Supporting Online Material for

## Adaptive Evolution of Pelvic Reduction in Sticklebacks by Recurrent Deletion of a Pitx1 Enhancer

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## Supporting Online Material

## Material and Methods

Fish Collection and Husbandry. Sticklebacks were collected in minnow traps from locations described in table S1. Laboratory crosses were generated, as described (S1).

BAC Screening. BAC libraries, generated from a pelvic-complete stickleback population, Salmon River, BC, Canada (CHORI-213; Children's Hospital Oakland Research Institute, Oakland, CA, USA); and from a pelvic-reduced population, Paxton Lake (benthic form), BC, Canada (CHORI-215) (S2) were screened on high-density filters with radioactively-labeled oligonucleotide probes as described (S3-4) using sequences near Pitxl:
-43kb-OVa 5'-CTCCCTGGATGGCTGGGAAACTGA-3', -43kb-OVb 5'-
GGTCTCTGGTCCCGGCCTCCCTGG-3', +38kb-OVa 5'-
TTCCGTAGTGTAGTGGTTATCACG-3', $+38 \mathrm{~kb}-\mathrm{OVb} 5$ '-
TCGCGTGTTAGGCGAACGTGATAA-3',-51kb-Ova 5'-
GAGGAGTGGAAACGAAAGAGTGAG-3',-51kb-OVb 5'-
ATAACATCTCCAGCGGCTCACTCT-3'. Positive clones were secondarily screened with microsatellite markers $\operatorname{Stn} 446$, $\operatorname{Stn} 454$, $\operatorname{Stn} 474$ and end sequenced with T7 and SP6 primers on an ABI3730xl automatic sequencer (Applied Biosystems, Foster City, CA, USA). BAC Clones CH213-164F21, CH213-118G22, CH215-217P05 and CH215-196J14 (Genbank accession numbers: GU130433-4, GU130436-7) were sequenced to completion as described (S2).

Allele-Specific Expression. Allele-specific expression of Pitxl was analyzed using FRIL as the control parent. All FRIL individuals show robust expression of the pelvic spines and girdle. Wild-caught females from FRIL were crossed to males from a pelvic-complete marine LITC population and the pelvic-reduced freshwater PAXB population. The progeny were raised to stage 29/30 (S5), a stage when Pitxl is strongly expressed in both head and pelvic regions (S6S8). Larvae were anesthetized in $0.0003 \%$ tricaine (Ethyl 3- aminobenzoate methanesulfonate, Sigma, St. Louis, MO, USA). Heads and pelvic regions were dissected and placed individually in tubes. Tissue samples were immediately frozen and stored at $-80^{\circ} \mathrm{C}$. RNA was isolated from head and pelvis samples with the RNeasy Micro Kit (Qiagen, Valencia, CA, USA). The remainder of each larva was used for DNA isolation. We analyzed twenty larvae from each cross type. Allele specific RT-PCR was carried out with gene-specific primers, the forward primer was labeled with a 6-FAM fluorophore (Forward: 5’-CCCCTTCTCTTCAAGCTGAG-3'. Reverse: 5'-AGCTGCTGGCTGGTGAAG-3'). We distinguished the alleles using a six base-pair deletion in a GCG repeat, resulting in a poly-alanine polymophism in the poorly conserved N -terminal region of the Pitxl protein. The PCR product is approximately 250 bp from spliced RNA, depending on the length of the GCG repeat. Any residual DNA in the samples would result in an approximately 2.2 kb product that was not visible in our assays. RT-PCR reactions were done with the OneStep RT-PCR (Qiagen). Each $50 \mu \mathrm{l}$ reaction contained: $16 \mu \mathrm{l}$ water, $10 \mu \mathrm{l} 5 \mathrm{X}$ Qiagen PCR buffer, $10 \mu \mathrm{l}$ Q solution, $2 \mu \mathrm{l}$ dNTPs, $3 \mu \mathrm{l}$ each primer, $2 \mu \mathrm{l}$ OneStep RT-PCR enzyme mix, and $4 \mu$ RNA from pelvis samples or $2 \mu \mathrm{l}$ RNA from head samples plus an additional $2 \mu$ l of water. To ensure accurate quantification of peak intensity, $5 \mu$ l aliquots were
removed during the extension stage every 5 cycles between 25 and 40 cycles. Of each $5 \mu \mathrm{l}$ aliquot, $1 \mu \mathrm{l}$ was mixed with $10 \mu \mathrm{HiDi}$ formamide (Applied Biosystems) and $0.25 \mu \mathrm{l}$ of GeneScan 500 LIZ (Applied Biosystems) and analyzed with an ABI3730xl and GeneMapper v3.5 software (Applied Biosystems).
Reaction products were detectable in pelvis samples by cycle 30, robust at cycle 35 and off-scale by cycle 40 . Head samples reached saturation by cycle 35 and were analyzed at cycle 30 . For each sample, allele ratios were computed as follows: the peak height (arbitrary units of peak intensity) of the experimental allele (LITC, PAXB) was divided by the height of the control allele (FRIL).

