piggyBac transformation of the New World screwworm, Cochliomyia hominivorax, produces multiple distinct mutant strains

M.L. Allen

A.M. Handler
US Department of Agriculture, Gainesville, U.S.A.

Dennis Berkebile
USDA-ARS, DENNIS.BERKEBILE@ars.usda.gov

S.R. Skoda
USDA-ARS

Follow this and additional works at: http://digitalcommons.unl.edu/entomologyfacpub


http://digitalcommons.unl.edu/entomologyfacpub/29

This Article is brought to you for free and open access by the Entomology, Department of at DigitalCommons@University of Nebraska - Lincoln. It has been accepted for inclusion in Faculty Publications: Department of Entomology by an authorized administrator of DigitalCommons@University of Nebraska - Lincoln.
piggyBac transformation of the New World screwworm, *Cochliomyia hominivorax*, produces multiple distinct mutant strains

M. L. ALLEN, A. M. HANDLER *, D. R. BERKEBILE and S. R. SKODA
Midwest Livestock Insects Laboratory, US Department of Agriculture, Lincoln and *Center for Medical and Veterinary Entomology, US Department of Agriculture, Gainesville, U.S.A.

Abstract. Sterile insect technique (SIT) programs are designed to eradicate pest species by releasing mass-reared, sterile insects into an infested area. The first major implementation of SIT was the New World Screwworm Eradication Program, which successfully eliminated the New World screwworm (NWS), *Cochliomyia hominivorax* (Coquerel) (Diptera: Calliphoridae), from the Continental US, Mexico and much of Central America. Ionizing radiation is currently used for sterilization, but transgenic insect techniques could replace this method, providing a safer, more cost-effective alternative. Genetic transformation methods have been demonstrated in NWS, and verified by Southern blot hybridization, PCR and sequencing of element insertion junctions. A lethal insertional mutation and enhancer detection-like phenotypic expression variations are presented and discussed. In addition to supporting the eradication efforts, transformation methods offer potential means to identify genes and examine gene function in NWS.

Key words. *Cochliomyia hominivorax*, green fluorescent protein, piggyBac, screwworm, sterile insect technique (SIT), transformation, transgenic insect

Introduction

The New World screwworm (NWS), *Cochliomyia hominivorax* (Coquerel), was once a costly pest of livestock and other warm-blooded animals, including humans, in the US and Mexico. The insect damages its host by feeding as a maggot on living tissue at a wound site. NWS has been successfully eradicated from the entire North American continent through a multinational programme; this eradication programme was the first implementation of the sterile insect technique (SIT). SIT theory states that a wild population can be eliminated if the pest insect can be economically mass-reared, effectively sterilized and dispersed at a sufficiently high ratio of sterile to wild males, there is no immigration and the sterilized insects are able to mate competitively (Knipling, 1955). The annual estimated benefits of NWS eradication to livestock producers in the US and Mexico exceed $1 billion (Wyss & Galvin, 1996) (http://www.aphis.usda.gov/bad/refbook2000/Screwworm.pdf). NWS are prevented from reinfesting North America by the ongoing release of sterile adult flies at the current barrier zone in Panama. Sterile flies for release are produced by irradiation in a highly secure mass-rearing facility in Tuxtla Gutierrez, Mexico (Wyss, 2000).

Transposable element-mediated insect transformation has been proposed as a means to improve and augment SIT programmes (Heinrich & Scott, 2000; Alphey, 2002; Handler, 2002; Horn & Wimmer, 2003). Applications for insect transformation include: unambiguous marking of released insects; restrictive rearing conditions (diet-suppressed conditional lethal); genetic sexing strain through female lethal selection (Heinrich & Scott, 2000; Horn & Wimmer, 2003), preferably at the larval stage; and genetically sterile insects, preferably males.

Correspondence: Margaret L. Allen, Midwest Livestock Insects Laboratory, US Department of Agriculture, Agricultural Research Service, 305 Plant Industry Building, University of Nebraska East Campus, Lincoln, NE 68583, U.S.A. E-mail: mallen2@unl.edu

© 2004 The Royal Entomological Society
Unambiguously marked strains of insects used in a SIT programme would benefit eradication efforts by clearly and quickly identifying production facility-reared vs. wild insects in cases of outbreaks or expanded releases. NWS is still indigenous to South America and the Caribbean, and if world-wide eradication of NWS is pursued, monitoring of progress would be facilitated by simple and quick identification of mass-reared insects. The current climate of world concern over bioterrorism demands that any outbreaks of NWS in non-endemic regions be rapidly evaluated for origination of infestation. Additionally, mass-reared screwworms capable of surviving only under specific permissive dietary conditions in artificial rearing would add to the measures of biosecurity, by eliminating survival of any insects that escape the rearing facility.

