Next-generation plasmids for transgenesis in zebrafish and beyond

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Summary Statement

We describe the generation, validation, and application of several new plasmid vectors suitable for transgenesis in zebrafish and other species.

Abstract

Transgenesis is an essential technique for any genetic model. Tol2-based transgenesis paired with Gateway-compatible vector collections has transformed zebrafish transgenesis with an accessible, modular system. Here, we established several next-generation transgenesis tools for zebrafish and other species to expand and enhance transgenic applications. To facilitate gene-regulatory element testing, we generated

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Gateway middle entry vectors harboring the small mouse *beta-globin* minimal promoter coupled to several fluorophores, CreERT2, and Gal4. To extend the color spectrum for transgenic applications, we established middle entry vectors encoding the bright, blue-fluorescent protein mCerulean and mApple as an alternative red fluorophore. We present a series of p2A peptide-based 3' vectors with different fluorophores and subcellular localizations to co-label cells expressing proteins of interest. Lastly, we established Tol2 destination vectors carrying the zebrafish *exorh* promoter driving different fluorophores as a pineal gland-specific transgenesis marker active prior to hatching and through adulthood. *exorh*-based reporters and transgenesis markers also drive specific pineal gland expression in the eye-less cavefish (*Astyanax*). Together, our vectors provide versatile reagents for transgenesis applications in zebrafish, cavefish, and other models.

Introduction

Transgenesis is a central technique across model systems for investigating biological processes *in vivo*. Integration of a transgene into the genome of a model organism enables experiments including live imaging, lineage tracing, generating conditional mutants, and disease modelling. Transgenic reporters permit labelling of cells, lineages, and structures *in vivo* using fluorescent proteins, making transgenesis an imperative tool in the study of developmental processes. Transgenesis can be paired with spatio-temporally controlled or ectopic gene expression to investigate the role of a particular gene within a molecular pathway. The zebrafish is an ideal system for transgenic applications due to its rapid development, optically transparent embryos, fecundity, and amenability to genetic manipulation. Numerous additions to the toolbox for creating transgenic lines in zebrafish and improving transferability of expression constructs between species have critically enabled the study of evolutionarily conserved genes and their role in developmental processes and disease progression (Amsterdam et al., 1995; Carney and Mosimann, 2018; Fisher et al., 2006a; Kawakami, 2004; Kawakami, 2005; Kikuta and Kawakami, 2009; Long et al., 1997; Meng et al., 1997; White et al., 2013).

In particular, Tol2-based transgenesis and accessible vectors for its application have transformed the zebrafish community with a simple, modular system for the efficient generation of transgene constructs (Kawakami, 2004; Kawakami, 2005). First isolated in medaka (Koga et al., 1996), Tol2-based transgenesis requires the delivery of transposase mRNA and a transgenesis vector containing a desired transgene cargo flanked by Tol2 transposon repeats, resulting in efficient random integration into the genome (Kawakami, 2005; Kawakami et al., 1998). This workflow is particularly suitable for use in zebrafish, which are injected with Tol2 transposase-encoding mRNA and a transgenesis vector at

the one-cell stage (Kawakami, 2005; Kawakami, 2007; Kikuta and Kawakami, 2009; Urasaki et al., 2008). Tol2 transposon activity for transgenesis is not restricted to zebrafish, as injection or electroporation-based delivery methods have been successfully used in several fish species and in other vertebrate models including *Amphioxus*, *Xenopus*, and chicken embryos (Hamlet et al., 2006; Kozmikova and Kozmik, 2015; Sato et al., 2007).

Despite its versatility, taking full advantage of Tol2 transgenesis hinges upon the broad availability, applicability, and functionality of its required components. The first iteration of the Tol2kit (Kwan et al., 2007) and other powerful plasmid collections (Don et al., 2017; Villefranc et al., 2007) provided several entry and destination vectors for Multisite Gateway cloning (Cheo et al., 2004; Hartley et al., 2000) or restriction enzyme-based assembly of [regulatory element(s)]–[effector coding sequence]–[3' trailer] constructs into a Tol2 transposon-flanked backbone. However, as transgenesis applications have become more widely applied and complex, an expanded set of transgenesis components has become essential.

First, transgenes based on isolated regulatory elements need a basic complement of validated parts to constitute a gene expression unit. Most prominent is the need for a minimal promoter sequence that is inert to position effects upon random Tol2-based integration and that features reproducible, robust interaction with numerous enhancer elements (Carney and Mosimann, 2018; Felker and Mosimann, 2016). Second, the widespread use of EGFP, mCherry, and other fluorophores that generate red and green fluorescent reporters has introduced a challenge in the generation of combinatorial transgenic lines (Don et al., 2017; Kwan et al., 2007; Villefranc et al., 2007). Third, marking the expression of a transgene carrying a gene of interest as a fusion protein is not always practical, and previous attempts to use internal ribosomal entry sites (IRES) sequences to generate bi-cistronic reporters have been generally unsuccessful for practical use in zebrafish (Kwan et al., 2007). Lastly, the limited collection of available destination backbones with validated transgenesis markers restricts combinatorial transgene use and at times limits observations in tissues dominated by common transgenesis markers such as the heart (promoter of myl7, formerly cmlc2) and the eye lens (various crystallin gene promoters) (Hesselson et al., 2009; Huang et al., 2003; Kwan et al., 2007; Villefranc et al., 2007).

To expand on previously available transgenesis tool sets including the Tol2kit (Kwan et al., 2007), we embarked on a collaborative effort to augment, diversify, and increase the functionality of available plasmid sets for transgenesis. Building upon components and applications generated by many before us (including, but not restricted to, Amsterdam et al., 1995; Asaoka et al., 2002; Bessa et al., 2009; Distel et al., 2009; Fisher et al., 2006b; Hesselson et al., 2009; Huang et al., 2003; Kawakami et al., 2000; Koga et al., 1996; Mosimann et al., 2011; Tamplin et al., 2011; Woolfe et al., 2004) and with the goal of accessible documentation and distribution to the community, we here present several plasmid vectors compatible with MultiSite Gateway and traditional restriction enzyme cloning. These constructs are geared towards facilitating gene-regulatory element discovery, transgene quality control, reporter analysis, and an expanded fluorophore spectrum for imaging. For all germline-transmitted, stable transgenic lines, we selected multiple founders and quality-controlled for expected Mendelian ratios in subsequent generations; we have been observing most transgenic lines for three or more generations if not noted otherwise. While our new backbones are geared towards Tol2-based transgenesis in zebrafish, the individual modules are deployable for numerous applications in diverse species, such as cavefish. Together, our vectors (Table 1, GenBank sequences and PDF maps in Supplemental Data File 1) expand the available modules for reproducible, quality-controlled Tol2 transgenesis and gene-regulatory element testing in zebrafish and other model systems.

Results

The mouse beta-globin (Hbb-bt) minimal promoter for universal enhancer testing

The discovery and isolation of gene-regulatory elements, such as enhancer elements, has been instrumental in decoding gene-regulatory input and in the creation of new transgenic tools to label and manipulate cell types of choice. Enhancers require a minimal promoter region associated with the transgene reporter to initiate RNA Pol II-based transcription (Haberle and Stark, 2018). Since reporter expression can be positively and negatively influenced by neighboring loci in the genome (Amsterdam et al., 1995; Jaenisch et al., 1981; Lalonde et al., 2022; Wilson et al., 1990), an expression construct designed to test an enhancer requires a minimal promoter that is not sensitive to other enhancers in the vicinity of the transgene insertion. Several minimal promoters have been applied in zebrafish transgenesis, including *Ef1a*, *E1b*, *hsp70l*, *c-fos*, *krt4*, *TK*, *gata2a* promoters that are several hundred to several kilo-bases in length (Amsterdam et al., 1995; Bessa et al., 2009; Johnson and Krieg, 1995; Kondrychyn et al., 2009; Li et al., 2010; Meng et al., 1997; Ogura et al., 2009; Scott et al., 2007; Shoji and Sato-Maeda, 2008; Weber and Köster, 2013).

The promoter activity of the mouse *beta-globin* minimal promoter (from the mouse *Hbb-bt* gene, 129 base pair sequence including 54 bases of 5' UTR) was initially characterized busing HeLa cells (Myers et al., 1986). Coupling of the mouse *beta-globin* minimal promoter to a promoter-less enhancer enables the generation of stable Tol2-based zebrafish transgenics with tissue-specific reporter activity, including fluorescent reporters and CreERT2 and KalTA4 driver lines (D'Agati et al., 2019; Kaufman et al., 2016; Prummel et al., 2019; Tamplin et al., 2011; Woolfe et al., 2004). Reporters including the mouse *beta-globin* minimal promoter show minimal to no background in transient injections and drive stable transgene activity in zebrafish, including when paired with enhancers from other species (Prummel et al., 2019; Tamplin et al., 2011; Tamplin et al., 2011; Woolfe et al., 2011; Tamplin et al., 2015; Woolfe et al., 2004). To enable testing and application of candidate enhancers in 5' entry vectors, we have generated middle entry vectors with mouse *beta-globin* minimal promoter-coupled m*Cerulean*, *EGFP*, *mCherry*, and *mApple* ORFs, as well as *creERT2* and the codon-optimized *Gal4*-activator, *KalTA4* (Distel et al., 2009) (**Fig 1A**, **Table 1** lists all generated plasmids used in this study).

To further illustrate the utility of the mouse beta-globin minimal promoter, we documented regulatory element testing of a mouse enhancer. The 668 bp-spanning downstream enhancer ECR2 in the mouse Bmp4 locus has previously been discovered to convey posterior lateral plate mesoderm activity when tested as a reporter in mouse embryos (Chandler et al., 2009). In zebrafish embryos, the ECR2 enhancer coupled with beta-globin-minprom:EGFP (mBmp4^{ECR2}_minprom:EGFP) (Fig. 1B) as stable transgene resulted in EGFP reporter expression in a dynamic expression pattern in different cell types, including trunk and tail muscles, pectoral fins, the heart, and the yolk-surrounding mesothelium (Fig. 1C). Notably, several of these reporter expression domains matched with the reported expression of zebrafish bmp4 (Fig. 1D-G) (Bisgrove et al., 2000; Lenhart et al., 2013; Nikaido et al., 1997; Stickney et al., 2007; Thisse and Thisse, 2005). The mouse beta-globin minimal promoter appears inert to position effects or non-specific episomal expression upon injection: testing of a reference construct harboring the phagederived T7 promoter that has no transcriptional activity in zebrafish (T7_minprom:mCerulean) reproducibly showed no discernible reporter expression at 2 dpf (n=136/136), while the transgenesis marker in cis revealed successful F0 injections (Fig. **1H-J**). Together with prior work, these data illustrate the utility of the mouse beta-globin minimal promoter for zebrafish transgenes based on isolated regulatory elements.