Association Mapping. Thirty-six fluorescent-labeled markers (Genbank GF100639- GF100673, GF101813, table S2) were designed with Primer3 as described (S1, S9), with an average spacing of 8 kb across a 259 kb Pitx1 region derived from BAC clones CH213-164F21 (5') and CH213118G22 (3'). Two populations segregating pelvic phenotypes, (PAXB/PAXL and WALC/WALR) were screened for alleles associated with pelvic reduction. In each case, 48 pelvic-reduced individuals and 48 pelvic-complete individuals were genotyped (Paxton Lake: Stn431, Stn445-9, Stn452-4, Stn456, Stn458, Stn462, Stn464, Stn471-2, Stn474, Stn47-9, $\operatorname{Stn} 482$; Wallace Lake: $\operatorname{Stn} 336, \operatorname{Stn} 445, S \operatorname{tn} 448, \operatorname{Stn} 450-1, S \operatorname{tn} 453-7, S \operatorname{tn} 459-S \operatorname{tn} 470, S \operatorname{tn} 472-$ $\operatorname{Stn} 480$ ). Microsatellites were amplified and genotyped on an ABI3730xl sequencer with GeneMapper v3.7 software as described (S1). Differences in microsatellite allele frequency distributions were expressed in $\log (p-v a l u e s)$ between the pelvic-complete and pelvic-reduced fish with a modified $\chi^{2}$ test on a coalesced $2 \times 2$ contingency table (T4) as implemented by CLUMP (S10). The candidate region was delimited by haplotypes strongly associated with pelvic reduction, as determined by PHASE v2.1 (S11).

Transgenic Assays. Transgenic sticklebacks were generated by microinjection (S12) with the following modifications to the injection mount: a glass plate measuring $20 \times 15 \mathrm{~cm}$ was used as the injection platform instead of a microfabricated injection chamber. Freshly fertilized embryos at one-cell stage were placed, with the blastodisc facing the microinjection needle, in the indentations between teeth of a 6 " plaster saw blade (HILTI, Tulsa, OK, USA: \#00374342, 6 teeth per inch) with an artists' paintbrush.
Plasmids were co-injected with tol2 transposon mRNA as described (S12). We synthesized mature tol2 mRNA by in vitro transcription using the mMessage mMachine SP6 kit (Ambion, Austin, TX, USA). For enhancer assays using EGFP as the reporter, we used fertilized embryos from either MATA or LITC pelvic-complete populations. For the functional rescue assays, we used fertilized embryos from the BEPA pelvic-reduced population. All larvae were raised under standard aquarium conditions to Swarup St 29/30 (S5), when pelvic bud development is initiated, for phenotyping. Larvae were anesthetized in $0.0003 \% \mathrm{w} / \mathrm{v}$ tricaine (Ethyl 3-aminobenzoate methanesulfonate, Sigma). Microscopic observation for EGFP and mCherry expression was conducted with a MZFLIII fluorescent microscope (Leica Microsystems, Bannockburn, IL) with GFP2, GFP3 and DsRed filters. Some larvae were also grown to adulthood for skeletal visualization by Alizarin Red staining, as described (S1).
pTol2[drHsp70:eGFP] (pBHR) vector. The $d r H s p 70$ promoter from pHSP70-4-EGFP (S1314) was released by digestion with AvaI and cloned into the AgeI site of pBH-mcs-YFP. EGFP was PCR amplified from pHSP70-4-EGFP with primers 5'-
AAAATCGTCGACGGTCGCCACCATGGTGAGCAAGG-3' and 5'-
AAAATCGATCGATTAAGATACATTGATGAGTTTGG-3'. The EGFP was then cloned into pBH mcs YFP using digestion with SalI and ClaI, replacing YFP as the fluoroflore, yielding the pBHR vector. Potential enhancer sequences were then PCR amplified, NotI digested, and cloned into pBHR at the $P s p O M I$ site.

Transgenic rescue vector. Individual fragments matching Pitxl exons were PCR amplified with primers pairs 5'-
AAAACTGTCGACATGGAGCTAAATCTTTCGAGCAAACAAAACGTCATTTCCGGCGC
GGACGCAGGTGGGATGAATTCGGACC-3" (designated "primer 1") and 5"-
CTCGGACGCGGAGCCGCCAGAGAAGGAGC-3'; 5'-
AGCCGCCAGAGAAGGAGCGCGGCG-3' and 5'-
CGGCGGTTCTTGAACCACACCCGCACTCGGGCCTCG-3'; 5’-
GAGTGCGGGTGTGGTTCAAGAACCGCCG-3' and 5'-
AAAACTGATCAGCTGTTGTACTGGCACGCG-3' (primer 5). For the PCR, we used Phusion Polymerase (Finnzymes, Woburn, MA) and the following cycling conditions: $98^{\circ} \mathrm{C}$ for $5^{\prime} 30^{\prime \prime}$; $98^{\circ} \mathrm{C}$ for $30^{\prime \prime}$; ( $58^{\circ} \mathrm{C}$ for $20^{\prime \prime}-0.5^{\circ} \mathrm{C}$ per cycle; $72^{\circ} \mathrm{C}$ for $2^{\prime}$ ) x 4 times; $\left(98^{\circ} \mathrm{C}\right.$ for $30^{\prime} ; 58^{\circ} \mathrm{C}$ for $20^{\prime \prime}$; $72^{\circ} \mathrm{C}$ for $2^{\prime}$ ) x 29 times; $72^{\circ} \mathrm{C}$ for $7^{\prime} ; 4^{\circ} \mathrm{C}$ to hold. All fragments were amplified from pooled genomic DNA extracted from 4 individuals from Bear Paw Lake, AK. A second PCR amplification was performed to fuse the individual fragments into a Pitxl minigene with identical cycling conditions, but only with primers 1 and 5. The fused Pitxl minigene fragment was then cloned into pBHR using digestion with SalI and ClaI, replacing eGFP from the vector pBHR described above. With the exception of the signature sequence located in primer 1 , the amplified region is identical in BEPA and SALR populations. Injected individuals were genotyped with primers 5'-CGGGCATTTACTTTATGTTGCT-3' (primer Geno-F) and 5'-GTCCGAATTCATCCCACCTG-3' (primer Geno-R) to confirm incorporation of the transgene, with the same overall PCR cycling conditions as above, but with only 30 " for the elongation step.

