Inverse PCR and Sequencing Protocol on 5 Fly Preps<br>For recovery of sequences flanking piggyBac elements<br>This protocol is an adaptation of<br>"Inverse PCR and Cycle Sequencing Protocols" by E. Jay Rehm<br>Berkeley Drosophila Genome Project<br>And<br>"Single-Fly DNA Preps for PCR" by Greg Gloor and William Engels<br>Dept. of Genetics, University of Wisconsin<br>By<br>Ross Buchholz, Wes Miyazaki, Nick Dompe<br>and<br>Maddy Demsky, Stephen Thibault<br>Exelxis, Inc.<br>170 Harbor Way<br>South San Francisco, CA 94083

To prep the DNA for use with this protocol use the " 5 Fly Drosphila Genomic Prep for iPCR in 96-well Format" protocol.

## I. Things to keep in mind before you start this protocol

- Read the whole protocol before you start to make sure each step is clear.
- Ensure that you have all reagents and primers before you start.
- Keep all reactions on ice until they go into incubators or tetrads. Use metal 96-well plate holders that have been cooled to $4^{\circ} \mathrm{C}$ for best results. Place the plate in the metal holder, which is sitting on the ice for the duration of the setup. Use these to keep the DNA/donor reaction cool during reaction setup.
- Always add enzyme last to reagent mixtures. Do this just before you are ready to add the DNA or aliquot from the previous step which then starts the reaction.
- After reagent mixture has been added to plate wells, be sure to quick spin the covered plate to pull all liquid to bottom of plate.
- The polymerase chain reaction (PCR) is covered by patents owned by Hoffman-La Roche, Inc. and F. Hoffman-La Roche Ltd. Users should obtain a license to perform the reaction.
- AT ALL STEPS USE AEROSOL TIPS


## II. Reagents

This protocol optimized with the following reagents.

| Reagent | Manufacturer | Catalog |
| :--- | :--- | :--- |
| 96-well plate | Axygen | PCR-96-HS-C |
| Tape Pad | Qiagen | 19570 |
| Adhesive PCR film | Abgene | AB-0558 |
| Aluminum Sealing Film | Axygen | $47734-816$ |
| Sau3A I enzyme | New England Biolabs | R0169 |
| HinP1 I enzyme | New England Biolabs | R0124 |
| T4 DNA Ligase | New England Biolabs | M0202 |
| Tetrad thermal cycler | MJ Research | PTC-225 |
| AmpliTaq DNA polymerase | Perkin Elmer | E09425 |
| ExoSAP-IT kit | USB Corp. | 78200 |
| 96-well working rack | Stratagene | 410094 |
| BigDye | Applied Biosystems | 4331186 |
| ABI Prism 3700 | Applied Bioystems | 4308058 |
| 70\% ethanol |  |  |
| multichannel pipet, aerosol tips, centrifuge, vortex |  |  |

## III. Digestions (Sau3A I and HinP1 I done separately)

## Sau3A I digests are for 5' iPCR

HinP1 I digests are for 3' iPCR
Protocol per reaction is as follows:
$20 \mu \mathrm{l}$ reactions done in 96-well plate

| Genomic DNA ( $\sim 0.5$ fly) | $10.0 \mu \mathrm{l}$ |
| :--- | ---: |
| 10X buffer (NEB Sau3AI or NEB 2) | $2.0 \mu \mathrm{l}$ |
| 10X BSA (Sau3AI only) | $2.0 \mu \mathrm{l}$ |
| Sau3AI or HinPI | 4 units Sau3A1 or 5 units HinP1 |
| ddH2 O | add to $20 \mu \mathrm{l}$ total |

1) Cover plate with Adhesive PCR film.
2) Incubate @ $37^{\circ} \mathrm{C}$ for 1 hr in MJ Tetrad.
3) Incubate @ $65^{\circ} \mathrm{C}$ for 20 min . to heat inactivate.
4) Briefly centrifuge plate to spin down condensation.
5) Remove film to aliquot for ligations.
6) For storage @ $-80^{\circ} \mathrm{C}$, use Aluminum Sealing Film. Apply sheet to plate and incubate again @ $65^{\circ} \mathrm{C}$ for 15 min to seal plate. Briefly centrifuge plate to spin down condensation. Store @ $-80^{\circ} \mathrm{C}$.