Components for fluorescent reporter generation

To expand available basic components for transgenic reporter generation, we revisited available fluorophores and 3' UTR constructs. The prevalent use of green fluorescent avGFP derivatives and red-fluorescent proteins (Chalfie et al., 1994; Rodriguez et al., 2017; Tsien, 2003) has greatly expanded the number of available transgenic zebrafish reporters, yet has led to a bottleneck in combining reporters for multi-color imaging. To extend the color spectrum available for transgenic applications with accessible tools, we have generated middle entry clones with cytoplasmic, membrane, and nuclear versions of mCerulean (Fig. 2D-F, Table 1). mCerulean is a monomeric, blue-fluorescent protein derived from avGFP and subsequent CFP with an excitation wavelength of 433 nm and an emission wavelength of 475 nm (Rizzo and Piston, 2005; Rizzo et al., 2004). Stable transgenes using mCerulean as reporter are well-tolerated in zebrafish (Fig. 2A) (Hesselson et al., 2009; Trinh et al., 2011; Zuniga et al., 2011). Spectral separation of mCerulean from GFP is easily achievable on common confocal and light sheet setups, either by using a 445 nm or 458 nm laser or using features of typical confocal software platforms to minimize spectral overlap (such as in Zeiss' ZEN software). We noted that depending on genetic background, mCerulean imaging on a scanning confocal microscope resulted in auto-fluorescence of skin or pigment cells, particularly in the head of zebrafish embryos and larvae (48 hpf onwards) when visualized with an Argon laser (488 nm) (Fig 2A-F). Use of the ideal excitation laser wavelengths greatly diminished this auto-fluorescence. Of note, mCerulean is easily visible on standard dissecting scopes with epifluorescence equipment, ideally using blue fluorescence filters and excitation spectra. mCerulean is excited and detectable by standard setups used for GFP fluorescence, requiring care to avoid misinterpreting mCerulean versus GFP signal in double-transgenic embryos.

To demonstrate the versatility of mCerulean, we imaged the transgenic reporter line *Tg(Hs-HOPX_4q12N3.1:minprom_mCerulean)* (subsequently abbreviated as *HOPX:mCerulean)*. *HOPX:mCerulean* includes the upstream region of the human *HOPX* gene together with the mouse *beta-globin* minimal promoter and resulted in stable expression in the notochord at 48 hpf (**Fig. 2A**). This expression is consistent with mammalian expression of *HOPX* in the developing primitive streak as well as in chordoma tumors (Loh et al., 2016; Nelson et al., 2012; Palpant et al., 2017). Notably, the human *HOPX* upstream region also drives expression in rhombomeres 3 and 5 in individual transgenic insertions (Chen et al., 2002). As a blue fluorescent reporter, *HOPX:mCerulean* combines well with mCherry-based reporters such as *drl:mCherry* that marks lateral plate mesoderm before restricting expression to cardiovascular lineages

(**Fig. 2A**) (Mosimann et al., 2015; Sánchez-Iranzo et al., 2018). Together, mCerulean provides a practical alternative fluorophore to expand the reporter spectrum in transgenic experiments.

Transgene constructs require a 3' UTR with polyadenylation signal (polyA) for productive mRNA transcription, stability, and translation (Felker and Mosimann, 2016; Fink et al., 2006; Matoulkova et al., 2012; Okayama and Berg, 1992). The most common 3' UTR with polyA sequences used for zebrafish transgenes are the bi-directional SV40 late polyA sequence (abbreviated as SV40 polyA) (Okayama and Berg, 1992) that was included in the original Tol2kit collection (vector #302) (Kwan et al., 2007) and the BGH polyA sequence (Higashijima et al., 1997). Several available Tol2 backbones for MultiSite Gateway assembly contain the SV40 polyA after the recombination cassette (Kwan et al., 2007); while practical, its presence can lead to assembly and sequencing issues when another SV40 polyA trailer is used in the 3' position. To expand the available 3' UTRs for transgene assembly, we isolated a 516 bp fragment of the 3' UTR of the zebrafish ubiquitin B (ubb, ubi) gene. We designed pA02 as p3E vector with versatile multiple cloning site to enable the generation of new 3' vectors by restriction enzyme cloning; transferring the ubb 3' region into pAP02 resulted in p3E_ubb-polyA. To test the p3E_ubbpolyA and compare it to the commonly used Tol2kit vector #302, we generated pCK005 ubi:mCerulean-ubb-polyA as well as pCK004 ubi:mCerulean-SV40-polyA, which provide convenient control vectors for co-injection along with newly generated reporter constructs of unknown activity (Fig. 2B,C, Table 1). In addition to this alternative SV40 polyA p3E vector, we also generated a p3E vector with the multiple cloning site from pBluescript KSII(+), providing single-cutter restriction sites for cloning inserts into the final plasmid (see Methods for details).

mCherry and multimeric dsRED versions are red fluorophores commonly used for zebrafish transgenesis (Davison et al., 2007; Kwan et al., 2007; Pisharath et al., 2007; Provost et al., 2007; Rodriguez et al., 2017; Traver et al., 2004). mApple (derived from *Discosoma sp.*) is a fast-maturing, monomeric red fluorophore with an excitation maximum of 568 nm, therefore less red-shifted than mCherry and still excitable by standard lasers on confocal microscopes (543 or 561 nm) (**Fig. 3A-G**) (Shaner et al., 2008). mApple also has a better theoretical quantum yield and a fast-folding mechanism, rendering it a potentially suitable, additional red monomeric fluorophore for any transgenesis toolkit. mApple is well-tolerated in zebrafish and has been successfully pioneered for red-fluorescent transgenic applications (Heim et al., 2014; Lou et al., 2015), yet remains less frequently used compared to mCherry or dsRED derivatives. Applied to

label endoderm membranes, mApple expression in stable *Tg(sox17:mApple-CAAX)* embryos showed faithful and well-detectable expression (**Fig. 3A,B**) akin to *Tg(sox17:EGFP-CAAX)* (**Fig. 3C,D**), complementing the existing and widely used cytoplasmic *sox17:EGFP* to label endoderm progenitors (Chung and Stainier, 2008; Sakaguchi et al., 2006). To expand the application of red-fluorescent mApple in zebrafish, we further generated combinations of *mApple* with the mouse *beta-globin* minimal promoter as middle entry clones (see Methods for details) with nuclear as well as membrane-localized versions (**Fig. 3E-G, Table 1**).

Together, our series of minimal promoter-coupled fluorophore vectors, multiple localization tags, plus *p3E polyA* with multiple cloning site vectors provide a basic set of reagents for versatile regulatory element discovery and simple fluorescent microscopy with standard equipment accessible in the field.

2A peptide clones for fusion protein reporters

Transgenic expression of a protein of interest in a specific cell population greatly benefits from fluorescently labeling of transgene-expressing cells. One method to directly label the expressed protein of interest is as fusion with a fluorescent tag; yet the impact of adding a large fluorescent protein to the protein of interest can result in mis-localization, sequestration, degradation, or inactivity of the fusion product (Giepmans et al., 2006). A widely deployed alternative is the translation of the protein of interest and a fluorescent reporter via a bi-cistronic message (Chan et al., 2011; Trichas et al., 2008). We previously released a series of vectors including IRES (internal ribosome entry site) sequences (Jang et al., 1988; Pelletier and Sonenberg, 1988) for such applications. However, IREScontaining constructs have been repeatedly reported to cause variable expression of the ORF (open reading frame) downstream of the IRES sequence (Ibrahimi et al., 2009; Wong et al., 2002). With IRES-containing constructs used in zebrafish, the ORF following the IRES (often a fluorescent reporter) is commonly expressed at a lower level than the first ORF. Individual users observed up to a 10-fold lower expression of the second ORF, while other data indicate that the presence of the IRES reporter can even dampen expression of the first ORF (Weber and Köster, 2013), severely limiting the practical utility of IRES constructs.

To provide an alternative strategy for fluorescent labeling, we have established 3' entry (p3E) vectors that include the PTV-1 2A peptide sequence, a well-characterized viral peptide that results in production of two proteins from the same message across model systems (Kim et al., 2011; Provost et al., 2007; Ryan et al., 1991). 2A-mediated "self-

cleavage" takes advantage of the ribosome not forming a glycyl-prolyl peptide bond at the C- terminus of the 2A peptide, allowing the translational complex to release the first polypeptide and continue translation of the downstream product without independent ribosome recruitment or additional host factors (Donnelly et al., 2001a; Donnelly et al., 2001b) (**Fig. 4A, Table 1**). The resulting equimolar protein amounts make the 2A peptide an ideal tool for the quantification of transgene expression or the controlled expression of multiple proteins from one transgene (Chi et al., 2008; Knopf et al., 2010; Knopf et al., 2011). We generated 3' entry constructs containing the 2A peptide-encoding sequence followed by different fluorophores (EGFP, mCherry, and mCerulean) in cytoplasmic, membrane, and nuclear forms (**Fig. 4B-D, Table 1**). These constructs are ready for use with middle entry clones that are in Gateway reading frame without a stop codon (see Methods, maps, and Gateway manual for details on primer design).

To evaluate and compare stoichiometry of protein production using 2A and IRES, we generated transgene expression constructs in which the same fluorophore (EGFP) was targeted to two different subcellular locations in transient injections: nuclear-localized EGFP (nls-EGFP) was in the first position and membrane-targeted EGFP (EGFP-CAAX) was in the second (**Fig. 4E, Table 1**). When separated by the 2A peptide, EGFP was easily visualized in both nuclear and membrane locations, even in cells expressing low levels of the transgene (**Fig. 4E**). When separated by the IRES, in most cells, only nuclear-localized EGFP was detectable; EGFP-CAAX was only detected in the brightest cells, confirming that the 2A peptide yields effective production of two separate proteins targeted to distinct subcellular locations (**Fig. 4F**). We quantified this result by measuring the nuclear:membrane fluorescence intensity ratio in individual cells (see Methods for details). IRES constructs yielded a much higher ratio of nuclear:membrane fluorescence compared to the 2A (Fig. 4G), indicating a greater expression difference of the two EGFP fluorophores from the same messenger RNA.

