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22 Abstract

23 The red flour beetle, Tribolium castaneum, is an important model insect and agricultural pest. 24 However, many standard genetic tools are lacking or underdeveloped in this system. Here, we 25 present a set of new reagents to augment existing Tribolium genetic tools. We demonstrate a 26 new GAL4 driver line that employs the promoter of a ribosomal protein gene to drive expression 27 of a UAS responder in the fat body. We also present a novel dual fluorescent reporter that 28 labels cell membranes and nuclei with different fluorophores for the analysis of cellular 29 morphology. This approach also demonstrates the functionality of the viral T2A peptide for 30 bicistronic gene expression in Tribolium. To facilitate classical genetic analysis, we created lines 31 with visible genetic markers by CRISPR-mediated disruption of the yellow and ebony body color 32 loci with a cassette carrying an attP site, enabling future ϕ C31-mediated integration. Together, 33 the reagents presented here will facilitate more robust genetic analysis in Tribolium and serve 34 as a blueprint for the further development of this powerful model's genetic toolkit.

35

36 Introduction

37 The red flour beetle (*Tribolium castaneum*), a pest of stored agricultural products, has emerged 38 as a promising system for biological research. It is a representative of the order Coleoptera, 39 which comprises approximately 40% of known insect species and 25% of all known animals [1]. 40 While Drosophila melanogaster is by far the most popular insect model system, many aspects 41 of its development and physiology are not representative of insects in general, and so findings 42 in Tribolium may be more broadly applicable to insects in many cases. Furthermore, Coleoptera 43 includes significant agricultural pests such as the corn rootworm, Colorado potato beetle, and 44 Asian longhorn beetle, and so using *Tribolium* as an insect model may lead to advances in pest 45 control.

46 Advances in genetic tools have cemented the status of *Tribolium* as the second model 47 insect of choice behind Drosophila. Transgenic Tribolium may be obtained using various 48 transposons [2, 3] and more recently via CRISPR/Cas9-mediated genome editing [4]. Eye-49 specific fluorescent markers have also been developed to aid identification of transgenics [2]. 50 Transposition techniques have been used in for insertional mutagenesis, allowing the 51 identification of essential genes as well as enhancer traps [5]. Another valuable tool for 52 functional genomics, RNA interference (RNAi) via injection of eggs, larvae, or adults, has been 53 implemented in Tribolium, both in targeted studies [6-8] and in a large-scale screen of the 54 protein-coding genome [9]. Lastly, the GAL4/UAS system, a popular choice for spatiotemporally 55 controlled expression of a gene of interest in Drosophila, has been demonstrated to function in 56 *Tribolium* in the presence of a species-specific basal promoter [10].

57 Despite the proliferation of tools for genetic analysis and manipulation of Tribolium, 58 notable gaps remain. In the case of the GAL4/UAS system, only two driver lines are available, 59 one using a heat shock promoter [10] and the other making use of the odorant receptor co-60 receptor (Orco) regulatory regions [11]. Furthermore, Tribolium strains with visible phenotypic 61 markers of known genetic location, which are staples of classical genetic analysis, are not 62 readily available. Here, we present a set of reagents to address these issues and enhance the 63 utility of Tribolium as a genetic model organism. We first present a GAL4 driver line that 64 employs a ribosomal gene promoter to direct expression in the fat body and can serve as an 65 second effective marker for transgenesis. In addition, we describe a GAL4-inducible cellular 66 reporter in which the nucleus and endomembrane system are labeled with different fluorescent 67 proteins, acting as a robust means by which to analyze cellular structure, particularly with 68 respect to neurons. Furthermore, both the GAL4 and UAS cloning vectors are designed to 69 accept any gene or genomic region of interest to generate new drivers and reporters. We also 70 address the lack of visible phenotypic markers in *Tribolium* by using CRISPR to disrupt two 71 genes involved in cuticle pigmentation via homologous recombination with cassettes containing

- 72 an attP site to facilitate future genomic insertion of DNA of interest using the ϕ C31 integrase.
- 73 The tools presented here represent a valuable resource for the *Tribolium* research community
- and serve as a general template upon which further tools can be based.

75

76 Materials and methods

77 Tribolium husbandry and strains

- 78 All animals were raised at 28°C on a standard flour yeast mix. The following strains were
- 79 utilized: v^{W} and m26 [3].