Pel enhancers. Fragments containing the Pel-2.5kb ${ }^{\text {SALR }}$ enhancer were cloned with a three-step process with primers 5' -AAACTGGCGGCCGCCAGTTATTAAGACGGTTATTAATGT-3' and 5'-AAGCACTAGTGTCAGGGCCGGACCATCTAACTC-3' (cut with NotI and SpeI); 5'-AACTGTCTAGACACACAGGAGATCTGAGGC-3' and 5'-
AAAACTGAATTCTGACGCGGCGCTCCATCACCGAGCC-3' (cut with EcoRI and XbaI). Fragments amplified from BAC clone CH213-164F21, all with the PCR cycling conditions described above, were individually cloned into a pBluescript KS+ vector that has been modified with linkers 5’-TCGACCTCGAGGGGGGGCGGCCGCCCGGTACCCAGC-3' and 5’-GCTGGGTACCGGGCGGCCGCCCCCCCTCGAGGTCGA-3' to introduce an additional NotI site between ApaI and KpnI. Using the endogenous NsiI site within the insert itself, the released fragment bounded by EcoRI and NsiI was then cloned into the other construct linearized by NsiI and NotI site. This fused fragment was further released using NotI and cloned into the PspOMI
site in the pBHR vector for an enhancer assay. The Pel- $\Delta 2.5 \mathrm{~kb}^{\text {PAXB }}$ construct from PAXB (1868bp deletion) was directly amplified from genomic DNA with primers $5^{\prime}$ -AAACTGGCGGCCGCCAGTTATTAAGACGGTTATTAATGT-3' and 5'-AAACTGCGGCCGCTGACGCGGCGCTCCATCACCGAGCC-3' with NotI-digested overhangs for the pBHR vector. The Pel-501bp ${ }^{\text {SALR }}$ insert was amplified from BAC clone CH213-164F21 as well, with primers 5'-AACTGTCTAGACACACAGGAGATCTGAGGC-3' and 5'-AAGCACTAGTCCTCAAATCTGCAGCGTT-3' (cut with XbaI and SpeI), and cloned into the aforementioned modified pBlueScript KS+ vector linearized with SpeI. The resulting plasmid was then re-cut with SpeI, now only re-opening on the original SpeI site. The concatenated insert was then released by NotI digest and cloned into pBHR at the PspOMI site.

Alignment against distantly related teleosts. Sequences from the stickleback Pitx1 region (derived from an alignment of the 5' and the 3' SALR Pitxl BACs, (CH213-164F21 \& CH213118G22, aligned reference submitted as Genbank: GU130435) were aligned pairwise against tetraodon (TETRAODON 7, Apr 2003), fugu (FUGU 4.0, Apr 2006), medaka (assembly 200506, Jun 2005) and zebrafish sequences (Zv6, Mar 2006) using mLAGAN (S15) following repeat filtering with RepeatMasker (S16) with a custom stickleback-specific repeat library, or with default options for other teleost sequences. The alignments were then visualized using mVISTA (http://genome.lbl.gov/vista/index.shtml; S15)

Transcription factor binding sites. Predicted transcription factor binding sites in the Pel-501bp enhancer sequence (SALR) were identified using MatInspector (version 8.01, July 2009, Genomatix, Munich, Germany). Matrix Family Library Version 8.1 (June 2009) was used to identify elements belonging to the classes General Core Promoter Elements (parameters: 0.75 for the core score and matrix similarity setting "optimized") and Vertebrates (0.75/Optimized), with "family matches" option.

TwistFlex. Complete Pitxl SALR BAC sequence (see above) and individual chromosomes from the stickleback genome assembly were analyzed by TwistFlex (S17). The following parameters were used for all regions: window size $=50 \mathrm{bp}$; leap size $=1 \mathrm{bp}$; threshold value $=13.7^{\circ}$; normalization value $=10 \mathrm{~kb}$; discrete display value $=5 \mathrm{~kb}$; maximal distance between adjacent peaks in a cluster $=2 \mathrm{~kb}$; minimal number of peaks in a cluster $=3$.

SNP genotyping. Complete BAC sequences from SALR, PAXB and the initial stickleback genome assembly from a pelvic-reduced BEPA individual (Broad gasAcu1, Feb 2006), were pairwise aligned with mLAGAN (S15) following repeat filtering with RepeatMasker (S16) with a custom stickleback-specific repeat library. Single nucleotide polymorphisms (SNPs) were identified from these comparisons. In addition, a number of SNPs were ascertained by amplification and resequencing of portions of the Pitxl coding and intergenic regions from a worldwide panel of sticklebacks. 149 SNPs (table S3; Genbank: ss158145672-849) from the Pitxl region were genotyped on 399 individuals on the Illumina GoldenGate 1536 platform (Illumina, San Diego, CA, USA) according to manufacturer's protocols with the BeadStudio v3.0 (Illumina) software. The raw results were individually inspected with the BeadStudio v3.0 software to minimize clustering errors.

Signature of Selection. Since SNP ascertainment may bias towards overestimates of frequencies of common alleles and may cause spurious selective sweeps, only SNPs with at least $10 \%$ minor allele frequencies amongst the ancestral marine populations were included in our calculation for various signatures of selection. Population genetics statistics for $\theta \pi$ and Fay and Wu's H were calculated as described (S18-20). Fay and Wu's H calculation was based on a sliding window of 11 consecutive SNPs, similar to (S19). Therefore the resulting values should not be compared against those obtained from re-sequencing data, due to an inflation of segregating sites (S) from the use of concatenated SNP genotypes in the place of sequences.