The utility of 2A-based fusions becomes apparent when applied with the Gal4/UAS system. Gal4-dependent expression of genes of interest as UAS-controlled transgenes is a powerful approach to test individual gene function and for disease modeling (Akitake et al., 2011; Distel et al., 2009; Halpern et al., 2008; Köster and Fraser, 2001; Provost et al., 2007). The ability to fluorescently label the UAS transgene-expressing cells provides an immediate phenotype readout; as 2A-based fusions harbor the fluorescent reporter C-terminally of the ORF, the fluorescence signal additionally provides an expression control of the entire coding sequence. Injection of UAS:kdr-2A-Cerulean into the cartilage-specific Gal4-driver transgenic line *col2a1aR2:KalTA4* (D'Agati et al., 2019; Dale and Topczewski,

2011) resulted in mosaic Cerulean expression in the developing notochord (**Fig. 4H**). Stable *UAS:kdr-2A-Cerulean* transgenic insertions (selected for with the *crybb1:TagBFP* transgenesis marker *in cis* on the transgene backbone *pCB24*) crossed to the same driver resulted in homogenous notochord expression (**Fig. 4I**).

Of note, using the 2A peptide in the Gateway system results in the addition of a 28 aa "tag" to the C-terminal end of the first protein (containing the translated *att* site and the 2A peptide) (Rothwell et al., 2010) (**Fig. 4A**). This residual amino acid sequence may disturb the protein of interest in the first position if it is sensitive to C-terminal tagging. To reverse the relative positions of the fluorophore and ORF of interest cloning, we also generated a 3' 2A entry vector containing the *MCS* from *pCS2* for cloning of genes of interest after the 2A peptide sequence that can be used with middle entry fluorescent reporters (see Methods for details). Together, our 2A-based vectors enable versatile tagging of proteins of interest in zebrafish transgenes and beyond.

A selective, inert transgenic marker labeling the pineal gland

Transgenesis requires rigorous quality control, which is facilitated by inclusion of an easily-to-screen, dominant reporter gene *in cis* with the intended transgene cargo. In practical terms, inclusion of an independent fluorescent transgene that is detectable for screening at a desired developmental stage enables identification of transgene-positive animals and tracking of the transgenic insertion across generations (Felker and Mosimann, 2016). The initial Tol2kit contained the destination vectors *pDestTol2pA2* as "empty" destination vector and *pDestTol2CG2* that included the *myl7* (formerly *cmlc2*) promoter driving EGFP in the developing myocardium from late somitogenesis stages on, as easily screenable by 24 hpf (Huang et al., 2003; Kwan et al., 2007). However, *myl7*-based transgenesis markers are not desirable for transgenes focused on heart development, a significant interest in the field. Further, the heart becomes increasingly obscured by skin pigmentation in wild type zebrafish strains, rendering transgene detection in adults challenging (White et al., 2008).

Eye lens-specific promoters of different *crystallin* genes have been used as powerful transgenesis markers (Davidson et al., 2003; Emelyanov and Parinov, 2008; Hesselson et al., 2009; Kurita et al., 2003; Marques et al., 2008; Waxman et al., 2008). *Alpha-* or *beta-crystallin* promoter-based transgenes (*cryaa* or *crybb1*) are visible throughout adulthood and are easily detected via UV flashlights and filter glasses (Felker and Mosimann, 2016). We previously adopted *cryaa:Venus* (Hesselson et al., 2009) to continuously mark effector transgenics such as CreERT2 drivers with a yellow fluorescent transgenic marker

(Mosimann et al., 2015). To complement existing reporter backbones, we generated $pCB59 \ crybb1:mKate2$ to serve as a red eye transgenesis marker (**Fig. 2A**) and $pCB24 \ crybb1:TagBFP$ as a blue eye transgenesis marker (**Fig. 4H,I**). The far-red mKate2 (Lin et al., 2012; Shcherbo et al., 2009) is still detectable using standard red fluorescence filters on dissecting scopes with excitation at 588 nm and emission at 633 nm, and is complementary to blue, green, or yellow fluorescent *crystallin* promoter-driven markers for combinatorial transgene experiments. TagBFP is a robust blue fluorophore with excitation at 402 nm and emission at 457 nm (Subach et al., 2008) that is well-tolerated in zebrafish as well (Singh et al., 2012). One drawback of *crystallin*-based reporters poses their expression initiation around 36-48 hpf (Emelyanov and Parinov, 2008; Hesselson et al., 2009; Kurita et al., 2003; Marques et al., 2008; Waxman et al., 2008), which is later than *myl7*-based reporters and close to the start of swimming that can render sorting of transgene-positive embryos challenging. Additionally, eye lens expression of fluorophores can interfere with imaging of head regions and is not desirable for labs and experiments focused on development or regeneration of the visual system.

To establish a transgenesis marker with a) early developmental expression to enable efficient screening, b) persistent fluorescent reporter expression that is unobtrusive to tissues of interest, c) focused expression in a small region in the zebrafish, and d) activity throughout the zebrafish life cycle, we sought to generate *Tol2* backbones with pineal gland-specific reporter expression. The deeply conserved pineal gland, pineal organ, or conarium is a small photosensitive endocrine gland forming at the anterior dorsal midline on the head and is involved in circadian rhythm control (Erlich and Apuzzo, 1985; Gheban et al., 2019). Importantly, in zebrafish the pineal gland condenses during somitogenesis, remains in place for the lifespan of the animal, is small, and at a different anatomical location compared to commonly used transgenesis markers (Asaoka et al., 2002; Kazimi and Cahill, 1999; Ziv et al., 2005).

The 1055 bp upstream region of the *exo-rhodopsin* (*exorh*) gene harbors a *pineal expression-promoting element* (*PIPE*) that drives specific, persistent transgene expression in the pineal organ as detectable by stereo microscopy from 22 somites on (Asaoka et al., 2002). We generated three destination vectors containing the *exorh* promoter driving *EGFP*, *mCherry*, and *mCerulean*, respectively (*pCK017 pDEST-exorh:EGFP*, *pCK053 pDEST-exorh:mCherry*, *pCK054 pDEST-exorh:mCerulean*) and tested their performance as transgenesis backbones (**Fig. 5A, Table 1**).

Upon injection with Tol2 transposase mRNA, all three reporter backbones resulted in prominent, robust, and reproducible pineal gland-restricted fluorophore expression starting from around 22 hpf, in line with the initial reports characterizing *exorh*-based reporters (**Fig. 5B-F**) (Asaoka et al., 2002). As stable transgenic insertions over at least three generations, *exorh:EGFP* recapitulated the transient expression (**Fig. 5B-D**, **Table 1**) with reporter activity that is easy to screen under a standard fluorescent dissecting scope from 24 hpf onwards while embryos remained in their chorions (**Fig. 5F**). We observed stable integration and expression of *exorh:EGFP* throughout zebrafish development and adulthood, when the pineal gland-specific fluorescence can be detected by UV filter glasses for simple, non-intrusive screening of transgene-positive adult zebrafish (**Fig. 5G,H, Supplemental Movie 1**). These data establish *exorh*-based transgenesis markers as a viable alternative to existing transgenesis markers in zebrafish.

A key application benefiting from transgene markers *in cis* is testing of gene-regulatory elements by transient injections and subsequent generation of stable transgenic lines (Felker and Mosimann, 2016; Fisher et al., 2006a; Fisher et al., 2006b; Kague et al., 2010). To facilitate the use of *exorh* reporter-based backbones for regulatory element testing beyond Multisite Gateway, we introduced the 2.4 kb upstream regulatory region of zebrafish *desmin a* (*desma*) together with *beta-globin*-coupled EGFP, mApple, mCherry, or mCerulean into Tol2 backbones carrying *exorh:EGFP*, *mCherry*, *or mCerulean* (*plus one version driving mCerulean in the cryaa:Venus-carrying Tol2 backbone*) (**Fig. 6A**, **Table 1**). These vectors enable traditional cloning for replacement of the *desma* regulatory sequences (flanked 5' by *EcoRI*, *EcoRV* and *NheI*, 3' by *Sall* and *EcoRI* sites) with other regulatory elements of interest in vectors that include a minimal promoter and a transgenesis marker *in cis*. Upon transient injections into wildtype zebrafish embryos, we observed robust somatic muscle expression of *pCK069 desmaMCS:minprom-mApple* (**Fig. 6B**, **Table 1**). Taken together, these experiments establish *exorh:fluorophore*-carrying Tol2 backbones as a new tool for marking transgenes with different cargo.

Enhancers can act at a distance on different promoters to activate transcription of target genes (Schaffner, 2015). This distant activation potential is detrimental when combining regulatory elements *in cis* on a confined transgenesis vector, such as a transgenesis reporter together with an enhancer-driven transgene. To assess the robustness and utility of the new *exorh*-based transgenesis backbones, we tested each *exorh:fluorophore* combination in assembled MultiSite Gateway constructs to determine if the *exorh* promoter cross-reacts with any reporters cloned *in cis*, in particular minimal promoter-containing reporters. In stable transgene integrations of HOPX_4Q12N3.1:minprom-

mCherry, exorh:EGFP, we observed faithful mCherry activity in the notochord and rhombomeres and EGFP expression in the pineal gland, as anticipated (**Fig. 6C,E,F**); however, we did not observe any mCherry signal in the pineal (**Fig. 6D**) or notochord expression of EGFP (**Fig. 6C**). While not comprehensive, this basic test indicates that both regulatory elements in our observed transgenic insertion were inert to the other's activities.