## IV. Ligations

Protocol per reaction is as follows:
$10.5 \mu \mathrm{l}$ reactions done in 96-well plate

| Digested genomic DNA ( $\sim 0.075 \mathrm{fly}$ ) | $3.0 \mu \mathrm{l}$ |
| :--- | :---: |
| NEB 10X T4 DNA Ligase Buffer (w/ 10mM ATP) | $1.0 \mu \mathrm{l}$ |
| ddH $_{2} \mathrm{O}$ | $6.0 \mu \mathrm{l}$ |
| NEB T4 DNA Ligase (200 Weiss units) | $0.5 \mu \mathrm{l}$ |

1) If doing PCR immediately following ligation;

Incubate @ Room Temp for 30 min (cover plate with Tape Pad)
Remove Tape Pad and aliquot to $1^{\text {st }}$ round PCR.
For storage @ $-80^{\circ} \mathrm{C}$, use Aluminum Sealing Film. Apply film to plate and incubate @ $65^{\circ} \mathrm{C}$ for 15 min to seal plate. Briefly centrifuge plate to spin down condensation, store @ $-80^{\circ} \mathrm{C}$.
2) If not doing PCR immediately following ligation;

Apply Aluminum Sealing Film to plate and incubate @ Room Temp for 30 min, then incubate @ $65^{\circ} \mathrm{C}$ for 15 min to seal plate. Briefly centrifuge plate to spin down condensation, store @ $-80^{\circ} \mathrm{C}$.

## V. iPCR

PCR to be done in 96-well plates covered with Adhesive PCR film.
$1^{\text {st }}$ round iPCR: $20.0 \mu \mathrm{l}$ reaction

| Ligated genomic DNA ( $\sim 0.035$ fly $)$ | $5.0 \mu \mathrm{l}$ |
| :--- | :---: |
| 10 X dNTP $(2 \mathrm{mM}$ each $)$ | $2.0 \mu \mathrm{l}$ |
| forward primer $(10 \mu \mathrm{M})$ | $0.4 \mu \mathrm{l}$ |
| reverse primer $(10 \mu \mathrm{M})$ | $0.4 \mu \mathrm{l}$ |
| 10 X PE AmpliTaq buffer w/ $15 \mathrm{mM} \mathrm{MgCl}{ }_{2}$ | $2.0 \mu \mathrm{l}$ |
| ddH $_{2} \mathrm{O}$ | $9.9 \mu \mathrm{l}$ |
| PE AmpliTaq (2 units) | $0.3 \mu \mathrm{l}$ |

## piggyBac iPCR

1) $95^{\circ} \mathrm{C} 5 \mathrm{~min}$
2) $95^{\circ} \mathrm{C} 30 \mathrm{sec}$
3) $55^{\circ} \mathrm{C} 1 \mathrm{~min}$
4) $72^{\circ} \mathrm{C} 2 \mathrm{~min}$
5) GOTO $2 \times 34$
6) $72^{\circ} \mathrm{C} 10 \mathrm{~min}$
7) $12^{\circ} \mathrm{C}$ hold
8) Cycle on MJ Tetrad using "piggyBac iPCR" program with heated lid.
9) Centrifuge briefly to spin down condensation.
10) Do $1: 10$ dilution of $1^{\text {st }}$ Round PCR by adding $180 \mu 1 \mathrm{H}_{2} \mathrm{O}$.
$2^{\text {nd }}$ round iPCR: $20.0 \mu \mathrm{l}$ reaction.

| $1: 10$ diluted $1^{\text {st }}$ round DNA | $5.0 \mu \mathrm{l}$ |
| :--- | :---: |
| 10 X dNTP $(2 \mathrm{mM}$ each $)$ | $2.0 \mu \mathrm{l}$ |
| forward primer $(10 \mu \mathrm{M})$ | $0.4 \mu \mathrm{l}$ |
| reverse primer $(10 \mu \mathrm{M})$ | $0.4 \mu \mathrm{l}$ |
| 10 X PE AmpliTaq buffer w/ $15 \mathrm{mM} \mathrm{MgCl}{ }_{2}$ | $2.0 \mu \mathrm{l}$ |
| ddH $_{2} \mathrm{O}$ | $9.9 \mu \mathrm{l}$ |
| PE AmpliTaq (2 units) | $0.3 \mu \mathrm{l}$ |

4) Cycle on MJ Tetrad using "piggyBac iPCR" program ( $\sim 3 \mathrm{hr}$ run) with heated lid.
5) Optional: Examine $5 \mu 1$ of the $3^{\prime} 2^{\text {nd }}$ round and $5^{\prime} 2^{\text {nd }}$ round iPCRs on $1.0 \%$ agarose gel.