## Supporting figures



Fig. S1. Fine mapping of pelvic reduction to Pitxl upstream region. (A) Recombinant animals from a pelvic-complete (JAMA) by pelvic-reduced (PAXB) cross (S1, S3) define a minimal region that is always homozygous for PAXB alleles in F2 progeny completely missing a pelvis. Symbols: - microsatellite markers, red marker represents $\operatorname{Stn} 336$, peak marker for pelvic QTL (S3); blue bars, homozygous PAXB genotypes; red bars: PAXB/JAMA heterozygous genotypes; pink bars: transition zone between known genotypes; grey bar, candidate non-recombinant interval (124kb). (B) Association mapping of dimorphic pelvic phenotypes in Paxton and Wallace Lakes. Symbols ( $\bullet \square$ ) indicate markers and phenotype-genotype association at different positions in 259 kb interval surrounding Pitx1. Solid blue marker: Stn463. Log p-values on y-axis are based on a modified $\chi^{2}$ test, T4 as implemented in CLUMP (S10). Dotted vertical lines delimit a strongly associated 23 kb candidate region that maps to the $5^{\prime}$ intergenic region of Pitx1, and was used for both enhancer studies and statistical analyses in Fig. 5C \& S9F. (C) Names and locations of genes in region. Arrows indicate transcription orientation and abbreviations correspond to nomenclature of human orthologs where possible.


## Pelvic Score

Fig. S2. Phenotypic distribution in Wallace Lake, AK. Pelvic scores were assessed according to (S21). Zero ( 0 ) is absent, pelvic size and complexity increases to the right, and 8 is complete. The 96 specimens were collected in Summer, 2005. Stickleback drawings are modified after (S22).


SALR 612
$\begin{array}{ll}\text { Tetraodon } & 349 \\ \text { Fugu } & 368\end{array}$ $\begin{array}{ll}\text { Fugu } & 668 \\ \text { Medaka } & 632\end{array}$

|  |  |
| :--- | :--- |
| SALR | 639 |
| Tetraodon | 395 |
| Fugu | 421 |
| Medaka | 737 |
| Zebrafish | 36 |
|  |  |
|  |  |
|  |  |
| SALR | 639 |
| Tetraodon | 395 |
| Fugu | 421 |
| Medaka | 847 |
| Zebrafish | 36 |



$\begin{array}{ll}\text { SALR } & 740 \\ \text { Tetraodon } & 782\end{array}$
$\begin{array}{ll}\text { Tetraodon } & 782 \\ \text { Fugu } & 891\end{array}$
$\begin{array}{ll}\text { Fugu } & 891 \\ \text { Medaka } & 141\end{array}$ Zebrafish 36


SALR 740






$\begin{array}{ll}\text { SALR } & 1765 \\ \text { Tetraodon } & 2190\end{array}$
Fugu 2395
$\begin{array}{ll}\text { Medaka } & 4591 \\ \text { Zehrafish } & 1112\end{array}$
Zebrafish

|  |  |
| :--- | :--- |
| SALR | 187 |
| Tetraodon | 2190 |
| Fugu | 239 |
| Medaka | 463 |
| Zebrafish | 1112 |


 ДGCATTAAGGTCTTTAAAGAAACCACGTATTCAGTCAGCTGTGAAGTCATTGTTAAATTCATCTTTGCATTCGTGGAAGACGCCACGTGTGAACAAACCTTCCTGGAAG

SALR 1939
Fugu 2408
Medaka 4797
Zebrafish 1123

| SALR | 1939 |
| :--- | :--- |
| Tetraodon | 2202 |
| Fugu | 2408 |
| Medaka | 4907 | Zebrafish 1123


|  |  |
| :--- | :--- |
| SALR | 193 |
| Tetraodon | 220 |
| Fugu | 240 |
| Medaka | 501 |
| Zebrafish | 112 |

[^0]




Fig. S3. Multiple sequence alignment between stickleback Pel-2.5kb ${ }^{\text {SALR }}$ sequence (Fig. S7) and the corresponding region from other teleost fish. Some sequences conserved between sticklebacks and zebrafish show sequence changes in pelvic-reduced pufferfish (fugu and tetraodon). The sequences conserved between distantly-related teleost species can not yet be aligned to most tetrapod genomes tested to date, including chicken, mouse, human, and various other mammals.


Fig. S4. Predicted binding sites of major developmental regulators located within the Pel-501bp (SALR) sequence. Predicted transcription factor binding sites are indicated by their names and bars above the sequence. The wider bars indicate sequences matching the position weight matrices as predicted by MatInspector (see methods). Strong bars indicate the core of the matrix. Greyed out sequences indicate overlap with the 488bp area shared between pelvic-reduced populations (see Fig. 4A, S7).


Fig. S5. Distribution of pelvic phenotypes in injected sticklebacks and uninjected control siblings. Transgenic animals were generated as described in Methods, and pelvic scores were assessed according to (S21).


Fig. S6. PCR validation of transgenesis. Primers Geno-F and Geno-R (see Transgenic Rescue Vector in methods) were used to amplify the fragment containing the transgenic Pitxl minigene. The transgene ( $\mathrm{Tg}+$ ) yields a product of $\sim 300 \mathrm{bp}$.