Crossing stable transgenics of *myl7:mCerulean,exorh:EGFP* with the *hsp70l:Switch* line that contains the *cryaa:Venus* lens marker in the backbone (Felker et al., 2018; Hesselson et al., 2009) further documented how the pineal gland marker can be used in combination with commonly applied transgene reporters (**Fig. 6G,H**). *exorh*-based fluorophore reporters such as *exorh:EGFP* are reproducibly strong and bright, enabling their use in ubiquitous fluorescent backgrounds, as documented with easily detectable *exorh:EGFP* expression and ubiquitous EGFP (**Fig. 6I,J**). Together, these observations establish *exorh:fluorophore* Tol2 backbones as new transgenesis vectors for a multitude of applications.

Zebrafish exorh: fluorophore reporters are functional in Astyanax

Establishing a new model organism greatly benefits from access to cross-compatible transgenesis reagents. The Mexican cavefish (*Astyanax mexicanus*) is an increasingly applied model system to study evolutionary selection and metabolism (McGaugh et al., 2020; Rohner, 2018). While transgenesis has been successfully introduced in the system including the use of originally zebrafish-derived regulatory sequences (Elipot et al., 2014; Lloyd et al., 2022; Stahl et al., 2019), a more widespread use has been hampered by the lack of functional transgenesis markers: *myl7*-based cardiac reporters seem not functional when tested in *Astyanax* (Elipot et al., 2014) and the diverse *crystallin* gene-based eye lens transgenesis markers cannot be used due to the eye degeneration phenotype in cavefish. We therefore sought to test the functionality of *exorh*-based reporters in the model.

Injection of a Tol2 expression construct carrying *exorh:EGFP* (**Fig. 7A**) into one-cell stage embryos of different populations of *Astyanax mexicanus* (derived from surface and Tinaja cave populations) resulted in easily detectable pineal gland expression beginning around hatching stage (1 dpf) in both Tinaja cave and surface morphs (**Fig. 7B-E**). At 5 dpf, we continued to observe robust expression in both morphs (**Fig 7C,E**), detectable from different imaging angles (lateral-dorsal-ventral) (**Fig. 7F-H**). Pineal expression of *exorh*-driven fluorophores continued throughout the larval stages of surface and cave morphs into adulthood (**Fig. 7I,J**). Tol2-based destination vectors engineered with the

exorh promoter driving a fluorescent marker (tested here as mGreenLantern and mCherry) (Campbell et al., 2020) also recapitulated the pineal gland expression pattern in transient injections (**Fig. 7K-O**) and can be applied to generate transgenesis constructs using MultiSite Gateway. Of note, in our transient injections we observed weak, mosaic expression in retinal cells of surface morph animals (**Fig. 7M**). Taken together, these observations document the activity of zebrafish *exorh* promoter-based reporters in *Astyanax* as a new transgenesis marker as supported by the developmental retention of the pineal gland in cave morphs.

Discussion

Tol2-based transgenesis and the wide accessibility of the required basic tools has led to a revolution of transgenic reporter and effector strains in the zebrafish field (Kawakami, 2005; Kawakami, 2007). The industriousness and collegiality of the zebrafish community has facilitated expansion beyond the basic tools as well as extensive sharing and distribution. While by no means comprehensive, our data and reagents presented here augment, diversify, and increase the functionality of the transgenesis toolkit for zebrafish and other systems. All plasmid sequences and maps are available with this manuscript (**Supplemental Data File 1**) and all reagents (**Table 1**) have been submitted for distribution by Addgene.

The widespread use of green and red fluorophores, in particular EGFP, mCherry, and dsRED2, increasingly hinders the combinatorial use of reporter transgenes. We provide a set of middle and 3' entry vectors containing mCerulean, a blue-fluorescent avGFP derivative, which provides a complementary fluorophore to common green and red fluorophores. mCerulean is detectable with standard GFP filters on fluorescent dissecting scopes as routinely used in the field, yet does benefit from using a dedicated blue filter set and matching excitation light source (**Fig. 2**). As an alternative to mCherry and dsRED2, mApple provides a complementary red fluorophore that is detectable with standard RFP filters (**Fig. 3**); we also found that antibodies against dsRED detect mApple to stain for its expression (data not shown). Combined with its better theoretical quantum yield and fast folding mechanism, mApple constructs provide a versatile red-fluorescent alternative to mCherry and dsRED derivatives.

Regulatory element testing is a powerful approach in zebrafish; the discovery and testing of regulatory elements from different species can reveal fundamental insights into upstream gene regulation as well as into the ontogeny of cell types (Fisher et al., 2006a;

Fisher et al., 2006b). As enhancer sequences need to act on a promoter region to support assembly of productive RNA Pol II complexes, transgene constructs for enhancer testing require a sensitive, yet position effect-inert minimal promoter region that is also ideally short to keep assembled vector sizes down (Felker and Mosimann, 2016). Several minimal promoter regions have been successfully deployed in zebrafish, including sequences from other species. The Xenopus Ef1a minimal promoter has been repeatedly used for basic transgenes, yet remains sensitive to silencing (Amsterdam et al., 1995; Tamplin et al., 2011). The zebrafish gata2a promoter region has been successfully applied in a variety of settings including enhancer trap vectors and reporters (Bessa et al., 2009; Meng et al., 1997); however, its comparatively large size (over 2 kb) can result in large transgene vectors. Previous work has provided solid evidence for the potency of the mouse beta-globin minimal promoter to test regulatory elements including notochord and hematopoietic enhancers (Tamplin et al., 2011; Tamplin et al., 2015; Woolfe et al., 2004). We have previously incorporated this promoter for testing enhancers active in the lateral plate mesoderm and neural crest across species (Kaufman et al., 2016; Parker et al., 2019; Prummel et al., 2019). Here, we provide vectors pairing the mouse beta-globin minimal promoter with EGFP, mCherry, mApple, and mCerulean for versatile generegulatory element testing (Fig. 1). While required for enhancers, upstream regions that contain the transcription start of a gene of interest might not require an additional minimal promoter as the provided sequences might suffice to assemble a functional RNA Pol II complex. Incorporation of the mouse beta-globin minimal promoter is advisable when testing regulatory elements from divergent species or when establishing basic regulatory element activity (Bergman et al., 2022). Notably, we also present here the pAP02 p3E MCS 3' entry vector that enables cloning of enhancer elements downstream of the reporter cassette, resulting in the enhancer getting incorporated into the resulting mRNA before being terminated and polyadenylated. This vector design will allow the potential application of methods to discover active regulatory elements by sequencing, such as successfully performed in cell culture systems as STARR-seq (Arnold et al., 2013; Neumayr et al., 2019). The pAP02 vector can also be used to clone cassettes for 3' barcoding (McKenna et al., 2016) or different polyadenylation sequences beyond the ones available to date.

Our collection of 2A peptide fusions with different fluorophores and localization tags is suitable to generate bi-protein constructs with minimal C-terminal tagging (**Fig. 4**). Previous attempts to broadly apply IRES sequences for bi-cistronic transgenes and fluorescent labeling have widely been unsuccessful for as-of-yet unclear reasons (Kwan et al., 2007). Viral p2A or similar peptides that break the emerging polypeptide chain during

translation have been widely applied to express several genes of interests at equal protein levels in diverse cell culture systems, such as for cardiac reprogramming (Muraoka and leda, 2015; Song et al., 2012; Zhang et al., 2019). The direct fluorescent labeling of cells expressing a gene of interest for modifier studies, functional tests, and disease modeling provides a potent phenotype readout and imaging tool (**Fig. 4**). The simple cloning requires omission of the stop codon in the N-terminal ORF of interest, cloned in-frame with the following *attB2*Gateway repeat to connect with a 3' p2A-fluorophore. However, the polypeptide breakage leaves a residual p2A sequence as well as the encoded Gateway repeat at the C-terminal end of the ORF of interest. Mechanistic tests are therefore warranted to ensure proper functionality of this tagging method in individual transgenes.

Versatile, easy-to-screen transgenesis markers greatly augment any model organism tool kit by enabling rigorous quality control and transgene identification. A frequently used transgenesis marker in zebrafish is based on the myl7 promoter that drives fluorescence in cardiomyocytes routinely from 20-24 hpf onward (Huang et al., 2003). However, the heart is obscured in the growing and adult zebrafish. crystallin gene-based reporters drive fluorescence in the eye lens, providing an elegant and simple reporter property for screening in adults using UV filter goggles and handheld flashlights. We here add two crybb1-based reporters driving mKate2 and TagBFP in Tol2-based Multisite Backbones to the roster (Fig. 2A, Fig. 4G,H). A drawback of crystallin-based reporters is their comparatively late onset of detectable fluorophore expression from around 36 hpf or later (Davidson et al., 2003; Kurita et al., 2003), which can render sorting of transgene-positive embryos challenging. Our exorh-based reporters documented here (Fig. 5,6) circumvent this issue and provide reporters that are reproducibly expressed from 22ss and stay active throughout adulthood (Asaoka et al., 2002); exorh-marked backbones are thereby screenable while embryos remain in their chorions as well as with UV filter goggles in adults (Supplemental Movie 1). The confined, small area of fluorescence dorsal and anterior in the head does not obscure major organs during early development and should still enable imaging of significant brain regions in fluorescent transgenic reporters or stained embryos. exorh:fluorophore cassettes are less than 2 kb in size, and we here documented several successfully germline-transmitted transgenes using such Tol2 backbones (Fig. 5,6). Notably, exorh-marked transgenes are easily combined with other transgenes with myl7-, crystallin-, or even ubiquitously expressed reporters (Fig. 6E-H). We here generated three distinct exorh:fluorophore combinations to facilitate planning and execution of such combinatorial experiments, and we provide versions compatible with restriction enzyme-based cloning and MultiSite Gateway (Fig. 5A). The provided

backbones that harbor *desma*-driven fluorophores enable restriction site-based exchange of the *desma* regulatory region to test different regulatory elements (**Fig. 6A**). In addition, *desma*-driven reporters express well in somatic muscle and are simple to detect (**Fig. 6B**), providing potent reagents to troubleshoot injections, to test *Tol2* mRNA batches, or to train new researchers in transgenesis basics.