Primers for $1^{\text {st }}$ and $2^{\text {nd }}$ round iPCR:

| Primer Name | PCR <br> Round | piggyBac <br> end | Primer Sequence 5' to 3' |
| :---: | :---: | :---: | :---: |
| 5F1 | $1^{\text {st }}$ | 5' end | 5' GAC GCA TGA TTA TCT TTT ACG TGA C 3' |
| 5R1 | $1^{\text {st }}$ | 5' end | 5' TGA CAC TTA CCG CAT TGA CA 3' |
| 5F2 | $2^{\text {nd }}$ | 5 ' end | 5' GCG ATG ACG AGC TTG TTG GTG 3' |
| 5R2 | $2^{\text {nd }}$ | 5' end | 5' TCC AAG CGG CGA CTG AGA TG 3' |
| 3F1 | $1^{\text {st }}$ | 3' end | 5' CAA CAT GAC TGT TTT TAA AGT ACA AA 3' |
| 3R1 | $1^{\text {st }}$ | 3' end | 5' GTC AGA AAC AAC TTT GGC ACA TAT C 3' |
| 3F2 | $2^{\text {nd }}$ | 3' end | 5' CCT CGA TAT ACA GAC CGA TAA AAC $3^{\prime}$ |
| 3R2 | $2^{\text {nd }}$ | 3' end | 5' TGC ATT TGC CTT TCG CCT TAT 3' |

## VI. Pre-Sequencing Preparation

Strong and unique bands as well as smears from the iPCRs can be directly sequenced without extensive purification. Prior to sequencing, use the USB ExoSAP-IT kit to clean up an aliquot of the $2^{\text {nd }}$ round reactions. This kit uses Exonuclease I (degrades primers) and Shrimp Alkaline Phosphatase (degrades unincorporated nucleotides) to prepare the template for sequencing.

The protocol per reaction is as follows:

## ExoSAP protocol

Done in 96-well plates covered with Adhesive PCR film Make a master mix per reaction of:

| Exonuclease I $(10 \mathrm{U} / \mu \mathrm{l})$ | $1 \mu \mathrm{l}$ |
| :--- | ---: |
| Shrimp Alkaline Phosphatase $(2 \mathrm{U} / \mu \mathrm{l})$ | $1 \mu \mathrm{l}$ |
| $\operatorname{ddH}_{2} \mathrm{O}$ | $3 \mu \mathrm{l}$ |

1) Remove $5 \mu 12^{\text {nd }} \mathrm{Rd}$ iPCR and add to $5 \mu \mathrm{l}$ SAP mix (to make $10 \mu \mathrm{l}$ total).
2) Run on "SAP" program on tetrad using heated lid.

SAP

1) $37^{\circ} \mathrm{C} 30 \mathrm{~min}$.
2) $85^{\circ} \mathrm{C} 15 \mathrm{~min}$.
3) $4^{\circ} \mathrm{C}$ hold
4) Do not hold @ $4^{\circ}$ overnight. The SAP prep should be done on the day that the sequencing reactions are to be done.

## VII. Cycle Sequencing Protocol for $\mathbf{3 7 0 0}$ ABI Machine

The protocol per reaction is as follows
$10 \mu 1$ reaction done in 96 -well plate

| DNA $(1 \mu \mathrm{l}$ from $10 \mu \mathrm{l}$ SAP prep $)$ | $1.0 \mu \mathrm{l}$ |
| :--- | ---: |
| Primer $(0.8 \mu \mathrm{M})$ | $4.0 \mu \mathrm{l}$ |
| 5X BigDye buffer | $1.5 \mu \mathrm{l}$ |
| ABI BigDye $(\mathrm{v} 3.0)$ Mix | $1.0 \mu \mathrm{l}$ |
| $\mathrm{ddH}_{2} \mathrm{O}$ | $2.5 \mu \mathrm{l}$ |