80 **Vectors**

All vectors will be made available through the *Drosophila* Genomics Resource Center at Indiana
University (https://dgrc.bio.indiana.edu/Home).

83 **P119der**

119der, a derivative of pSLfa[UAS-Tc'Hsp-p-tGFP-SV40] (kindly provided by Dr. Gregor Bucher)
[10], was constructed by excision of tGFP with KpnI and NotI followed by replacement with the
sequence ACTAGTGAATTCAAAGTACCACTCGAGAGCGGCCGCG. This replacement
destroyed the KpnI site but preserved the unique NotI site and added a unique XhoI site.
(DGRC # XXXX)

89 **P130der**

p130der, a derivative of pSLfa[Hsp-p-Gal4Delta-SV40_attp] (kindly provided by Dr. Gregor
Bucher) [10], was constructed by digestion with BamHI and addition of the sequence
GGATCCAGGTACCAGCGGCCGCAGGATCC, containing unique KpnI and NotI sites. (DGRC
XXXX)

94 *pGZ286 (pBac-3xP3-EGFP-pTC006550-GAL4Δ)*

95 The basal *hsp68* promoter of PCR-linearized p130der was replaced with the *TC006550* 96 promoter amplified from blaAmp-Tc6550Pro-GFPZeo-Luciferase-HSP-Orange-pIZT (kindly 97 provided by Dr. Yoonseong Park) [12] with NEBuilder HiFi assembly (NEB). The resulting 98 p*TC006550-GAL4* Δ -SV40 polyA coding sequence was then amplified and inserted into PCR-99 linearized pBac[3xP3-EGFP] with NEBuilder HiFi assembly. (DGRC # XXXX)

100 pTC241 (pBac-3XP3-EGFP-UAS-nls-EGFP-T2A-mCherry)

101 The nls-EGFP-T2A-mCherryCAAX insert was amplified from pSYC-102 (a gift from Seok-Yong 102 Choi, Addgene plasmid #74790) [13] as two fragments, and assembled into the Notl site of 103 p119der using NEBuilder HiFi assembly. UAS-nls-EGFP-T2A-mCherryCAAX was then excised 104 using flanking AscI sites and ligated into pBac[3xP3-GFP]. (DGRC # XXXX)

105

106 Immunofluorescence and imaging

Tribolium larvae were dissected in PBS and tissue was fixed in 4% paraformaldehyde/PBS. The tissue was washed 2X with PBT (PBS and 0.8% Triton-X 100) and incubated with DAPI (final concentration of 0.1 μg/ml) for 10 minutes and then washed with PBS before mounting for imaging (modified from [14]). Confocal images were captured on a Leica SP8 confocal utilizing a Leica 63X oil immersion objective with a numerical aperture of 1.40. Light and fluorescent whole animal images were collected on a LeicaMZ10F dissecting microscope. All pictures were processed in Adobe Photoshop.

114

115 CRISPR

116 gRNAs were designed using CRISPRdirect [15] using the Tcas3 genome assembly for the 117 specificity check. Only high-quality gRNAs were selected. 20-mer protospacer sequences were 118 cloned into Bsal-digested pU6b-Bsal-gRNA [4] by NEBuilder HiFi-mediated ssDNA oligo

119 bridging as desbribed (https://www.neb.com/-/media/nebus/files/application-notes/construction-120 of-an-sgrna-cas9-expression-vector-via-single-stranded-dna-oligo-bridging-of-double-stranded-121 dna-fragments.pdf) using an ssDNA oligo consisting of the protospacer flanked by 25 bp regions 122 of vector homology. Protospacer sequences used were CCGGAAAATAATCTCCCAGT (vellow. 123 TC000802) and TTTCGTAAAAGTTTGAATCG (ebony, TC0011976) (DGRC #XXXX and 124 #XXXX). Homology donors consisting of an attP site and 3xP3-DsRed-SV40 polyA flanked by 125 loxP sites in the same orientation between 800 bp (yellow; left arm: ChLG2:7,663,877-126 7,664,676, right arm: ChLG2:7,664,677-7,665,476) or 726/739 bp (ebony; left arm: 127 ChLG9:13,340,543-13,341,268, right arm: ChLG9:13,341,269-13,342,007) homology arms were 128 synthesized by IDT and delivered in pUCIDT-amp. Mixtures consisting of 400-500 ng/mL each 129 of Cas9 plasmid [4], sqRNA plasmid, and donor plasmid were injected into v^{w} embryos.