Use of our vector tools is not confined to zebrafish, as many can be applied in other models and Gateway-compatible cloning reactions. We have previously used mouse *beta-globin*-containing enhancer constructs used for zebrafish transgenes without modifications for electroporation in chick (Prummel et al., 2019), underscoring the utility of the mouse *beta-globin* minimal promoter for enhancer testing. Relevant to other aquatic models, *exorh*-based reporter utility expands to *Astyanax*, an emerging model for the study of evolutionary adaptation (**Fig. 7**) (Elipot et al., 2014; McGaugh et al., 2020; Rohner, 2018). While the cave forms of *Astyanax* have lost their eyes, they do form functional pinealocytes and a pineal gland, although their impact on behavior and circadian rhythms remains to be fully elucidated (Lloyd et al., 2022; Mack et al., 2021; Omura, 1975; Yoshizawa and Jeffery, 2008). *exorh*-based reporter transgenes have the potential to facilitate transgene quality control, visual genotyping, and combinatorial transgene experiments such as Cre/*lox* approaches in the model. As the pineal gland remains visible on the skull of fishes and amphibians, *exorh*-based reporters have the potential to support transgenesis experiments in a wide variety of aquatic models.

Materials and Methods

Zebrafish Husbandry and Procedures

Animal care and procedures were carried out in accordance with the veterinary office of the University of Zürich, Zürich, Switzerland; the IACUC of the University of Colorado School of Medicine (protocol #00979), Aurora, Colorado, USA; and the IACUC of the University of Utah (protocol #21-01007), Salt Lake City, Utah, USA. All zebrafish embryos were incubated at 28.5 °C in E3 medium unless indicated otherwise. Staging was performed as described by Kimmel et al. (Kimmel et al., 1995).

Cavefish Husbandry and Procedures

Astyanax husbandry and care was conducted identical to reported procedures (Xiong et al., 2022) and as approved by the Institutional Animal Care and Use Committee (IACUC) of the Stowers Institute for Medical Research on protocol 2021-122. All methods described here are approved on protocol 2021-129. Housing conditions meet federal

regulations and are accredited by AAALAC International.

Plasmid Construction

All plasmids generated for this study are in the process of being deposited, along with maps and sequence files, with Addgene for non-profit distribution. First-time users of Tol2 technology are asked to connect with Dr. Koichi Kawakami to set up a material transfer agreement (MTA). Plasmid sequences and maps are available online accompanying this manuscript (**Supplemental Data File 1**). The full list of deposited vectors is found in **Table 1**.

Entry Clones

All entry clones were verified by colony PCR and sequencing with *M13 fw*: 5'-GTAAAACGACGGCCAG-3' and *M13 rv*: 5'-CAGGAAACAGCTATGAC-3'.

p5E

For *p5E* entry clones, all PCR products were amplified using Roche Expand High Fidelity PCR kit (Sigma 11732641001), which contains a 3'-adenine overhang-depositing Taq. PCR products were purified using the QIAquick PCR Purification Kit (Qiagen 28104) or the QIAquick Gel Extraction Kit (Qiagen 28706) using ddH2O elution as last step. PCR products were then TOPO-cloned into MultiSite Gateway entry vectors with the *pENTR5'-TOPO TA* Cloning Kit (Invitrogen K59120): 4 µL of purified PCR product was immediately combined with 1 µL *pENTR5'-TOPO TA* and 1 µL of salt solution and incubated at room temperature for a minimum of 2 hours to overnight to ligate. 3 µL of the TOPO reactions were then transformed with One ShotTM TOP10 Chemically Competent *E. coli* (Invitrogen C404010) and plated on Kanamycin-infused agar plates. Half reactions can also be used. Primer sequences for *pE5'* enhancer/promoter fragments are as follows:

pCK079 p5E Mm BMP4ECR2:

oBuS014 Mm BMP4 ECR2 fw: 5'-GGGGATGAAAGTAGCATCCTG-3' oBuS015 Mm BMP4 ECR2 rv: 5'-TTCCACTTTGCTTCCCAAACTGG-3' pAB019 pE5 Hs HOPX_4q12N3.1:

oAB019 Hs HOPX fw: 5'-GTGTTGGGTTAGTTTGAGC-3'oAB019 Hs HOPX rv: 5'-GTTCTGTTGGGGATATGTCC-3' pCK033 p5E desma MCS:oCK024 Dr desma MCS fw: 5'-gatatcACTGATgctagcTCCTTGAGGCACTTTCGG-3' oCK025 Dr desma MCS rv: 5'gtcgacTACGCTGTGTGAATGCTGG-3'

pME, pENTR/D, mouse beta-globin minimal promoter, and fluorophore ORFs

For middle entry vectors, blunt-end PCR products were generated using Phusion High Fidelity DNA Polymerase (NEB M0530S) and forward primers were designed including 5'-CACC-3' on the 5' end for directional cloning and then ligated into MultiSite GatewayTMcompatible *pME* entry vectors using *pENTR/D-TOPO* (Invitrogen K240020). 4 μ L of purified PCR product was combined with 1 μ L *pENTR/D-TOPO* and 1 μ L of salt solution and incubated at room temperature for a minimum of 2 hours to overnight to ligate. 3 μ L of the TOPO reactions were then transformed with One Shot TOP10 Chemically Competent *E. coli* and plated Kanamycin-containing agarose plates. Half reactions can also be used.

The mCerulean ORF sequence can be amplified using mCerulean fw: 5'-5'-ATGGTGAGCAAGGGCGAGGAGC-3' and mCerulean rv. CAGCTCGTCCATGCCGAGAGTG-3'. The mApple ORF sequence can be amplified using 5'-ATGGTGAGCAAGGGCGAGG-3' 5'mApple fw: and mApple rv: CTTGTACAGCTCGTCCATGC-3'.

р3Е

pAP02_p3E-MCS was generated by amplifying the multiple cloning site from *pBluescript KSII(+)* with the primers *AP13fw*: 5'-*GGGGACAGCTTTCTTGTACAAAGTGGGCGGCCGCTCTAGAA-3*' and *AP14rv*: 5'-*GGGGACAACTTTGTATAATAAAGTTGGGTACCGGGCCCCC-3*' and inserted into *pDONRP2R-P3* via BP reaction.

P3E_ubb-polyA was cloned by amplifying a 516 bp fragment of the zebrafish *ubiquitinB* (*ubb, ubi*) locus using the primers *ubb polyA fw*: 5'-TAGAACCGACAGTCTTAGGGATGG-3' and *ubb polyA rv*: 5'-GAATTCATTGCCATCAAGTGTTAGC-3' with Phusion High Fidelity DNA Polymerase (NEB M0530S), subcloned into the Zero Blunt TOPO PCR Cloning vector (Invitrogen K283020), digested out with BgIII and ligated into the single cutter BamHI site in *pAP02_p3E-MCS*.

Of note, frequent users of both the #302 p3E-polyA from the original Tol2kit (Kwan et al., 2007) and p3E_ubb-polyA introduced here have experienced decreased Multisite Gateway reaction efficiencies after multiple freeze-thaw cycles of these p3E vectors. For optimal colony numbers and reaction efficiency, we recommend freshly preparing

minipreps, facilitated by frequent retransformations of the vector and storage of the plasmid in small aliquots. The reason for this reduction in cloning efficiency with prolonged storage of these p3E vectors is currently unknown, but could possibly be due to structural factors such as secondary structure formation detrimental to Multisite Gateway assembly or bacteria-based propagation.

2A constructs were built via a two-step process. First, the 2A peptide-encoding sequence was added to the N-terminus of fluorescence protein ORFs using PCR and inclusion of the 2A peptide sequence in the 5' upstream primer. The fusion product was conventionally cloned into *pCS2FA* using the *FseI* and *AscI* sites (enzymes from NEB). This placed the fusion product upstream of the *SV40 polyA* signal sequence. Ligation reactions were transformed using Subcloning Efficiency *DH5a* chemically competent *E. coli* (Invitrogen/ThermoFisher) and plated onto Ampicillin- or Carbenicillin-containing agar plates. Cloning was confirmed by restriction digest and subsequent sequencing.

This 2A-FP-polyA cassette was then cloned into the 3' entry vector using common att PCR primers (2A-attB2F and CS2pA-attB3R). For p3E-2A-CS2MCS-pA, pCS2P+ was used as a template for a PCR reaction in which the 2A peptide-encoding sequence was included in the att2F forward primer. Care was taken to ensure that all primers used "Gateway reading frame", in which bases listed in the manual as "N' in the primer were included as "G". This ensures maintenance of reading frame between the middle entry clone (which must not include a stop codon) and the 3' clone containing the 2A peptide sequence and reporter. PCR products were gel-purified (Qiagen Gel Extraction Kit) and quantified. For the recombination, equimolar amounts (usually 50 fmoles each) of purified PCR product were combined with pDONR-P2R-P3 donor vector and Gateway BP Clonase II Enzyme Mix (Invitrogen/ThermoFisher). Reactions were incubated at room temperature for several hours and transformed using Subcloning Efficiency DH5a chemically competent *E. coli* (Invitrogen/ThermoFisher) and plated onto Kanamycincontaining agar plates. Cloning was confirmed by restriction digest and subsequent sequencing.