Fig. S7. Sequence of Pel enhancer and deletion end points in pelvic-reduced populations. (A) The sequence of the region located within the Pel-2.5kb ${ }^{\text {SALR }}$ is displayed in background grey color. Homopolymer tracks are indicated by black letters, with repeats, alternating purine/pyrimidine tracts, and microsatellites indicated in blue. The underlined blue sequence indicates a 48bp sub-telomeric repeat motif that is specific to threespine sticklebacks. Red brackets and numberings indicate cloned breakpoints, with the red letters underneath each breakpoint indicating micro-homologies. (4) indicates BEPA, with a CA micro-homology. (6) indicates HUMP. 77 indicates PAXB, with a CTG micro-homology. Shaded sequence indicates minimal region of 488 bp shared in multiple deletions. This region shows substantial overlap with the Pel-501bp ${ }^{\text {SALR }}$ fragment (black underlined sequence). (B) A model of a possible deletion event. A fragile site (indicated by wavy line) induces a spontaneous deletion via double strand break during replication fork stalling. To resolve breakage, re-ligation is attempted, with chewback at the deletion boundary, according to the non-homologous end joining (NHEJ) pathway. Short identical sequences at the junction can assist the annealing and ligation process, indicated here as CA overhangs. After repair, the deleted sequence shows a single instance of CA, whereas it appears at both mapped boundaries in the ancestral sequence.


Fig. S8. SNP allele frequencies in the Pitxl region by population. Each column corresponds to an individual SNP, and each row shows genotyping results collected from 12 individuals of a particular population. Only markers with at least $10 \%$ minor allele frequency in marine populations are included in the analysis. The positions of these SNPs are marked by lines above each column, indicating the physical location of the SNP with respect to the transcription start site of Pitxl ( 0 kb ). Each genotype tab is colored according to the ancestral-derived gradient, as indicated, with the most common allele among marine populations taken as the marine ancestral allele, and the alternate allele as the derived allele. Markers that are polymorphic within a population are shown with intermediate reddish tints. Failed genotypes are indicated by "-" dashes. Red lines and dashes indicate locations falling within the 23 kb candidate interval defined by association studies in Fig. S1B. The order of populations plotted is as follows: BIGR_3.63, TYNE_1, RABS, NVRO_1, KODK, GORT, GJÖG, LITC, BJDG for marine pelvic-complete populations. WALC, PAXL, PRIB, PRIL, WMSO, TYNE_8, OLNY, FTC, FRIL, FRIC, FLCK_12.8, BIGR_20.33 for freshwater pelvic-complete populations. ORPH, BOUL, SCAD, DOLO, [break], HUMP, BOOT, BEPA, WHAL, PAXB, MVRO, VIFI, FADA, WALR for freshwater pelvic-reduced populations. The small break between pelvic-reduced populations separates populations with and without detected deletions.


Fig. S9. Signatures of positive selection in Pel enhancer region of pelvic reduced fish. (A and B) Expanded view of Fig. 5A \& B. Fay and Wu's H and relative heterozygosity $(\theta \pi)$ statistics across the Pitxl region. Blue (freshwater pelvic-reduced) and green (freshwater pelvic-complete) data points and LOESS smoothed ( $\alpha=0.2$ ) line represent the phenotypic mean-averaged summary statistics calculated initially by stickleback populations. The Pel-containing regulatory region of Pitxl (grey candidate region from Fig. S1B) shows both negative H values, indicating an excess of derived alleles; and reduced heterozygosity in pelvic-reduced fish, consistent with positive selection (see main text). $\theta \pi$ values are plotted relative to the grouped marine mean (per SNP) to control for variation in ascertainment between SNPs. (C, D and E) Relative
heterozygosity $(\theta \pi)$, normalized to grouped marine mean value. Each panel represents the heterozygosity of the group indicated plotted along the Pitxl region. Curves correspond to LOESS regressions by population ( $\alpha=0.2$, split plots shown below main figure). Red dotted line indicates the marine mean value (normalized to 1). The lower dotted line in each panel indicates a relative heterozygosity of 0.25 . (F) Position of minimum heterozygosity relative to Pel enhancer, shown for individual freshwater and marine populations. Minimum heterozygosity colocalizes with the Pel region in most pelvic reduced populations, but not in freshwater or marine populations with a complete pelvis. *** indicates $P<0.001$.

## Signatures of selection displayed as individual population plots:

Freshwater, pelvic-reduced







Freshwater, pelvic-complete











## Supporting tables

Table S1. Stickleback populations used in this study.

| Population | Country | Code | Habitat | Class | Phenotype | Lat | Long | Year | Collectors | Pop \# in Figs. 4 and 5 | Collecting Ref. |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Bear Paw <br> Lake, AK | USA | BEPA | Lake | Freshwater | Reduced | 61.615 | -149.757 | 2006 | M.A. Bell | FW/R-4 | This study |
| Big River (Site 20.33), CA | USA | $\underset{3}{\text { BIGR_20.3 }}$ | River | Freshwater | Complete | 39.317 | -123.686 | 2007 | F.C. Jones \& C. Brown | FW/C-1 | This study |
| Big River (Site 3.63), CA | USA | BIGR_3.63 | Estuary | Marine | Complete | 39.304 | -123.780 | 2007 | F.C. Jones \& C. Brown | Mar-2 | This study |
| Bodega Bay, CA | USA | BDGB | Brackish water | Marine | Complete | 38.325 | -123.041 | 2005 | H. Zhang | Mar-1 | This study |
| Boot Lake, AK | USA | BOOT | Lake | Freshwater | Reduced | 61.718 | -150.130 | 2006 | M.A. Bell | FW/R-5 | This study |
| Boulton <br> Lake, BC | Canada | BOUL | Lake | Freshwater | Reduced, atypical | 53.783 | -132.098 | $\begin{gathered} 2003 \\ - \\ 2006 \end{gathered}$ | M.E. <br> Marks, B.R. Summers \& Y.F. Chan | FW/R-12 | This study |
| Dolomite <br> Lake, AK | USA | DOLO | Lake | Freshwater | Reduced, atypical | 60.721 | -151.136 | 2006 | M.A. Bell | FW/R-10 | This study |
| Fish Trap Creek, WA | USA | FTC | River | Freshwater | Complete | 48.931 | -122.487 | 2005 | C.T. Miller | FW/C-4 | (S24) |
| Flynn Creek (Site 128), CA | USA | FLCK_128 | River | Freshwater | Complete | 39.161 | -123.583 | 2007 | F.C. Jones \& C. Brown | FW/C-2 | This study |
| Friant Dam (Completeplated), CA | USA | FRIC | River | Freshwater | Complete | 36.980 | -119.731 | $\begin{gathered} 1994 \\ - \\ 2007 \end{gathered}$ | Kingsley Lab | FW/C-3 | (S25) |
| Friant Dam (Lowplated), CA | USA | FRIL | River | Freshwater | Complete | 36.980 | -119.731 | $\begin{gathered} 1994 \\ - \\ 2007 \end{gathered}$ | Kingsley Lab | FW/C-11 | (S25) |
| Gasterosteus | USA | WMSO | River | Freshwater | Complete | 34.435 | -118.198 | 2001 | C. Peichel | FW/C-10 | (S25) |


