HT further suggests that the Gli1 and Gli3T generated upon 4-HT-dependent splicing retain their native properties.

To determine whether protein splicing is necessary for the correct localization of proteins containing the evolved intein, the localization of the nonsplicing Cys→Ala Gli3T-intein(3-2)mut was examined (Figure 4D). As with Gli3T-intein(3-2), the nonsplicing mutant protein is cytoplasmically localized in the absence of 4-HT and nuclearly localized in the presence of 4-HT. Therefore, nuclear localization of Gli3T-intein(3-2) is not contingent upon protein splicing but is instead likely dependent on the abrogation of interactions with Hsp90 upon treatment with 4-HT.

Evolved Inteins Render Gli-Mediated Osteoblast Differentiation Dependent on 4-HT. The above results demonstrate that 4-HT-dependent splicing can be used to generate protein that is both active and correctly localized, but they do not reveal if the amounts of active protein generated are sufficient to fulfill the proteins’ endogenous roles in their native biological contexts. We therefore next evaluated the function of an intein-controlled mammalian protein in its endogenous context. To achieve this goal, we used the well-characterized mouse embryonic 10T1/2 cell line as a test system for the biological function of Gli1-intein(3-2) and Gli1-intein(2-4).

Activation of the Hedgehog signaling pathway causes 10T1/2 cells to differentiate into osteoblasts, which can be quantitatively measured by the production of alkaline phosphatase.\(^{31}\) In nature, the Hedgehog pathway is activated by the presence of Sonic Hedgehog (Shh) protein, which binds to an extracellular receptor to initiate a signaling cascade that results in the modulation of Gli transcription factor activity. As Gli1 is a downstream component of the Hedgehog pathway, transfection of 10T1/2 cells with a construct expressing Gli1 induces osteoblast differentiation in the absence of Shh. When cells are transfected with Gli1-intein(3-2) or Gli1-intein(2-4), however, osteoblast differentiation becomes dependent on 4-HT in the absence of Shh (Figure 5). Importantly, in the presence of 4-HT, both Gli1-intein(2-4) and Gli1-intein(3-2) promote differentiation to the same extent as wild-type Gli1, even though the intein fusions generate only 25% and 35% as much activity, respectively, as measured by the luciferase reporter assay. These results demonstrate that the quantities and activity of Gli1 generated by 4-HT-mediated splicing of Gli1-intein are sufficient to fulfill its endogenous role in osteoblast differentiation to an extent comparable to that of wild-type Gli1, and in a manner that is no longer dependent on Shh.

**Discussion**

Comparison of Evolved Intein Properties in Yeast and Mammalian Cells. In this work we have shown that two evolved intein variants are capable of rendering mammalian protein activity dependent on a cell-permeable small molecule. In general, the properties of the inteins in mammalian cells parallel their properties in yeast. Protein splicing is dependent on 4-HT in both systems, and intein(3-2) generates more spliced product in the presence of 4-HT than intein(2-4). A graded dose-
dependent response to 4-HT is also observed in both yeast and mammalian cells. These similarities of behavior further underscore the generality of the evolved inteins, even though the environment of the mammalian cell is significantly different than the yeast cell environment used to evolve the inteins.

One difference observed between the evolved intein’s behavior in yeast and its behavior in mammalian cells is that spliced GFP appears to accumulate slower in mammalian cells expressing GFP-intein(3-2) than in yeast expressing the same protein. While the onset of splicing is rapid in mammalian cells, only 5% of detected protein is spliced after 2 h. In yeast, this percentage is significantly higher. This discrepancy may be due to differences in the stability of the unspliced protein, its folding ability, or its rate of degradation in the two cellular contexts. Although the accumulation of spliced protein over a 24-hour period is consistent with the product accumulation observed using an engineered rapamycin-dependent trans intein system in mammalian cells, further optimization of the evolved intein’s activity in mammalian cells may be possible through additional rounds of directed evolution.

**Basis of Ligand-Dependent Activation of Gli1-Intein and Gli3T-Intein.** The cytoplasmic sequestration of Gli1-intein(3-2) and Gli3T-intein(3-2) in the absence of 4-HT complicates the analysis of the mechanism of their evolved ligand-dependent activities. In contrast to GFP, Gli1 and Gli3T are active only in the nucleus. Therefore, it could be argued that cytoplasmic sequestration rather than the presence of the intein is responsible for the lack of Gli1 and Gli3T activity in the absence of 4-HT. Indeed, the Hsp90-mediated cytoplasmic sequestration of protein fusions containing hormone receptor ligand-binding domains has served as a basis for previously engineered small-molecule-dependent proteins, including the Cre recombinase variants described above. However, the above results indicate that a nonsplicing evolved intein mutant leads to cytoplasmic sequestration in the absence of 4-HT and localization identical to that of native protein in the presence of 4-HT (Figure 4D).