For 2A-FP cloning into *pCS2FA* (restriction sites in bold, sequence complementary to template underlined):

2AEGFP5Fse: 5'-gaac**GGCCGGCC** GGATCCGGAGCCACGAACTTCTCTCTGTTAAAGCAAGCAGGAGACGTGGAAGAAAA CCCCGGTCCT ATGGTGAGCAAGGGCGAGGAGCTG-3' 2AmCherry5Fse: 5'-gaac**GGCCGGCC** GGATCCGGAGCCACGAACTTCTCTCTGTTAAAGCAAGCAGGAGACGTGGAAGAAAA

CCCCGGTCCT ATGGTGAGCAAGGGCGAGGAGGAC-3' 2Anls5Fse: 5'-gaacGGCCGGCC GGATCCGGAGCCACGAACTTCTCTCTGTTAAAGCAAGCAGGAGACGTGGAAGAAAA CCCCGGTCCT ATGGCTCCAAAGAAGAAGCGTAAGG-3' EGFP3Asc (rev): 5'-gaacGGCGCGCCTCACTATAGGGCTGCAGAATCTAGAGG-3' mCherry3Asc (rev): 5'-gaacGGCGCGCCTTACTTGTACAGCTCGTCCATGCCGC-3' CAAX3Asc (rev): 5'-gaacGGCGCGCCTCAGGAGAGCACACACTTGCAGCTC-3' For *p3E* cloning from *pCS2FA*: 5'-2A-attB2F: qqqqACAGCTTTCTTGTACAAAGTGGGGGGATCCGGAGCCACGAACTTCTCTC-3' 2A-CS2PMCS-attB2F: 5'-ggggACAGCTTTCTTGTACAAAGTGGGG GGATCCGGAGCCACGAACTTCTCTCTGTTAAAGCAAGCAGGAGACGTGGAAGAAAA CCCCGGTCCT

AGGGATCCCATCGATTCGAATTCAAGGCCTC-3'

CS2pA-attB3R (rev): 5'-

ggggACAACTTTGTATAATAAAGTTGGAAAAAACCTCCCACACCTCCCCTG-3'

Expression Constructs

Multisite Gateway recombination reactions were principally performed as described in the Invitrogen Multisite Gateway Manual, with minor modifications. 10 fmol of pE5', pME, and pE3' vectors were combined with 20 fmol of destination vector, 1 µL of LR Clonase II Plus Enzyme Mix (Invitrogen 12538120, vortexed 2x for two seconds prior to use), and ddH2O for a final reaction volume of 5 µL. Vector calculations for molarity were performed using the Multisite Gateway Excel spreadsheet (Mosimann, 2022). Reactions were incubated at 25°C overnight and treated the following day with 1 µL 2 µg/µL Proteinase K for 15 minutes at 37°C and 3 µL of the reaction was transformed with One Shot TOP10 Chemically Competent E. coli (Invitrogen C404010) and plated on Ampicillin selection plates. T4 ligations into multiple cloning site plasmids were performed by amplifying fragments via PCR with Roche Expand High Fidelity Polymerase (Sigma 11732641001) with primers containing 5' restriction enzyme sequences for digestion following PCR purification. Digested backbones were treated with shrimp alkaline phosphatase (NEB M0371S) for 2 hours at 37 °C prior to gel extraction. T4 ligations were all performed with a 1:3 pmol ratio of backbone to insert at 16 °C overnight. 7.5 µL of T4 reactions were then transformed with One Shot TOP10 Chemically Competent E. coli (Invitrogen C404010) and Ampicillin selection. All assembled gateway reactions were tested by diagnostic digest and confirmed by sequencing using the primers below (Mosimann, 2022):

attB1 fw: 5'-CAAGTTTGTACAAAAAGCAGGCT-3' attB1 rv: 5'-AGCCTGCTTTTTTGTACAAACTTG-3' attB2 fw: 5'-ACCCAGCTTTCTTGTACAAAGTGG-3' attB4 fw: 5'-CAACTTTGTATAGAAAAGTTG-3'

Zebrafish exorh promoter element

The exorh promoter: EGFP was amplified from the pCR2.1 TOPO vector containing a 1055 bp upstream region and short coding sequence of the zebrafish exorh gene driving EGFP (a kind gift from Dr. Yoshitaka Fukuda and Dr. Daisuke Kojima) (Asaoka et al., 2002) with 5' Asp718I (Acc651) restriction site containing primers. Of note, exorh lies on the edge of a genomic contig in the sequenced zebrafish genome, rendering the promoter isolation challenging without the pioneering work by Asaoka et al.. The Asp718l-flanked exorh:EGFP fragment was subcloned into pENTR5 -TOPO as stated in the Entry Clone section above to generate pCK074 pE5' exorh:EGFP. The shorter 147 bp exorh promoter fragment driving EGFP was amplified from the same vector (Asaoka et al., 2002) and subcloned in the same way to generate pCK072 pE5' 147bp exorh:EGFP. The exorh promoter alone was also subcloned into pENTR5'-TOPO to generate pCK011 pE5' exorh. The vectors containing the exorh promoter driving mCerulean and mCherry ORFs were cloned using PCR fusion of Asp718I- flanked fragments and subcloned into pENTR5'-TOPO to generate pCK051 pE5' exorh:mCerulean and pCK052 pE5' exorh:mCherry, respectively. Of note, the exorh promoter sequence contains a GT-rich repeat region, resulting in challenging PCR amplification. Primer sequences are stated below:

pCK011 p5E exorh:

oCK001 exorh fw 5'- GCTCAGCTGGCAGTACTACC-3' oCK009 exorh rv 5'-GATGGAGAAGTGGACGATCG-3'

pCK072 p5E 147bp Exorh:EGFP:

oCK018 ASP7181-147bpexorh:EGFP fw 5'- agatctggtaccGCGAGCGCTGCTGTGTCTCC-3'

oCK017 ASP7181-exorh:EGFP Rv 5'-agatctggtaccTTACTTGTACAGCTCGTCC-3'

pCK074 p5E Exorh:EGFP:

oCK016 ASP7181-exorh:EGFP fw: 5'-agatctggtaccGCTCAGCTGGCAGTACTACC-3' oCK017 ASP7181-exorh:EGFP rv: 5'-agatctggtaccTTACTTGTACAGCTCGTCC-3'

pCK051 p5E Exorh:mCerulean:

oCK016 ASP7181-exorh:EGFP fw: 5'-agatctggtaccGCTCAGCTGGCAGTACTACC-3' oCK054 mcerulean-STOP-Asp7181 rv: 5'-GGTACCtaCAGCTCGTCCATGCCGAGAG-3'

pCK052 p5E Exorh:mCherry:

oCK016 ASP7181-exorh:EGFP fw: 5'-agatctggtaccGCTCAGCTGGCAGTACTACC-3' oCK056 mcherry-STOP-Asp7181 rv: 5'-GGTACCtaCTTGTACAGCTCGTCCATGC-3'

Destination Vectors

To generate destination vectors containing the *exorh* promoter driving various fluorophores as secondary transgenic markers, digested *Asp718l-exorh:fluorophore-Asp718l* fragments were T4-ligated as described above into the Asp718l-digested and dephosphorylated *pCM326 pDEST_cryaa:Venus* (Mosimann et al., 2015) following gel extraction of the digested backbone minus the *cryaa:Venus* cassette to generate *pCK017 pDEST exorh:EGFP*, *pCK053 pDEST exorh:mCherry*, and *pCK054 pDEST exorh:mCerulean*. Clones were verified by test digest and full plasmid sequencing. All destination vectors were transformed with One Shot *ccdB* Survival 2 T1R Chemically Competent Cells (Invitrogen A10460) with both Ampicillin and Chloramphenicol selection.

pCB59_crybb1:mKate2 was amplified from *pDEST-Tol2_Fli1ep-mCherry-GM130* (a kind gift from Dr. Holger Gerhard) (Franco et al., 2015) including the zebrafish *crybb1* promoter, parts of the minimal *CMV* promoter, and the mKate2 ORF. The PCR product was cloned into the *Tol2*-based Multisite Gateway backbone *pDestTol2CG2* (Tol2kit #395) (Kwan et al., 2007) via the *BgIII* restriction site. Of note, the reporter uses the terminal Tol2 repeat downstream of the insert as possible polyadenylation signal.

pCB54_crybb1:TagBFP, containing the zebrafish *crybb1* promoter and parts of the minimal *CMV* promoter expressing a zebrafish codon-optimized *TagBFP* ORF flanked by *Bgll1* restriction sites, was synthesized (GENEWIZ). After sequence verification, the fragment was cloned into the *Tol2*-based Multisite Gateway backbone *pDestTol2CG2* (Tol2kit #395) (Kwan et al., 2007) via the *Bgll1* restriction site.

The construction of cavefish-tested *pDEST_exorh:mGreenLantern* and *pDEST_exorh:mCherry* was performed using the NEBuilder HiFi DNA Assembly Cloning Kit (NEB E5520S). The original vector *pDestTol2A2* (Tol2kit #394) (Kwan et al., 2007) was digested with BgIII (NEB R0144S) and HpaI (NEB R0105S) restriction enzymes. The digested plasmid was purified using the Wizard SV GeI and PCR Clean-Up System (Promega A9281). The zebrafish *exorh* promoter region was amplified from *pCK029* (this manuscript) using the following primers

HpalBgl2_NEB_exorhprom:

ofwd:

GCCACAGGATCAAGAGCACCCGTGGCCGTATCTTCGCAGCTCAGCTGGCAGTACTA CCGC-3'

orev:5'-cccttgctcaccatggtggcGATGGAGAAGTGGACGATCGG-3'.

The mGreenLantern and mCherry fluorescent protein-coding sequences were amplified from *pTol2_tg:mGreen Lantern/mCherry* (a kind gift from Dr. Sumeet Pal Singh, IRIBHM, Belgium) using the following designed primers:

HpalBgl2_NEB_mGL_

ofwd: 5'- CGATCGTCCACTTCTCCATCgccaccatggtgagc-3'

orev: 5'- TATTTGTAACCATTATAAGCTGCAATAAACAAGTTttacttgtacagctcgtccatgtca-3' HpalBgl2_NEB_mCherry

ofwd: 5'-CGATCGTCCACTTCTCCATCgccaccatggtgagc-3';

orev:

5'-

TGCTTTATTTGTAACCATTATAAGCTGCAATAAACAAGTTttacttgtacagctcgtccatgcc-3' Purified amplicons were ligated to the digested *pDestTol2A2* according to the NEB HiFi DNA assembly protocol.

RNA Synthesis

Tol2 transposase-encoding capped mRNA was created by *in vitro* transcription using the SP6 mMessage mMachine Kit (Ambion, AM1340) from *pCS2+Tol2* plasmid (Kwan et al., 2007) linearized with Notl restriction digest. RNA was purified with Lithium Chloride precipitation followed by the Megaclear Kit (Ambion AM1908).

Zebrafish Injections and Transgenesis

All plasmids were miniprepped with the ZymoPURE plasmid miniprep kit (ZYMO D4212). Embryos were injected into the early cell of one-cell-stage wildtype embryos (*AB* or *TU*) with 1 nL injection mix containing plasmid DNA concentrations of 25 ng/µL and Tol2 mRNA at 25 ng/µL. Adult F0 animals mosaic for the transgene were outcrossed to wildtype and their progeny analyzed for stable transgene expression. Tol2-based zebrafish transgenics were generated using standard experimental protocols as specified in Felker and Mosimann (2016) to achieve single-insertion transgenics and reproducible quality control. Our transgenesis rate is routinely between 15–50% percent for each injected Tol2 transgenesis vector. We investigate several clutches from at least three independent F1 founders. All presented stable transgenic lines are at least F3 generation or beyond and were selected to transmit the transgenes at Mendelian ratios as outlined in Felker and Mosimann, 2016.