| williamsonii |  |  |  |  |  |  |  | \& | \& D. |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  |  |  |  |  |  | 2002 | Kingsley E. |  |  |
| Gjögur | Iceland | GJÖG | Fjord | Marine | Complete | 65.980 | -21.440 | 2003 | Elfarsdóttir $\& B$. Jónsson | Mar-3 | (S25) |
| Gorten <br> Sands | UK | GORT_4 | Tidal beach | Marine | Complete | 56.909 | -5.852 | 2001 | F.C. Jones \& C. Brown | Mar-4 | This study |
| Hump Lake, AK | USA | HUMP | Lake | Freshwater | Reduced | 60.768 | -151.170 | 2006 | M.A. Bell | FW/R-6 | This study |
| Kodiak <br> Island, AK | USA | KODK | Open sea | Marine | Complete | 56.967 | -151.350 | 2004 | M. Wilson (NOAA) | Mar-5 | This study |
| Little |  |  |  |  |  |  |  |  | C.T. Miller |  |  |
| Campbell <br> River, BC | Canada | LITC | Estuary | Marine | Complete | 49.018 | -122.779 | 2004 | \& Y.F. <br> Chan | Mar-6 | (S25) |
| Loch Fada | UK | FADA | Lake | Freshwater | Reduced | 57.613 | -7.211 | 2001 | C. Peichel \& D.M. Kingsley | FW/R-9 | (S25) |
| Loch Scadavay | UK | SCAD | Lake | Freshwater | Reduced | 57.589 | -7.225 | 2001 | C. Peichel \& D.M. Kingsley | FW/R-11 | (S25) |
| Matadero <br> Creek, CA | USA | MATA | River | Freshwater | Complete | 37.386 | -122.165 | $\begin{gathered} 2004 \\ - \\ 2008 \end{gathered}$ | Y.F. Chan \& F.C. Jones | FW/C-12 | This study |
| Morvoro <br> Lake, AK <br> Navarro | USA | MVRO | Lake | Freshwater | Reduced | 61.602 | -149.786 | 2006 | M.A. Bell | FW/R-3 | This study |
| River (Site <br> 1), CA | USA | NVRO_1 | Estuary | Marine | Complete | 39.193 | -123.761 | 2003 | Kingsley Lab | Mar-7 | (S25) |
| Olney Creek, CA | USA | OLNY | River | Freshwater | Complete | 40.528 | -122.383 | 2007 | B.R. <br> Summers | FW/C-6 | This study |
| Orphia Lake, AK | USA | ORPH | Lake | Freshwater | Reduced, atypical | 60.388 | -151.199 | 2006 | M.A. Bell | FW/R-13 | This study, also (S. 26) |
| Paxton Lake (Benthic), BC | Canada | PAXB | Lake | Freshwater | Reduced | 49.712 | -124.525 | 2003 | D. Schluter | FW/R-7 | (S1, S27) |


| Paxton Lake (Limnetic), BC | Canada | PAXL | Lake | Freshwater | Complete | 49.712 | -124.525 | 2003 | D. Schluter | FW/C-13 | (S27) |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Priest Lake (Benthic), BC | Canada | PRIB | Lake | Freshwater | Complete | 49.746 | -124.567 | 2003 | D. Schluter | FW/C-7 | (S27) |
| Priest Lake (Limnetic), BC | Canada | PRIL | Lake | Freshwater | Complete | 49.746 | -124.567 | 2003 | D. Schluter | FW/C-8 | (S27) |
| Rabbit <br> Slough, AK | USA | RABS | River | Marine | Complete | 61.537 | -149.166 | 2005 | M.A. Bell | Mar-8 | (S28) |
| River Tyne (Site 1) | UK | TYNE_1 | Tidal rock pools | Marine | Complete | 55.999 | -2.520 | 2003 | F.C. Jones \& C. Brown | Mar-9 | (S29) |
| River Tyne (Site 8) | UK | TYNE_8 | River | Freshwater | Complete | 55.943 | -2.785 | 2003 | F.C. Jones \& C. Brown | FW/C-9 | (S29) |
| Vifisstaðavat <br> n | Iceland | VIFI | Lake | Freshwater | Reduced | 64.080 | -21.873 | $\begin{gathered} 2002 \\ \& \\ 2004 \end{gathered}$ | E. <br> Elfarsdóttir \& B. Jónsson | FW/R-8 | (S29) |
| Wallace Lake (Complete), AK | USA | WALC | Lake | Freshwater | Complete | 61.573 | -149.576 | 2004 | M.A. Bell | FW/C-14 | This study |
| Wallace Lake (Reduced), AK* | USA | WALR | Lake | Freshwater | Reduced | 61.573 | -149.576 | 2004 | M.A. Bell | FW/R-1 | This study |
| Whale Lake, AK | USA | WHAL | Lake | Freshwater | Reduced | 61.542 | -149.751 | 2006 | M.A. Bell | FW/R-2 | This study |
| FW/R - Fres <br> FW/C - Fres Mar - Marin | water, water, pelvic | vic-reduc ic-comp mplete |  |  |  |  |  |  |  |  |  |