130

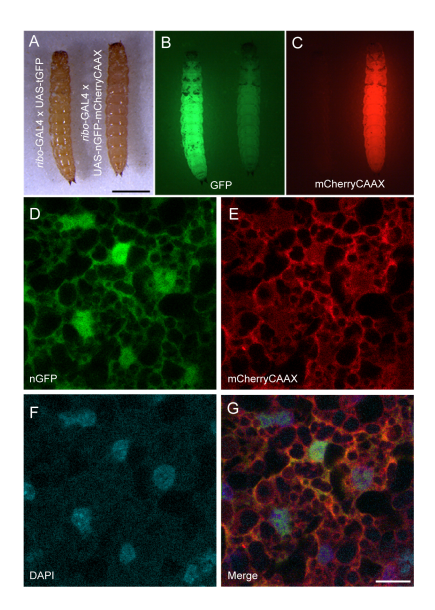
131 **Results**

132 Construction and implementation of a *ribo*-GAL4 driver

133 Inducible expression of a transgene of interest is a key capability in any genetic model 134 organism. In Drosophila, the bipartite Saccharomyces cerevisiae GAL4/UAS (upstream 135 activating sequence) system, in which the GAL4 transcription factor binds to a 17-bp motif within 136 the UAS to drive transcriptional activation, is frequently used for spatiotemporally controlled 137 expression of a gene of interest [16]. In Tribolium, the GAL4-UAS system has been 138 demonstrated to be functional when the UAS is coupled with the *hsp68* basal promoter and 139 driven by heat shock-inducible GAL4 [10]. However, no ubiquitous GAL4 drivers have been 140 reported for Tribolium. Such drivers are useful when screening for an organismal or 141 developmental phenotype of overexpression or knockdown.

142 In an attempt to generate a ubiguitous and constitutive GAL4 driver line, we considered 143 a previous study in which the promoter of a ribosomal protein gene (TC006550) was used to 144 drive high-level expression in the *Tribolium* TcA cell line [12]. We thus replaced the heat shock 145 promoter of our p130der vector (see Materials and Methods), bearing the basal hsp68 promoter 146 and GAL4 Δ , with the TC006550 promoter. GAL4 Δ is a variant of GAL4 in which the N- and C-147 termini, containing the DNA-binding and transcriptional activation domains, are directly fused 148 [17]. This variant of GAL4 has been shown to increase transactivation by ~2-fold in Drosophila 149 [18]. We then transferred the TC006550 promoter and GAL4 Δ coding region to a piggyBac 150 vector containing 3xP3-EGFP, enabling selection of transgenics by fluorescence in 151 photoreceptors. This vector was injected into a Tribolium line lacking eye pigmentation (vermillion^{white} (v^w)) with 3xP3-DsRed-marked piggyBac transposase integrated into the X 152 153 chromosome [3]. Resulting adults were outcrossed to v^{w} and progeny were assessed for GFP 154 expression in the retina. TC006550-GAL4 Δ transformants (hereafter referred to as ribo-GAL4) 155 were identified and the DsRed-marked transposase was removed through subsequent crosses 156 and two independent insertions were generated.

157 To assay the functionality of the *ribo*-GAL4 driver line, we crossed it to a previously 158 described UAS-GFP responder line [10]. Larvae displayed strong whole-body fluorescence (Fig. 159 1A-C) and fluorescence is maintained throughout pupal development and into adulthood as 160 expected for a ubiquitous expression. However, further examination revealed fluorescence was 161 only detected in the putative fat body (Fig. 1D-G) and absent from other tissues (e.g. the gut, 162 muscle, and CNS). We speculate that the lack of ubiquitous GAL4 expression in the ribo-GAL4 163 line may reflect tissue-specific differences in ribosomal protein gene expression [19] and that, 164 given the apparent in vivo expression profile of TC006550, the TcA cell line may be derived 165 from fat cells.



166

167 **Figure 1. Characterization of GAL4 and UAS vectors.**

(A-C) Tribolium larvae expressing either GFP (left) or nls-EGFP-T2A-mCherryCAAX (right)
driven by *ribo*-GAL4. (A) White light illumination; (B) GFP illumination. The nls-EGFP signal is
not detectable as compared to cytoplasmic GFP. (C) mCherry illumination. Scale bar = 1 mm.
The mCherry illumination mimics the cytoplasmic GFP expression. (D-G) Images of fat cells
expressing nls-EGFP (D) and mCherryCAAX (E), counterstained with DAPI (F), and the merge
of the three labels (G). Each represents a single confocal section. Scale bar = 10 μm.