5'-

Imaging

Embryos or larvae were anesthetized with 0.016% Tricaine-S (MS-222, Pentair Aquatic Ecosystems, Apopka, Florida, NC0342409) in E3 embryo medium. Dissecting scope fluorescence imaging was performed on a Leica M205FA with a DFC450 C camera and 1.0x PlanApo M-Series objective, illuminated with a TL5000 light base and CoolLED pE-300^{white} Illumination System. We use the following Leica filter sets for fluorescence: ET GFP 10447408, ET CFP 10447409, and ET DSR 10447412 for imaging and sorting, and TXR LP 10450590 for routine sorting of red fluorescence. Laser scanning confocal microscopy was performed on a Zeiss LSM880 following embedding in E3 with 1% low-melting-point agarose (Sigma Aldrich, A9045) on glass bottom culture dishes (Greiner Bio-One, Kremsmunster, Austria, 627861). Images were collected with a 10/0.8 air-objective lens with all channels captured sequentially with maximum speed in bidirectional mode, with the range of detection adjusted to avoid overlap between channels. Maximum projections of acquired Z-stacks were made using ImageJ/Fiji69 and cropped and rotated using Adobe Photoshop 2022.

Nulear:Membrane Fluorescence Intensity Ratio Quantification

Fluorescence intensity of the membrane and nuclear EGFP, comparing the 2A and IRES was assessed by drawing a line across the cell in ImageJ/FIJI and measuring the fluorescence intensity profile along the line. Signal maxima in the nucleus and at the membrane were recorded; measurements were not used if signal was saturated. The fluorescence intensity ratio (nucleus:membrane) was calculated for each individual cell. Welch's t test was used to calculate significance. Box and whisker plots were generated using the ggplot2 package in RStudio. The upper and lower hinges correspond to the first and third quartiles. The upper whisker extends from the upper hinge to the highest value within (1.5×IQR), where IQR is the inter-quartile range. The lower whisker extends from the lower hinge to the lowest value within (1.5×IQR). Data points outside of the ends of the whiskers are outliers.

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Competing Interests

No competing interests declared

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Data availability

All plasmids described as basic vector collection (Table 1) have been deposited with Addgene for distribution for non-commercial use in research. All vector sequences and maps are included in this manuscript in Supplemental Data File 1. Transgenic lines, additional vectors described in the manuscript, and all other reagents are available from the authors upon reasonable request.

Author contributions

C.M., A.B., K.M.K. conceptualized the manuscript and the overall experiments, assisted in experiments, acquired funding, supported data gathering and interpretation, coordinated the collaborators, and co-wrote and finalized the manuscript draft; C.L.K. and plasmid vectors including exorh-based H.R.M. cloned reporters, performed microinjections, generated transgenic lines, assembled plasmid maps, gathered data and performed imaging, assembled figures and figure panels, and co-wrote the manuscript draft; K.M.K., B.F.M. and A.S. cloned constructs, performed microinjections, acquired imaging data and expression patterns, generated vector maps, and assembled figure panels; J.R.K., R.L.R., E.L., and C.B.L. generated the sox17-based reporters, acquired imaging data, assembled figure panels, and provided manuscript text; S.B. and N.R. performed and documented all Astyanax experiments, assembled figure panels, and provided manuscript text; S.N, S.B., G.D., A.P., A.F., K. D., S. B. and A.B. generated individual vectors including quality control by sequencing, generated vector maps, and

performed individual validations; C. B. and M. A. generated the Destination vectors with crybb1-based reporters, performed validations, and supported the manuscript draft.

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Figures and Table



Fig. 1. Incorporating the small mouse *beta-globin* minimal promoter into transgenic reporters. (A) Workflow of reporter generation or enhancer testing with Multisite Gateway recombination and Tol2 transgenesis. An enhancer fragment is cloned into a *p5E* vector and Gateway- or restriction enzyme-cloned into an expression construct composed of 5'

fragment, middle entry mouse beta-globin minimal promoter.fluorophore/KalTa4/creERT2, 3' SV40 polyA, transgenesis marker (optional). The resulting Tol2 expression construct is injected along with Tol2 transposase-encoding mRNA into one-cell stage zebrafish, randomly integrated into the genome, and screened to establish transgenic lines. Schematic generated using BioRender. (B) Schematic of mBmp4^{ECR2}:minprom:EGFP containing the distal mouse BMP4 ECR2 element, the mouse beta-globin minimal promoter driving EGFP and the SV40 polyA. (C-G) mBmp4^{ECR2} reporter expression in a variety of zebrafish tissues linked to endogenous bmp4 expression. Lateral view of stable transgenic line of *mBmp4^{ECR2}:minprom:EGFP* at 20 somite stage, asterisk marks developing heart, arrowhead the olfactory bulb (C). Lateral (D,E,G) and ventral (F) views of 3 dpf stable mBmp4^{ECR2}:minprom:EGFP transgenic larva (D) show expression in the heart (asterisk) and pectoral fin (arrowhead), as well as in muscles and fins (D,E), parts of the head musculature (F), and in the mesothelium surrounding the yolk as well as trunk musculature and fin fibroblasts (G). (H-I) The *mHbb-bt* minimal promoter has little to no background activity in transient injectios. (H) Schematic of expression vector T7 minprom:mCerulean using the bacterial phage T7 promoter (inactive in zebrafish), followed by the *mHbb-bt* minimal promoter and *mCerulean* ORF as reporter. Eye lensspecific cryaa: Venus cassette in cis as transgenesis control. (I,J) Representative 2 dpf embryo of a control or reference injection repeat with T7_minprom:mCerulean, imaged for mCerulean (I) and Venus (J) as reference for successful injection. Note the absence of detectable mCerulean expression (representative among 136 individually injected embryos in this replicate).

Stable transgenic, 48 hpf



Fig. 2. mCerulean as versatile blue fluorophore for zebrafish transgenesis. (**A**) Stable transgenic line for *HOPX4q12N3.1:minprom_mCerulean,crybb1:mKate2*, shows mCerulean expression in both the vacuolar and sheath cells of the zebrafish notochord at 2 days post-fertilization and eye lens-specific mKate2 expression (asterisk) when crossed with *drl:mCherry* transgenic, marking lateral plate mesoderm-derived lineages (cardiovascular system, blood at 2 dpf). (**B**,**C**) Transient injection of zebrafish *ubiquitin* (*ubb*) promoter-driven mCerulean with 3' polyadenylation trailer (*polyA*) from *SV40* (**B**) or *ubb* (**C**). (**D-F**) mCerulean constructs for different cellular localization. Transient expression of cytoplasmic mCerulean (**D**), nuclear nls-mCerulean (**E**), and membrane-bound mCerulean-CAAX (**F**).



Fig. 3. mApple as bright red fluorophore for zebrafish transgenesis. (**A-D**) Comparison of membrane-localized mApple-CAAX with EGFP-CAAX in stable sox17:mAppleCAAX and sox17:EGFPCAAX lines show matching expression in the developing pharyngeal endoderm at 28 hpf. Pharyngeal pouches shown as close-up with anterior to the left in max projection (**A**,**C**) and single slice views (**B**,**D**); scale bars 50 μm. (**E-G**) mApple fusion proteins for different localizations. Transient expression of cytoplasmic mApple (**E**), nuclear nls-mApple (**F**), and membrane bound mApple-CAAX (**G**).



Fig. 4. 2A peptide linkers for fluorescent protein tagging. (**A**) Schematic of protein product resulting from a Tol2 construct containing two protein sequences separated by a 2A peptide following the *attB2* site of a Multisite Gateway expression construct. Translation of the first protein occurs, the first polypeptide is released at the site of peptide separation, and translation of the second ORF proceeds with the same ribosome resulting in equimolar protein amounts. (**B-F**) Transient injections of *bactin2* promoter-driven fluorophore combinations separated by a 2A peptide tag; *nlsmCherry-2A-EGFPCAAX* (**B**), *nlsEGFP-2A-mCherry* (**C**), and *EGFP-2A-mCherryCAAX* (**D**). Transient injection of *bactin2:nlsEGFP-2A-EGFPCAAX* (**E**) demonstrates expression of EGFP in both the nucleus and membrane of cells whereas injection of *bactin2:nlsEGFP-IRES-EGFPCAAX* (**F**) results in poor EGFP-CAAX detection in cells positive for nuclear EGFP. (**G**) Ratio of nuclear to membrane fluorescence intensity with n = total number of cells assayed (numbers at base of graph); for each condition, cells from 5 different embryos were analyzed. Welch's t test (unequal variance), first and third quartiles boxed, bars to highest

value within 1.5 x inter-quartile range. (H,I) Use of 2A fusions in Gal4/UAS transgene experiments. Injection of UAS:kdr-2A-mCerulean,crybb1:TagBFP into stable transgenic col2a1aR2:KalTA4,myl7:EGFP that expresses codon-optimized Gal4 in the developing notochord and subsequent cartilage lineages shows mosaic mCerulean expression in the developing notochord (H), with the secondary myl7:EGFP transgenic marker noted with an asterisk and the crybb1:TagBFP marker noted with an arrowhead. Homogenous notochord expression of mCerulean in stable F2 UAS:kdr-2A-mCerulean,crybb1:TagBFP crossed with col2a1aR2:KalTA4, myl7:EGFP (I).



Fig. 5. Labeling the zebrafish pineal gland with the *exorhodopsin* (*exorh*) **regulatory region as transgenesis marker**. (**A**) Schematic of the Tol2 secondary transgenic marker backbones *pCK017 pDEST-exorh:EGFP*, *pCK053 pDEST-exorh:mCherry-*, and *pCK054 pDEST-exorh:mCerulean* including the *attR4-* and *attR3-*flanked *ccdB* cassette followed by the1055 bp *exorh* promoter driving *EGFP*, *mCherry*, or *mCerulean*. (**B-D**) Stable transgenic lines for EGFP-, mCherry-, and mCerulean-based *exorh-*driven fluorophore reporters; lateral, dorsal, and ventral views at 2 dpf showing specific, easily detectable reporter activity in the pineal gland. (**E-H**) Stages and appearances of *exorh-*driven transgenesis reporters. Starting at 22 somite stage (**E**), *exorh:EGFP* expression (black arrowheads) is easily detectable in individual embryos and in groups under a standard dissecting scope (**F**) while embryos are still in their chorions. *exorh-*driven fluorophore expression persists into adulthood as seen in dorsal (**G**) and lateral views (**H**); dissecting scope images.