Table S2. Microsatellite Primer Sequences. Position from TSS — Position of the microsatellite marker relative to the Pitxl transcription start site (base pair 163184 in the 5' BAC clone CH213-164F21).

| Name | GenBank ID | Position from TSS | Sequence |
| :--- | :---: | :---: | :---: |
| Stn446-F | GF100639 | -162540 | GATGAACTTGCCATGAAAGG |
| Stn446-R | GF100639 | -162540 | CTTCCTCTGGAAACGACAGC |
| Stn447-F | GF100640 | -158901 | GAGAACCAACAGGCAGTATCG |
| Stn447-R | GF100640 | -158901 | CCACCGACAGTAAGTTAGACG |
| Stn448-F | GF100641 | -149303 | CGAAGCTCTCGCTGTTCC |
| Stn448-R | GF100641 | -149303 | CTCAGACTGCGTGGTAATGC |
| Stn449-F | GF100642 | -142358 | AATTGAGCTGGATGATGACG |
| Stn449-R | GF100642 | -142358 | TTTGATTCATCTCGGACTCG |
| Stn482-F | GF101813 | -140195 | TTTCTCAGAGGCATGTTTCC |
| Stn482-R | GF101813 | -140195 | GGCCAGAGACACTGAGTCC |
| Stn450-F | GF100643 | -131074 | GGTCCCATTTAAGTCCAATCC |
| Stn450-R | GF100643 | -131074 | GACAGAGACGTGGAAACACC |
| Stn451-F | GF100644 | -126002 | CCACTACATCCCTGTTTGAGC |
| Stn451-R | GF100644 | -126002 | ACGTGTGTCTTGCCTCTCC |
| Stn452-F | GF100645 | -122910 | ATCCTCCTCTTCCTCAGTCC |
| Stn452-R | GF100645 | -122910 | CTGGGCAGACGATATTTGC |
| Stn453-F | GF100646 | -118775 | AACCTGCTACATGTCGTCTCC |
| Stn453-R | GF100646 | -118775 | CTGACTGATGTCGGATAGATGC |
| Stn454-F | GF100647 | -108135 | TGGTCGAACATTTGGTTCC |
| Stn454-R | GF100647 | -108135 | CTACCATCACACCCAACAGG |
| Stn455-F | GF100648 | -82405 | CTGTTCTTTCGACACGTTGC |
| Stn455-R | GF100648 | -82405 | TGAATGAGGTCCTGTTACCC |
| Stn456-F | GF100649 | -66428 | CAGTTCAGAGACGACAATGAGG |
| Stn456-R | GF100649 | -66428 | CATCAGATGCAGAACAGACG |
| Stn445-F | GF089698 | -60520 | TCTTCCGCTCTGTGATTGG |
| Stn445-R | GF089698 | -60520 | CTTCACGCAACACGTTTACC |
| Stn457-F | GF100650 | -42385 | CCATCGTTAGATTCTCTTCTCC |
| Stn457-R | GF100650 | -42385 | CGGGTCCAGGACTCTAGG |
| Stn458-F | GF100651 | -42139 | CCCACATTGTGAAACTAACACC |
| Stn458-R | GF100651 | -42139 | CAGATAGTTGTGTTGCCTTTCG |
| Stn459-F | GF100652 | -39713 | TGCACACATTTCCTACATGC |
| Stn459-R | GF100652 | -39713 | ACAGGAAGCAGCAGAGTCG |
| Stn460-F | GF100653 | -39602 | ACGGGAAGGTCCCTAACG |
| Stn460-R | GF100653 | -39602 | AGCTGATAAGAGCTGAGAAAGC |
| Stn461-F | GF100654 | -38595 | TTCTACGTCGCCACTAAACG |
| Stn461-R | GF100654 | -38595 | CCGTGAAGTCTTCTCACAGC |
| Stn462-F | GF100655 | -37233 | TCAGCAGTGTTTGAGCTTCC |
| Stn462-R | GF100655 | -37233 | TGATTCACTAAACGCCATCG |
| Stn463-F | GF100656 | -35661 | AGACCTGCTCCTCCAGACC |
| Stn463-R | GF100656 | -35661 | CTTTCAAGTGTTGCCTGACG |
| Stn464-F | GF100657 | -30046 | CACATCTGCTGCACAAGC |
| Stn464-R | GF100657 | -30046 | TTCTTTCCTCCATCGATCC |
| Stn465-F | GF100658 | -23735 | GGAGAGGCTTTGATCTGAGG |