Any method of identifying and removing female screwworms from the insects reared for release would be of great value to the eradication process. If adult females were removed prior to release, more males (the effective sex in SIT) could be irradiated and packaged. If females could be identified and removed from rearing early in development, the cost of feeding the growing larvae in mass rearing could be greatly reduced. Also, the removal of females from the released population would reduce the expense for irradiating females. If females could be removed prior to release, more males (the effective sex in SIT) could be irradiated and packaged. If females could be evaluated for origination of infestation. Additionally, mass-reared screwworms capable of surviving only under specific permissive dietary conditions in artificial rearing would add to the measures of biosecurity, by eliminating survival of any insects that escape the rearing facility.

An ancillary use for insect transformation is the identification and testing, in vivo, of functional genes. The vast majority of insect species, including economically and medically important insects, will not be candidates for genomic sequencing in the near future. Therefore, if an insect such as NWS is amenable to genetic manipulation by transformation, position effects of the presumably random insertions of markers may provide interesting and useful genetic information (Bellen et al., 1989).

Here we describe the first transformation and establishment of stable colonies of transgenic NWS, C. hominivorax. We also discuss the unique characteristics of several distinct transgenic NWS strains, and their implications.

Materials and methods

Insect strains and rearing

Transformed C. hominivorax were derived from the P95 strain, which is the current strain used in the SIT programme. Insects were reared according to USDA-ARS and APHIS established protocols (Melvin & Bushland, 1936; Taylor et al., 1991), modified somewhat to accommodate rearing of small numbers of insects. Other NWS strains used as controls were J1, J5 and J14, colonies established in 1998 (J1) and 2002 (J5, J14) from specimens collected in Jamaica. Other species used in PCR studies were collected locally (Lincoln, Nebraska, U.S.A.); colonies were maintained temporarily in the Biosecure Facility in Lincoln and reared according to standard protocols.

Plasmids

A solution of 250 μg/mL helper plasmid (transposase-encoding) phsp-pBac, and 250 μg/mL vector pB[PUbnsEGFP] (Handler & Harrell et al., 1999, 2001a) in injection buffer was used for all microinjections. Plasmids were suspended in 6.25 mM Bis-Tris buffer with blue indicator dye. The pB[PUbnsEGFP] vector has the humanized enhanced green fluorescent protein (EGFP) variant gene (Clontech, Palo Alto, CA) under the regulation of the Drosophila melanogaster Meigen polyubiquitin promoter (Lee et al., 1988), which is normally active in all tissues throughout development. It is further linked to the SV40 T-antigen nuclear localizing sequence (nls) (Lanford et al., 1986).

Microinjection

Adult colonies of NWS were provided with substrate (raw ground beef) on which females deposited eggs for 15 min. Eggs were washed immediately with 1% NaOH solution to separate the masses (Berkebile & Skoda, 2002), and arranged on double-sided adhesive tape. The eggs were not dechorionated by this treatment, but the chorion was softened. Standard procedures for both D. melanogaster (Rubin & Spradling, 1982) and Aedes aegypti (Lineaus) (Morris, 1997) include injecting under oil; however, other insects have been transformed successfully without this step (Allen et al., 2001). NWS embryos were microinjected with plasmid solution at the posterior end, without oil, using bevelled borosilicate needles and an Eppendorf Transjector 5246™ system. Although a 37°C heat shock is normally applied to embryos after injection when using the piggyBac helper plasmid (Handler et al., 1998; Handler & Harrell, 1999), the NWS were reared at 37°C, so heat shock was unnecessary. All surviving injected insects were individually mated to wild-type P95 flies.