2) Cycle Sequence ( $\sim 2.5$ hours)

BigDye

1) $96^{\circ} \mathrm{C} 4 \mathrm{~min}$
2) $96^{\circ} \mathrm{C} 30 \mathrm{sec}$
3) $50^{\circ} \mathrm{C} 15 \mathrm{sec}$
4) $60^{\circ} \mathrm{C} 4 \mathrm{~min}$
5) GOTO 2 X24
6) $12^{\circ} \mathrm{C}$ hold
7) To purify reactions add $75 \mu \mathrm{l} 70 \%$ ethanol, cover, let stand 30 minutes at room temp in the dark. Centrifuge for 30 minutes @ 2,470 RCF. Remove cover, invert on paper towel and spin @ 700 RCF for 1 min.
8) Register plate in the LIMS for runs on ABI 3700 machines.

## Sequencing Primers:

| Primer <br> Name | piggyBac <br> End | Sequence <br> $\mathbf{5 '}^{\prime}$ to 3' |
| :---: | :---: | :--- |
| pB-5SEQ | 5' end | 5'-CGC GCT ATT TAG AAA GAG AGA G-3' |
| pB-3SEQ | 3' end | 5'-CGA TAA AAC ACA TGC GTC AAT T-3' |

## Figures:

## piggyBac 5' end



## piggyBac 3' end:

HinP1I
$\begin{array}{lllll}10 & 20 & 30 & 40 & 50\end{array}$
GCGCGATAATATCTCTAATATTTTGCCAAATGAAGTGCCTGGTACATCAG CGCGCTATTATAGAGATTATAAAACGGTTTACTTCACGGACCATGTAGTC
$60 \quad 70 \quad 80 \quad 90 \quad 100$

ATGACAGTACTGAAGAGCCAGTAATGAAAAAACGTACTTACTGTACTTAC TACTGTCATGACTTCTCGGTCATTACTTTTTTGCATGAATGACATGAATG
$110 \quad 120 \quad 130 \quad 140 \quad 150$
TGCCCCTCTAAAATAAGGCGAAAGGCAAATGCATCGTGCAAAAAATGCAA ACGGGGAGATTTTATTCCGCTTTCCGTTTACGIAGCACGTTTTTTACGTT

3R2
$160170 \quad 180 \quad 190 \quad 200$
AAAAGTTATTTGTCGAGAGCATAATATTGATATGTGCCAAAGTTGTTTCT TTTTCAATAAACAGCTCTCGTATTATAACTATACACGGTTTCAACAAAGA

$210220230240 \quad 250$
GACTGACTAATAAGTATAATTTGTTTCTATTATGTATAAGTTAAGCTAAT CTGACTGATTATTCATATTAAACAAAGATAATACATATTCAATTCGATTA
$260 \quad 280 \quad 290 \quad 300$
TACTTATTTTATAATACAACATGACTGTTTTTAAAGTACAAAATAAGTTT ATGAATAAAATATTATGTTGTACTGACAAAAATTTCATGTTTTATTCAAA

310320330340
ATTTTTGTAAAAGAGAGAATGTTTAAAAGTTTTGTTACTTTATAGAAGAA TAAAAACATTTTCTCTCTTACAAATTTTCAAAACAATGAAATATCTTCTT
360
$370 \quad 380$
390
400

ATTTTGAGTTTTTGTTTTTTTTTAATAAATAAATAAACATAAATAAATTG TAAAACTCAAAAACAAAAAAAAATTATTTATTTATTTGTATTTATTTAAC

410420430440 450
TTTGTTGAATTTATTATTAGTATGTAAGTGTAAATATAATAAAACTTAAT AAACAACTTAAATAATAATCATACATTCACATTTATATTATTTTGAATTA


ATCTATTCAAATTAATAAATAAACCTCGATATACAGACCGATAAAACACA TAGATAAGTTTAATTATTTATTTGGAGCTATATGTCTGGCTATTTTGTGT


TGCGTCAATTTTACGCATGATTATCTTTAACGTACGTCACAATATGATTA ACGCAGTTAAAATGCGTACTAATAGAAATTGCATGCAGTGTTATACTAAT