175 Construction and implementation of a reporter for cell 176 structure

177 There are numerous reporters available for highlighting cell structure and function. Our goal was 178 to test whether these reporters could be simply swapped into a universal UAS cloning vector for 179 Tribolium with zero or minimal changes to the already existing sequence. We chose nls-EGFP-180 T2A-mCherryCAAX [13] to test the utility of bicistronic fluorescent reporter expression for 181 studying cell morphology as well as the use of the viral T2A peptide in Tribolium. When 182 combined with our *ribo*-GAL4 line, mCherry expression could easily be detected in whole larvae. 183 mimicking the spatial and temporal pattern obtained with cytoplasmic GFP (Fig 1A-C). To 184 confirm the expression and localization of both the nuclear GFP and the endomembrane linked 185 mCherry we examined the subcellular localization of each in fat cells. Colocalization with DAPI 186 confirmed the subcellular localization of GFP in the nucleus with mCherryCAAX bound to 187 membranes (Fig 1D-G). These results indicate that existing fluorescent reporters can be easily 188 implemented in Tribolium using our UAS vector and that the T2A peptide can be used for 189 multicistronic gene expression in Tribolium.

190

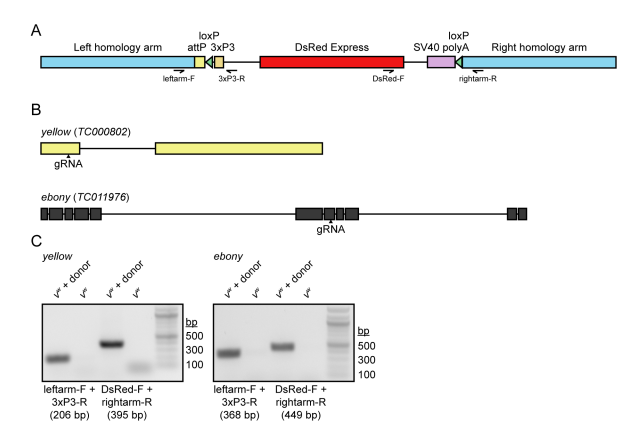
191 CRISPR-based generation of lines with visible phenotypic

192 markers

Defined phenotypic visible markers are essential for facilitating even basic tasks such as establishment of stocks of genetically modified organisms or tracking of specific chromosomes through multiple generations. While the scope of genetic reagents available in *Tribolium* has markedly increased over the past few decades, there are notable limitations and gaps in existing tools. For instance, the use of visible phenotypic markers for genetic mapping is quite limited, especially compared to *Drosophila*, where variations in wings, bristles, eye and body

pigmentation, and body shape are available. Additionally, while dominant mutations giving rise
to visible phenotypes in *Tribolium* have been documented [20-22], they have not been mapped
to a tractable genetic interval in most cases and so cannot be used in such analyses.

202 In order to create a general strategy for the expansion of a pool of visible phenotypic 203 markers for Tribolium, we employed CRISPR based homologous recombination. We designed 204 gRNAs against the coding regions of the yellow (TC000802) and ebony (TC011976) genes, as 205 well as a disruption cassette with useful genetic features. In many insects, the disruption of 206 either ebony and yellow leads to viable and fertile animals that are easily identifiable from their 207 wild-type counterparts. With respect to our CRISPR procedure, we chose a homologous 208 recombination strategy that would permit detection of a disruption in either locus regardless of 209 whether a change in pigmentation resulted. For each targeting construct, we utilized 700-800 210 bp homology arms, and between them we enclosed an attP site to facilitate future ϕ C31 211 integrase-mediated insertion of DNA of interest at a defined location [23]. More importantly, we 212 included the DsRed Express fluorescent protein under the control of the eye-specific 3xP3 213 regulatory element to facilitate screening of CRISPR based recombinants; and two loxP sites in 214 the same orientation, enabling future Cre-mediated excision of 3xP3-DsRed from the genome 215 (Fig. 2A). The positioning of the loxP sites is such that they flank only the fluorescent marker 216 and so excision would leave the attP site intact, effectively recycling the DsRed marker for 217 further use.