Screening and photography

Insects were screened with a Leica MZFLIII™ fluorescence stereomicroscope using the GFP2 filter set. The most efficient stage to screen was the crawler stage (fully mature third instars). Fluorescence could not be detected in the pupal or adult stage without dissection. Although fluorescence was visible in transgenic eggs, it was more apparent when the embryo was ready to hatch, but not in all strains. Removal of larvae from rearing medium prior to the crawler stage was often lethal. The crawler stage occurred after Day 5 of incubation, and once the larvae exited the rearing medium they pupated within 8 h. Thus, the screening step was restricted to a single day. Photographs were obtained using a Roper CoolSnap cf™ digital camera (Roper Scientific Inc., Tucson, AZ) accompanying Roper software, and Adobe Photoshop 6.0™ software (Adobe Systems Inc., San José, CA).
Genomic Southern blot hybridization

Genomic DNA (gDNA) was extracted from washed pupae using the Promega Wizard™ kit (Promega, Madison, WI), according to instructions provided by the manufacturer, incorporating a Proteinase K overnight digestion step. The gDNA specimens were digested with the following enzymes: HindIII, Sall, NotI and BglII, and size-fractionated on agarose gels, then transferred to a nylon membrane using standard procedures (Sambrook et al., 1989). A 580 bp product was PCR amplified from the vector plasmid pB[PUbnlsEGFP], 3’ piggyBac end (primers 1514F 5’-TGTTTTGACGGACCCCTTAC-3’ and 2094R 5’-CCAGAAACAACCTTGGCACA-3’). PCR was performed using ABI AmpliTaq Gold™ at the following conditions: 95°C 5 min, 55°C 15 s, 72°C 30 s) 35 times, 72°C 7 min, 4°C. The ampiclon was radioactively labelled using Amersham rediPrimeII™ random prime kit (Amersham Biosciences, Piscataway, NJ), and used to probe the genomic blot.

PCR

For inverse PCR, specimens of gDNA (approximately 2 mg/sample) were digested with Sau3A1 restriction endonuclease for 16–20 h. Half of the digested sample was cleaned (Zymo Research DNA Clean & Concentrator-5™, Orange, CA), and 1/50th of the material was ligated in a 40 µL reaction volume using T4 DNA ligase, again for 16–20 h. The ligations (2 µL used as template) were subjected to PCR amplification, using primers (in outward orientation) designed from the ends of the piggyBac element, under conditions recommended in the Expand Long Template PCR system (Roche Diagnostics Corporation, Indianapolis, IN) instructions: 93°C 2 min (94°C 10 s, 55°C 30 s, 68°C 6 min) 10 times (94°C 10 s, 55°C 30 s, 68°C 6 min + 20 s/cycle) 25 times, 72°C 7 min, 4°C. Nested primers were designed using Primer3 (Rozen & Skaltsky, 2000) for both the 5’ and 3’ ends of the piggyBac vector (5’ external primers used for first round PCR: 162R 5’-CACCGCGTCTGTATAGGTTCA-3’ and 357F 5’-TCCTCTCCTGC TCTCTCCTGAA-3’; 5’ internal primers used for second round PCR: 61R 5’-ACGGATT CCGC TATTTAGA-3’ and 380F 5’-GATGACGAGCTT TTGGG TGA-3’; 3’ external primers used for first round PCR: 1648R 5’-CCACTCCGCCCTTTAGTTTGA-3’ and 2099F 5’- TGTGCACAAAGTGTGTTCTG-3’; 3’ internal primers used for second round PCR: 1514R 5’-GATGACGAGCTT TTGGG TGA-3’ and 2385F 5’-CCTCGATATACAGACCGATA-3’). Amplicons were excised from agarose gels after brief visualization, and gel purified (Qiagen MinElute™, Valencia, CA). Some amplicons (approximately 1 ng template) were reamplified using nested primers and Applied Biosystems (Foster City, CA) AmpliTaq Gold™ PCR system (ABI) at the following conditions: 95°C 5 min (95°C 15 s, 55°C 15 s, 72°C 50 s) 35 times, 72°C 7 min, 4°C. Amplicons were sequenced by the University of Nebraska, Lincoln Genomics Core Facility, and compared to the known piggyBac terminal sequences and other published genomic sequences using NCBI BLAST programmes (Altschul et al., 1997).

Specimens of gDNA from NWS strains P95, J5 and J14, and gDNA extracted (as above) from Cochliomyia maccellaria (F.) and Lucilia sp. were digested with NotI restriction endonuclease. Primers were designed from the putative NWS genomic sequences, obtained by PCR amplifications described above, for insertion sites from two transgenic lines. These primers were designed to amplify across the TTA4 piggyBac insertion sites (GIZA 33F 5’-TCCTCT GATTTTGTCTTCA-3’, GIZA 366R 5’-AACGTTTCCT GCATAATATCCTG-3’; CLAY 53F 5’-AATGCTAC GACTCTTGC-3’, CLAY 241R 5’-AGTTGTTGCTCTT GCCGTGG-3’). PCR was performed using ABI AmpliTaq Gold™, for 35 cycles as above.