Fig. 6. Application examples of an exorh-based, pineal gland-specific transgenesis reporter. (A,B) Injection-based, transient testing and stable transgenesis of regulatory elements with Tol2 vectors that harbor a transgenesis marker in cis for quality control. The desmin a (desma) upstream region paired with the mouse beta-globin minimal promoter driving different fluorophores with exorh- or cryaa-based (eye lens) transgenesis markers in *cis* in Tol2 backbones (A, see also Table 1). In all vector versions, the *desma* regulatory region is flanked by restriction sites as simple Multiple Cloning Site (MCS), enabling restriction enzyme- based replacement of *desma* with a regulatory element of interest (A). Injection shows muscle-selective desma activity (mCherry) and pineal-specific transgenesis reporter expression (EGFP) (B). (C-F) Regulatory element testing with exorh:EGFP as transgenesis marker in cis. Stable transgene integration of HOPX^{4Q12N3.1}:minprom-mCherry,exorh:EGFP at 3 dpf shows mCherry expression in the notochord and rhombomeres, while EGFP expression is confined to the pineal gland (C, arrowhead). The transgenes do not cross-activate mCherry (D,F) or EGFP (E,F) (inserts, white arrowheads) despite being encoded in cis and in close proximity, suggesting the exorh regulatory region is inert. The white asterisk (E) indicates faint, patchy EGFP signal in the eye. (G,H) Combinatorial use of transgenesis markers. Ventral (G) and lateral (H) views of a stable myl7:mCerulean, exorh: EGFP transgenic crossed to cryaa: Venus and imaged at 2 dpf; note distinct expression patterns of the secondary transgenic markers

labeling the heart (*myl7*), pineal gland (*exorh*), and eye lens (*cryaa*), respectively. (**I**,**J**) *exorh:EGFP* fluorescence is discernible in ubiquitous green-fluorescent transgenics. Ubiquitous *ubb:EGFP* (*ubi:Switch*) expression (**I**) still allows detection of *exorh:EGFP* expression (white arrowhead) (**J**).



Fig. 7. The exorh-based transgenesis marker functions in Mexican cavefish (*Astyanax mexicanus*). (A) Tol2 expression vector using the zebrafish *exorh* promoter driving the expression of EGFP; see also Figure 5. (B-E) Transient injection of *exorh:EGFP* shows reporter activity in the pineal gland of both Tinaja cavefish at 27 hpf (B) and 5 dpf (C), as well as in surface fish at 36 hpf (D) and 5 dpf (E). Pineal expression indicated with an arrowhead and embryo morphology traced with a dashed line. (F-H) Lateral, dorsal, and ventral views at 5 dpf showing specific, easily detectable EGFP reporter activity in the pineal gland (arrowheads). (I,J) Grayscale images of live *exorh:EGFP*-injected animals, detection of EGFP (white dots) in surface and cavefish adults (arrowheads). (K) Schematic of Tol2-based transgenesis backbones *pDEST-exorh:mGreenLantern* and *pDEST-exorh:mCherry* including the *attR4* and *attR3*-flanked *ccdB* cassette and the 1055 bp zebrafish *exorh* promoter driving *mGreenLantern* or *mCherry* and for Multisite Gateway cloning. (L,M) Injection of the *exorh:mGreenLantern*-containing Tol2 Destination vector shows pineal gland expression (arrowhead) as well as

patchy retinal expression in developing surface fish (asterisk). (**N**,**O**) Injection of the *exorh:mCherry*-expressing Tol2 destination vector, showing pineal gland expression in developing cavefish (arrowheads).

Table 1. Plasmid list

Plasmid #	Plasmid Name	Desription	Source
	5' entry vectors (<i>p5E</i> , KanR)		
pCK011	p5E exorh	exorh regulatory element (reg. elem.)	Mosimann
pCK074	p5E exorh:EGFP	exorh:EGFP	Mosimann
pCK072	p5E exorh_short:EGFP	exorh minimal (min., 147 bp) reg. elem. and EGFP	Mosimann
pCK051	p5E exorh:mCerulean	exorh:mCerulean	Mosimann
pCK052	p5E exorh:mCherry	exorh:mCherry	Mosimann
pCK033	p5E desma-MCS	desma (reg. elem.) in a multipe cloning site (MCS)	Mosimann
	middle entry vectors (<i>pME</i> , KanR)		
pCK002	pME mCerulean	mCerulean (cytoplasmic) no stop	Burger
769	pME mCerulean-CAAX	mCerulean-CAAX (membrane) stop	Kwan
765	pME nls-mCerulean	<i>mCerulean</i> (nuclear) stop	Kwan
763	pME mApple	mApple (cytoplasmic) stop	Kwan
768	pME mApple-CAAX	mApple-CAAX (membrane) stop	Kwan
764	pME nls-mApple	<i>mApple (</i> nuclear) stop	Kwan
pCK007	pME KalTa4	KalTA4 (optimized Gal4-activator) stop	Burger
pKD001	pME minprom_EGFP	mouse <i>b-glob</i> min. promoter (prom.) <i>EGFP</i> no stop	Mosimann
pSN001	pME minprom_mCerulean	mouse <i>b-glob</i> min. prom. <i>mCerulean</i> no stop	Burger
pCK006	pME minprom_mCherry	mouse <i>b-glob</i> min. prom. <i>mCherry</i> no stop	Mosimann
pCK068	pME minprom_mApple	mouse <i>b-glob</i> min. prom. <i>mApple</i> no stop	Mosimann
pGD001	pME minprom KalTa4	mouse <i>b-glob</i> min. prom. <i>KaITA4</i> stop	Burger
pGD002	pME minprom_creERT2	mouse <i>b-glob</i> min. prom. <i>creERT</i> 2 stop	Mosimann
	3' entry vectors (<i>p3E</i> , KanR)		
pAF020	p3E 2A-mCerulean-pA	2A-mCerulean-SV40polyA (cytoplasmic)	Mosimann
pAF018	p3E2A-mCerulean-CAAX-pA	2A-mCerulean-CAAX-SV40pA (membrane)	Mosimann
460	p3E2A-mCherry-pA	2A-mCherry-SV40pA (cytoplasmic)	Kwan
612	p3E2A-mCherry-CAAX-pA	2A-mCherry-CAAX-SV40pA (membrane)	Kwan
766	p3E2A-nls-mCherry-pA	2A-nlsmCherry-SV40pA (nuclear)	Kwan
457	p3E2A-EGFP-pA	2A-EGFP-SV40pA (cytoplasmic)	Kwan
458	p3E 2A-EGFP-CAAX-pA	2A-EGFP-CAAX-SV40pA (membrane)	Kwan
459	p3E 2A-nls-EGFP-pA	2A-nlsEGFP-SV40pA (nuclear)	Kwan
791	p3E 2A-pCS2MCS-pA v2	2A-MCS-SV40pA	Kwan
pAP02	p3E MCS	MCS	Mosimann
pGD003	p3E ubbpA	ubbpA (dr_ubb 3' UTR)	Mosimann
	destination vectors (pDEST, AmpR,	CmIR, ccdB)	

pCK017	pDEST exorh:EGFP	exorh:EGFP-SV40pA transgenesis marker	Mosimann
рСК053, pNR002	pDEST exorh:mCherry	exorh:mCherry-SV40pA transgenesis marker	Mosimann, Rohner
pCK054	pDEST exorh:mCerulean	exorh:mCerulean-SV40pA transgenesis marker	Mosimann
pNR001	pDEST exorh:mGreenLantern	exorh:mGreenLantern-SV40pA transgenesis marker	Rohner
pCB24	pDEST crybb1:TagBFP	crybb:TagBFP-SV40pA transgenesis marker	Affolter
pCB59	pDEST crybb1:mKate2	crybb:mKate2-SV40pA transgenesis marker	Affolter
	expression vectors (AmpR)		
pCK004	ubi:mCerulean-SV40pA	ubi:mCerulean-SV40pA	Mosimann
pCK005	ubi:mCerulean-ubbpA	ubi:mCerulean-ubbpA	Mosimann
pCK029	exorh:EGFP-SV40pA	exorh:EGFP-SV40pA	Mosimann
pCK036	desmaMCS: minprom_mCerulean- SV40pA,cryaa:Venus	<i>desma</i> reg. elem., mm <i>beta-globin</i> min.prom. <i>mCerulean, cryaa:Venus</i>	Mosimann
pCK069	desmaMCS: minprom_mApple- SV40pA,exorh:EGFP	<i>desma</i> reg. elem., mm <i>beta-globin</i> min.prom. <i>mApple</i> , <i>exorh:EGFP</i>	Mosimann
pCK081	desmaMCS:minprom_mCerulean- SV40pA,exorh:mCherry	<i>desma</i> reg. elem., mm <i>beta-globin</i> min.prom. <i>mCerulean</i> , <i>exorh:mCherry</i>	Mosimann
pCK082	desmaMCS:minprom_mCherry- SV40pA, exorh:mCerulean	<i>desma</i> reg. elem., mm <i>beta-globin</i> min.prom. <i>mCherry</i> , <i>exorh:mCerulean</i>	Mosimann
pCK083	desmaMCS:minprom_EGFP- SV40pA, exorh:mCherry	<i>desma</i> reg. elem., mm <i>beta-globin</i> min.prom. <i>EGFP</i> , exorh:mCherry	Mosimann



Movie 1. Adult *exorh:EGFP*-transgenic zebrafish observed using a NightSEA hand-held UV flashlight and UV filter glasses ("goggles"), demonstrating the persistent pineal gland expression of the reporter and the simple detection.

Table S1.

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Dataset S1. All sequences for the vectors outlined in Table as GenBank (.gbk) and additional PDF (.pdf) maps.

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