219

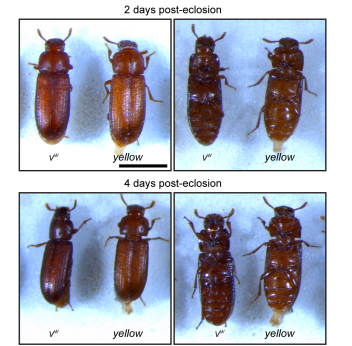
Figure 2. Construction and validation of the CRISPR gene disruption strategy.

(A) Schematic representation of the gene disruption cassette used in this study. The positions
and directionalities of primers used for genomic PCR validation are included in the schematic.
(B) Positions of gRNAs used to disrupt the *yellow* and *ebony* loci. (C) Genomic PCR
demonstrating the presence of each gene disruption cassette in the genome of transgenic but
not parental beetles.

226

 v^{w} embryos were injected with a mixture of Cas9 plasmids, gRNA expression plasmid, and repair template plasmids. Surviving adults were crossed to v^{w} and resulting progeny were screened upon eclosion for DsRed expression in the retina. For *yellow*, 308/356 injected larvae survived to adulthood and of these, two tested positive for germline transmission of the disrupted gene (0.65%). For *ebony*, 184/217 injected larvae survived to adulthood and of these, one tested positive for germline transmission of the disrupted gene (0.54%).

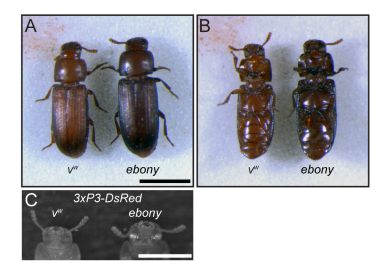
233 Characterization of *yellow*-edited adults revealed that cuticles of newly eclosed beetles 234 were noticeably lighter than those of their v^{w} counterparts, but their color darkened over time 235 until they were difficult to distinguish from the parental line (Fig 3). In contrast, adults with 236 homozygous disruption of ebony displayed substantially darker cuticular pigmentation than 237 parental v^{w} individuals (Fig. 4A-B) and DsRed fluorescence in their eyes (Fig. 4C). While *ebony* 238 CRISPR individuals were clearly phenotypically distinct from the parental line, their cuticle was 239 individuals injected with ebony dsRNA (http://ibeetle-base.uninot as dark as 240 goettingen.de/details/TC011976). We speculate that this difference is due to the location of the 241 gRNA target site. It lies within the seventh exon of ebony, which falls after the sequences 242 encoding all but one predicted functional domain of the protein (Fig. 5). However, this gRNA 243 falls at the start of the last domain of the protein, and disruption of this domain likely explains the 244 hypomorphic phenotype observed.



246

247 Figure 3. Phenotypic characterization of *yellow* CRISPR beetles.

- 248 Dorsal and ventral views of parental v^w and transgenic *yellow* CRISPR beetles at 2 and 4 days
- 249 post-eclosion. Scale bar = 1 mm.



250

251 Figure 4. Phenotypic characterization of *ebony* CRISPR beetles.

(A) Dorsal and (B) ventral views of parental v^w and transgenic *ebony* CRISPR beetles. Scale bar = 1 mm. (C) DsRed fluorescence microscopic image of the eyes of v^w and transgenic *ebony*

254 CRISPR beetles. Scale bar = 0.5 mm.

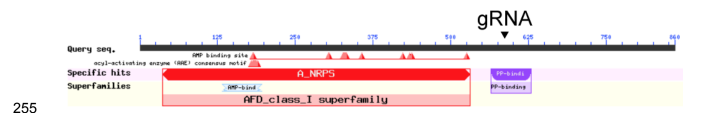


Figure 5. NCBI conserved domain search results for the *ebony* protein. The location of the gRNA relative to the mature protein sequence is indicated.

258

259 **Conclusions**

Here, we present a series of reagents aimed at increasing the genetic tractability of *Tribolium*. These are (1) a GAL4 driver plasmid and *Tribolium* line expressing GAL4 in fat cells; (2) a UAS plasmid and UAS-inducible dual-nuclear/endomembrane fluorescence reporter for analyzing cell structure; (3) a template gene disruption cassette for the generation of mutants and insertion of attP sites; and (4) *Tribolium* lines carrying disruptions of two body color loci, *yellow* and *ebony*.