Results

Injection and strain establishment

A total of 2180 injected C. hominivorax (strain P95) embryos yielded a relatively low number of 49 adults (Table 1). Combined factors, including injection trauma

<table>
<thead>
<tr>
<th>Week</th>
<th>Total embryos injected</th>
<th>Total pupae collected</th>
<th>Percent pupated</th>
<th>Total adult survival</th>
<th>Adults M/F</th>
<th>Fertile adults M/F</th>
<th>GI* offspring screened</th>
<th>Transgenic adults M/F</th>
<th>Transformation frequency (%) (transgenic/fertile)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>260</td>
<td>20</td>
<td>7.7</td>
<td>3</td>
<td>3/0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0/0</td>
</tr>
<tr>
<td>2</td>
<td>898</td>
<td>32</td>
<td>3.6</td>
<td>12</td>
<td>5/7</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0/0</td>
</tr>
<tr>
<td>3</td>
<td>710</td>
<td>89</td>
<td>12.5</td>
<td>31</td>
<td>19/12</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0/0</td>
</tr>
<tr>
<td>4</td>
<td>312</td>
<td>14</td>
<td>4.5</td>
<td>3</td>
<td>2/1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0/0</td>
</tr>
<tr>
<td>Total</td>
<td>2180</td>
<td>155</td>
<td>7.1</td>
<td>49</td>
<td>29/20</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0/0</td>
</tr>
</tbody>
</table>

*Estimate based on 100 eggs per mass.

M = male, F = female

© 2004 The Royal Entomological Society, Medical and Veterinary Entomology, 18, 1–9
and decreased larval density in rearing medium, contributed to this relatively low survival. However, of the 22 adults that were successfully mated, three yielded transgenic offspring. Both of the transgenic matings in week 3 produced larvae positive for green fluorescence (EGFP+), 12 larvae from one mating and four from the other. In contrast, the week 4 transgenic mating produced more than 50 EGFP+ maggots. The former were individually outcrossed to P95 (wild-type) adults, but the latter were outcrossed in pools. Offspring of the two transformation-positive adults produced in week 3 were identified with the letters X and Y, and those of the transformation-positive adult produced in week 4 were designated as Z. The Z transgenic offspring appeared phenotypically homogenous, with fluorescence in the gut and salivary glands. The X progeny were less homogenous, but mainly characterized by fluorescence detectable in the salivary glands and in cells just under the cuticle. The Y progeny were the least homogenous, with some showing expression near the cuticle, the gut, salivary glands and the fat body, in varying combinations. Several generations of outcrosses resulted in eight stable colonies, representing seven distinct phenotypes, designated as: CL*YX, CLAY, COTY, FOLY, GARY, GIZA and SUEZ (* = I or O). Letters were added to the (X, Y, Z) labels to indicate the main expression pattern (C = cuticle, F = fat body, G = gut, S = spiracles) and to form pronounceable names.

The characteristics of the strains selected as putative single transgene insertions are shown in Fig. 1. CLIX and CLOX fluoresce in the salivary glands and in cells near the cuticle. Cuticular expression is most distinct at intersegmental areas, but is moderately bright throughout. CLAY is characterized by very bright expression under the entire cuticle, in the salivary glands, and pharyngeal filter. COTY fluoresces in both the salivary glands and at the intersegmental areas of the cuticle, but with much less intensity than CL/IOX or CLAY. The FOLY strain exhibits fluorescence in what appears to be fat body tissue and in the gut. GARY expresses fluorescence in the antennomaxillary palpi, the pharyngeal filter, salivary glands and in a central portion of the larval gut. GIZA expresses fluorescent protein in the salivary glands and a large segment of gut, and SUEZ expresses only around the anterior and posterior spiracles. Each strain has been stable for over 16 generations. Each appears to be homozygous, based on no wild-type individuals found in screened offspring for more than eight generations, except for the GIZA strain, which is lethal in the homozygous state (Table 2).

The GIZA strain was observed to produce substantial numbers of wild-type individuals every generation when all wild-types were discarded each generation. This led to our hypothesis that the piggyBac insert interrupted a vital gene, resulting in death of the homozygous transgenic offspring. Screened larvae were counted and compared to expected Mendelian ratios for a lethal gene mutation, and support the hypothesis of a lethal insertion (Table 2).

Molecular analysis

Southern blot analysis confirmed integration of the piggyBac vector cassette into the genome of the transgenic strains CLAY, CLIX, CLOX, COTY, GARY and GIZA (Fig. 2). Strains CLAY, CLIX, COTY, GIZA and GARY each contain a single copy of the transgene, whereas CLOX contains a duplicate insertion.