Importantly, nonsplicing Gli1-intein or Gli3T-intein mutants demonstrate no activity by luciferase assay either in the presence or in the absence of 4-HT (Figure 3C,D). These results indicate that, while 4-HT releases proteins containing the evolved intein from cytoplasmic sequestration, splicing is required for protein activity. Therefore, 4-HT-dependent splicing of the evolved intein rather than a 4-HT-mediated change in localization is responsible for 4-HT-dependent activity of Gli1- and Gli3T-inteins in mammalian cells. This conclusion is consistent with our earlier studies in yeast cells in which we proposed that Hsp90 complexation may stabilize the unspliced protein but is not responsible for the ligand dependence of splicing.

The possibility exists that intein splicing observed in the absence of 4-HT may be triggered by endogenous estrogen. However, growth in estrogen-free media did not affect the levels of background splicing of Gli1- and Gli3-intein(3-2) observed by Western blot (data not shown). The responsiveness of the evolved inteins to estrogen and estrogen analogues other than 4-HT will be the subject of future experiments, and selectivity for 4-HT over estrogen could be optimized through further directed evolution.

**Application of Evolved Ligand-Dependent Inteins to the Study of Mammalian Signaling Pathways.** The demonstration that an evolved intein inserted into Gli1 renders osteoblast differentiation dependent on 4-HT suggests the possibility of target protein-specific, post-translational, small-molecule-dependent, dose-dependent, and temporally controlled protein activation in a transgenic mammalian system. Such a system would enable control of target protein function to be decoupled from natural upstream signaling partners. For example, a transgenic cell expressing Gli1-intein(3-2) in place of Gli1 will not activate Gli1 in response to Shh but will do so in response to 4-HT. Achieving small-molecule-dependent control over the function of a specific protein in order to study cellular processes is a central goal of chemical genetics. Traditional chemical genetic approaches require the discovery of a small-molecule activator or repressor of each protein of interest. The use of the evolved intein confers 4-HT dependence on proteins of interest in mammalian cells without requiring small-molecule synthesis and discovery, albeit with the additional significant requirement of genetic intervention.

The evolved intein is a particularly attractive tool for the study of signaling pathways because it requires minimal disruption of regulatory networks. No changes to the promoter or other regulatory regions of the gene of interest are required. Many signaling pathways involve feedback inhibition loops in which a downstream product of the pathway represses transcription of an upstream protein to attenuate the signal. An inducible promoter can replace the endogenous promoter of an upstream signaling protein in order to activate the signaling pathway on demand, but natural feedback transcriptional repression in this case cannot occur, precluding deactivation of the pathway by its normal regulatory mechanism. In contrast, the evolved intein exerts control over protein activity at the post-translational level. As a result, feedback regulation of protein-intein expression can proceed normally because no changes to regulatory promoter elements are required.

Because our results indicate that the evolved intein can exhibit a linear dose-dependent response to varying concentrations of
4-HT in mammalian cells, it is well suited for studying the response of a pathway to different levels of target protein activity. For example, transcription factors involved in stem cell differentiation have been shown to induce different cell fates based on modest changes to their cellular concentrations. Smith and co-workers showed that a 50% increase in the expression of Oct-4, a transcription factor involved in regulating pluripotency in embryonic stem cells, induces differentiation to endoderm or mesoderm, while a 50% decrease in expression causes de-differentiation to trophoderm.34 Likewise, Yamamoto and co-workers reported that complete inactivation of GATA-1, a transcription factor required for erythroid differentiation, both inhibits differentiation and induces apoptosis in erythroid progenitor cells, while expression of GATA-1 at 5% of wild-type levels inhibits differentiation without inducing apoptosis.35 While these studies required the generation of multiple cell lines to achieve different levels of protein activity, the use of the evolved intein may enable regulation of the level of target protein activity in a single cell line simply by altering the concentration of 4-HT in the medium.

Conclusion

In conclusion, we have shown that the evolved 4-HT-dependent intein can function as a tool for the relatively rapid, post-translational manipulation of protein function in mammalian cells. We inserted evolved inteins into three target proteins to create 4-HT-dependent GFP, Gli1, and Gli3T, all of which exhibited native-like function in the presence of 4-HT. Expression of Gli1-intein fusions in embryonic mouse cells enables 4-HT to induce a cellular response that mimics the effect of Hedgehog pathway activation. These results collectively suggest that evolved small-molecule-dependent inteins may provide a general method for achieving small-molecule-dependent control over protein activity in mammalian systems. The ability to control protein activity in a post-translational, rapid, dose-dependent, and gene-specific manner without requiring non-native promoters or heterologous control elements could prove valuable in the study of mammalian proteins and signaling pathways.

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