266 We surmise that the failure of the *ribo*-GAL4 line to drive ubiguitous reporter expression 267 is attributable to its late pupal origin [24], reflecting cell type variability in TC006550 expression. 268 Indeed, heterogeneity in ribosomal protein expression across cell types has been widely 269 reported [19, 25-27]. Other potential candidates for the establishment of a ubiguitous GAL4 line 270 include the α -Tubulin1 promoter, which has been shown to drive ubiquitous GFP expression 271 throughout the Tribolium life cycle [28], and the Polyubiquitin promoter [29]. The p130der 272 plasmid permits efficient insertion of any potential genomic sequence for designing future GAL4 273 lines. Furthermore, our data demonstrates that any established reporter can be cloned into our 274 modified version of pSLfa[UAS-Tc'Hsp-p-tGFP-SV40] [10], p119der, for direct expression in 275 Tribolium. Future variations/deviations of both p130der and p119der will include an attB site for direct insertion into known genomic positions as well as a fluorescent marker to enable rapidscreening of transgenics.

278 While we successfully disrupted the *yellow* and *ebony* loci via CRISPR, the visible 279 phenotypes associated with these editing events were unpredictable. In particular, disruption of 280 yellow resulted in a slow-tanning phenotype, with young adults displaying a visibly lighter cuticle 281 than the parental strain that then darkened until it was indistinguishable from that of non-edited 282 beetles. In the case of *ebony*, we were able to achieve a marked darkening of the cuticle using 283 our disruption strategy, but our mutation was potentially hypomorphic when compared to *ebony* 284 RNAi, which yielded a darker black-body phenotype. In order to maximise the visible 285 phenotypes obtainable with CRISPR disruption of an eye or body color locus, we therefore 286 recommend pre-screening of candidate visible marker genes with RNAi prior to initiating 287 genome editing. Several candidates for body eye and body color genes have been assessed by 288 RNAi in the literature [30-33] and may serve as suitable targets for our CRISPR disruption 289 approach. Moreover, our data demonstrate that the inclusion of an independent marker for 290 CRISPR-based modifications is invaluable in recovering transformants and thus can mitigate 291 the uncertainty associated with the targeting of other potential candidate loci for visible markers.

292

293

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301

Primer	Sequence (5' \rightarrow 3')	Application
pTC006550-F	GATCGATATCGGCGCGCCAACAGC	Amplification of the <i>TC006550</i>
pTC006550-R	CTCTAGTACCAACCTTACGTTAGAA	promoter from blaAmp-
p	TTGAGTTACGAG	Tc6550Pro-GFPZeo-Luciferase-
		HSP-Orange-pIZT
p130der-F	ACGTAAGGTTGGTACTAGAGGTACA	Linearization of p130der for
	CGTCTCCC	insertion of the TC006550
p130der-R	TTGGCGCGCCGATATCGATCGCGC	promoter
	GCAGA	
pTC006550-	CATAGGCCACGGCGCGCCAACA	Amplification of the ribo-GAL4
GAL4-SV40-F		coding sequence from p130der
pTC006550-	CGGAGTGGACAGATACATTGATGA	
GAL4-SV40-R	GTTTGGACAAACCAC	
pBac-F	CAATGTATCTGTCCACTCCGCCTTT	Linearization of pBac[3xP3-
	AGTTTGATTATAATACA	EGFP] for insertion of the ribo-
pBac-R	TTGGCGCGCCGTGGCCTATGGCAT	GAL4 coding sequence
	TATTGTACGGA	
nls-EGFP-F	ACTAGTGAATTCAAAGTACCACTCG	Amplification of nls-EGFP from
	AGAGCATGGCTCCAAAGAAAGAAG	pSYC-102
	CGTAAGGTAAAT	
nls-EGFP-R	CCTTAAGCTTGTACAGCTCGTCCAT	
	GCCGA	
T2A-mCherry-	TCGGCATGGACGAGCTGTACAAGC	Amplification of T2A-mCherry-
CAAX-F	TTAAGG	CAAX from pSYC-102
T2A-mCherry-	GTGGTATGGCTGATTATGATCTAGA	
CAAX-R	GTCGCTCAGGAGAGCACACACTTG	
	CAGCTCATGCA	

302

303 S1 Table. DNA oligonucleotides used in this work.

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