Inverse PCR was used to amplify the genomic sequence surrounding the piggyBac insertion site (Fig. 3). All sequences revealed the duplicated TTAA insertion site expected from piggyBac-mediated integration (Elick et al., 1996). Junction sequences from both 5' and 3' ends were obtained from strains CLAY, FOLY, GARY, GIZA and SUEZ. The putative genomic nucleotide sequences did not resemble one another or any other genomic sequences other than random 20 bp homologies (NCBI BLAST standard nucleotide comparisons); however, translated protein comparisons (BLASTx) of the sequences resulted in some homology. A total of 131 putative translated residues of GIZA sequence were 45% identical and 61% positive for an An. gambiae protein EAA12189 (gi: 21300044), and 42% identical and 63% positive for D. melanogaster CG6976-PB (gi: 24582545) and CG6976-PD (gi: 24582551), an unconventional myosin VII-like protein. A total of 132 putative translated residues of the CLAY sequence were 80% identical and 86% positive for D. melanogaster protein CG3996-PA (gi: 24655646) and 79% identical and 93% positive for An. gambiae protein EAA14637 (gi: 21302492).

If the amplified sequence surrounding the piggyBac insertion site is genomic NWS DNA, then primers based on sequences from each side (5' and 3') of a single insertion should amplify a predictable length of genomic DNA from wild-type NWS. Furthermore, if the insertion site is within a genomic region conserved between fly species, amplification of a similar length of genomic DNA might be obtained from related species. Using the NWS genomic sequences identified by inverse PCR, primers were designed that span the piggyBac insertion sites for the transgenic NWS strains GIZA and CLAY. These primers were used to amplify genomic DNA from wild-type NWS strains P95, J5 and J14 (two strains developed from recently collected Jamaican specimens), and two closely related blow flies, the secondary screwworm C. macellaria and a species of Lucilia (Phaenicia) found in Lincoln, Nebraska (U.S.A.). The GIZA primers amplified appropriately sized DNA fragments from all gDNA specimens, less efficiently in the non-conspecifics, and the CLAY primers only amplified from NWS specimens (Fig. 4).

Discussion

This is the first report of transformation of New World screwworm, and we have established eight stable colonies. The transformations have been confirmed by molecular analyses.
Fig. 1. Transgenic strains of Cochliomyia hominivorax, the New World screwworm (NWS).
The transposable element piggyBac has been used to genetically transform several insect species of economic importance. These include three species (of three genera) of fruit flies (Family Tephritidae) (Handler & Harrell, 1998, 2001c; Handler & McCombs, 2000), two lepidopterans (the pink bollworm and silkworm) (Peloquin et al., 2000; Tamura et al., 2000), three mosquitoes (Grossman et al., 2001; Kokoza et al., 2001; Lobo et al., 2002; Perera et al., 2002), the house fly (Hediger et al., 2001), the sheep blowfly (Heinrich et al., 2002), the red flour beetle (Berghammer et al., 1999) and a sawfly (Sumitani et al., 2003). The most closely related insect to have been transformed in the manner described here is Lucilia cuprina (Weidemann), the sheep blowfly (Heinrich et al., 2002). The transformation frequency reported for L. cuprina was lower than in NWS, but the number of surviving injected insects was much higher than our results. Transgenics of both species express EGFP in the embryo and larval stages, and in adult ovaries (data not shown). A great concern with NWS was the rapid embryonic development. The embryo hatches after 9 h at 37°C (standard mass-rearing protocol), and transformation protocol dictates that the plasmids are delivered to the developing embryo prior to cellularization of the syncitial blastoderm. Therefore, every effort was made to inject embryos as early as possible. We believe our injection technique improved over time (Table 1), as the number of GFP+ first generation transgenics was much higher (>50) in the final successful injection. Z. This leads us to speculate that an early germ line nucleus was transformed, which successfully replicated into many gametes. The X and Y transformation events may have occurred later in development, resulting in fewer sibling transformants.

The variation in expression patterns made it possible to isolate unique phenotypes associated with single transgene inserts that were amenable to inverse PCR and sequencing. Each of the insertions identified resulted in the canonical piggyBac TTAA integration specificity. Examination of the fluorescence expression can provide clues about the insertion site. Expression in cells near the cuticle appeared in four of the eight transgenic NWS strains, and was also described in PUbnlsEGFP transgenic Anastrepha suspensa (Loew) (Handler & Harrell, 2001b). This leads us to speculate that epidermal fluorescence may be a default expression pattern for PUbnlsEGFP in larval non-drosophilid dipterans. In adult tephritids (Handler & Harrell, 2001b) and drosophilid (Handler & Harrell, 1999) distinct fluorescence was noted in the thoracic muscles, but the NWS adult’s opaque cuticle made observation of fluorescence in live specimens impossible, and although dissected flight muscles of transformed specimens appeared fluorescent, similar fluorescence was observed in wild-type individuals (data not shown). Position effects will be analysed as more information about the insertion site characteristics becomes available. For future NWS transformations it would be useful to identify promoters expected to actively express in the embryonic and/or larval stages to facilitate transgenic screening.

The lethal insert in the GIZA strain was clearly validated by the ratios of GFP+ to wild-type larvae produced repeatedly after selecting only GFP+ individuals at every generation (Table 2). Although further sequencing will be required to identify and characterize the insertion site, the similarity of the genomic sequence surrounding the insertion site to a myosin gene also supports the assumption that the vector insert produced a lethal mutation. Myo28B1 in D. melanogaster is a

![Fig. 2. (A) Hybridization of vector probe to genomic Southern blots of restriction enzyme digested transgenic NWS. Lanes: 1, CLIX; 2, 4, GIZA; 3, 10, GARY; 5, P95 (wild type); 6, J1 (wild type), 7, CLAY, 8, COTY, 9, CLOX. (B) Diagrammatic representation of pB[PUbnlsEGFP] in transgenic NWS, showing location of restriction sites and probe. Single insertions should give single hybridization products larger than the vector lengths shown.](image-url)
single copy gene that encodes four isoforms of an unconventional myosin motor (FlyBase CG6976) (Yamashita et al., 2000). If this myosin motor is a component of basic cellular functions such as vesicle transport or actin organization, as has been suggested (Titus, 1999; Tuxworth & Titus, 2000), then insertional mutation would logically be lethal.

The genomic sequences identified through inverse PCR showed varying degrees of homology with closely related blowfly species. Clearly in the two primer sets tested, interspecies variation in homology was present (Fig. 4). Based on the translated BLASTx sequence homologies we predicted that both primer sets would amplify from closely related flies. It was unexpected that the more apparently conserved (79% identical residues) of the two translated sequences, CLAY, did not amplify products from C. macellaria or Lucilia. We intend to define and characterize these sequences further, and anticipate that they will be useful to identify and delineate populations of NWS.

The NWS eradication programme could benefit greatly from utilization of transgenic technology. The initial goal of producing genetically marked strains of NWS has been achieved as described here. Of the strains produced, one contains a lethal insertion, but other colonies appear vigorous. In-depth fitness comparisons of the new strains will elucidate both the appropriateness of transgenic strains for use in mass-rearing, and the potential for a genetically modified organism to persist, as it pertains to risk assessment (Ashburner et al., 1998). The production of genetically sterile strains or genetic sexing strains will depend on the availability of new transgenic tools such as inducible/repressible promoter systems, lethal (but environmentally safe) genes, and sex-specific genes and promoter systems.

In conclusion, the New World screwworm is an insect that lends itself well to piggyBac-mediated transformation. Laboratory colonies exist and rearing methods are well established in well-equipped biosecure facilities. With some effort even a single marker insertion can be utilized as an enhancer detection tool, in a similar fashion as methods used with D. melanogaster (O’Kane & Gehring, 1987; Bellen et al., 1989, 1990). Any use of transgenic insects in SIT or other pest control programmes will require the permission and cooperation of international regulatory agencies. Both the benefits to industry and wildlife, and environmental risks
should be thoroughly evaluated before mass release is implemented. Benefits of NWS SIT are well documented, and it is imperative that further benefits of transformation technology and associated risks be analysed carefully and rationally.

Acknowledgements

We thank the team of students and technicians at the USDA ARS MLIIRU BioSecure NWS Rearing Facility, and technicians at the USDA ARS CMAVE, Alan Christensen of the UNL Biology Department, and Ron Redder and Marilyn Jepson of the UNL Genomics Core Research Facility for their excellent assistance. David Stanley and Blair Siegfried of the UNL Entomology Department, and Ron Redder and technicians at the USDA ARS CMAVE, Alan Christensen of the UNL Biology Department, and Ron Redder and technicians at the USDA ARS CMAVE.

References


Accepted 26 November 2003