| Stn465-R | GF100658 | -23735 | CATCGTCAGTCAGAGATGTGG |
| :---: | :---: | :---: | :---: |
| Stn466-F | GF100659 | -22723 | GTGGGAACCTTCTCTGTGC |
| Stn466-R | GF100659 | -22723 | CCCTGTCCTTTCATCTCTGG |
| Stn467-F | GF100660 | -20477 | GAATATCCGCCTCGTTAGC |
| Stn467-R | GF100660 | -20477 | GGTTCTTTCTCCAACATCAGG |
| Stn468-F | GF100661 | -16787 | AACTTGAGAAGGACGCTTCG |
| Stn468-R | GF100661 | -16787 | GGTTAAACCTGCAGTCAAGC |
| Stn469-F | GF100662 | -16197 | CTGTCAATCACCGTTGTCG |
| Stn469-R | GF100662 | -16197 | ATGGGTGGAGCATTGACC |
| Stn470-F | GF100663 | 7567 | TCTGCACGTGTCTCTCACG |
| Stn470-R | GF100663 | 7567 | CCTGAGTCTCTTTCAGCATCC |
| Stn431-F | GF089697 | 8914 | ACCCAGAGGAGGAACATGG |
| Stn431-R | GF089697 | 8914 | AGCTTCACCTCCTAATTCTGC |
| Stn471-F | GF100664 | 15604 | TTCACTGGACGACGTTATGC |
| Stn471-R | GF100664 | 15604 | TCTCCGTGCTCTTCATCG |
| Stn472-F | GF100665 | 31503 | TGAAGGTCTGCGTCTGTCC |
| Stn472-R | GF100665 | 31503 | GCTCTGCAGATCACCAAGG |
| Stn473-F | GF100666 | 44744 | AGCATGTAGCATCTCACTATCG |
| Stn473-R | GF100666 | 44744 | GCATAGTGACTGGGAGCAGA |
| Stn474-F | GF100667 | 45612 | AGACGAGACGCAAATGTGG |
| Stn474-R | GF100667 | 45612 | AGGAAGTGACCTCACAGTCG |
| Stn475-F | GF100668 | 52160 | TCCTTATTGGTGAGTGTTCTCC |
| Stn475-R | GF100668 | 52160 | GGTGTGCAGAAATCCCAATT |
| Stn476-F | GF100669 | 58077 | ACTGTTCCCACCGAGAGC |
| Stn476-R | GF100669 | 58077 | GCACACACACACCAACACAC |
| Stn477-F | GF100670 | 60065 | GAATCCAGAGCCGTGAGC |
| Stn477-R | GF100670 | 60065 | TTCTGATCTGAGGAGCATCG |
| Stn478-F | GF100671 | 65042 | CATCATCAGCTCCGTACTTCC |
| Stn478-R | GF100671 | 65042 | TCTCGTCGGTCCGTTAGC |
| Stn479-F | GF100672 | 79311 | GTGTGGACGACACAATGC |
| Stn479-R | GF100672 | 79311 | CATCAAACGACAACAGAACC |
| Stn480-F | GF100673 | 103540 | TTTCAACCAACACACGTTCC |
| Stn480-R | GF100673 | 103540 | AATCCCTCCATTCAGACTGC |

Table S3. SNPs used for high density genotyping of the Pitxl region.

| SNP | Chr | Position | Alleles | "Anc" | $\begin{aligned} & 10 \% \\ & \text { MAF? } \end{aligned}$ | $\begin{aligned} & \text { Used } \\ & \text { in } \theta_{\pi} \text { ? } \end{aligned}$ | Pel | Sequence |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| ss158145849 | chriII | 6586435 | A/C | n.d. | N | Y | N/A | GATGAGCCG[A/C]AGGGCTTTT |
| ss158145672 | chriII | 6596376 | A/G | n.d. | N | Y | N/A | ACACAGACA[A/G]CAAAGGAAG |
| ss158145673 | chriII | 6599283 | A/G | n.d. | N | Y | N/A | ATGGTCATC[A/G]AATTTGGTA |
| ss158145674 | chriII | 6602429 | A/C | n.d. | N | Y | N/A | TAGCTGCAT[A/C]TATTCCCTC |
| ss158145675 | chriII | 6618228 | A/G | n.d. | N | Y | N/A | TTAAAATGC[A/G]CGACTTGTT |
| ss158145676 | chriII | 6625171 | A/C | n.d. | N | Y | N/A | CACGTGTTT[A/C]GAATATAGC |
| ss158145677 | chriII | 6630535 | A/T | n.d. | N | Y | N/A | GTCTACATA[A/T]GGGAGCCAT |
| ss158145678 | chriII | 6638976 | A/C | n.d. | N | Y | N/A | ATGCCTTCC[A/C]CTTGCCATT |
| ss158145679 | chriII | 6641414 | A/C | n.d. | N | Y | N/A | CTGAAACTA[A/C]AATTATAGC |
| ss158145680 | chriII | 6645878 | A/G | n.d. | N | Y | N/A | TAACAACAC[A/G]CGCAAATGT |
| ss158145681 | chriII | 6650410 | A/C | n.d. | N | Y | N/A | TGTTATCTA[A/C]CATACGTGT |
| ss158145682 | chriII | 6651054 | C/G | n.d. | N | Y | N/A | CCCTGAAAT[C/G]TCCCTGTTT |
| ss158145683 | chriII | 6661551 | A/G | n.d. | N | Y | N/A | TAATTTTGC[A/G]TGTCGTTTT |
| ss158145684 | chriII | 6675980 | A/G | n.d. | N | Y | N/A | ACCCAGTGC[A/G]ACTTTCATG |
| ss158145685 | chrVII-Pitx | -106860 | A/G | G | Y | N | N | CGTGTAGTC[A/G]GCTGGAGTC |
| ss158145686 | chrVII-Pitx | -104344 | C/G | N/A | N | N | N | ATACGCTAA[C/G]TGTACCTTT |
| ss158145687 | chrVII-Pitx | -103213 | A/C | N/A | N | N | N | TTGCCAATT[A/C]CTAAGCTAA |
| ss158145688 | chrVII-Pitx | -100714 | A/T | N/A | N | N | N | TTTGTCCTC[A/T]GCTCATACT |
| ss158145689 | chrVII-Pitx | -95697 | C/G | N/A | N | Y | N | TGAGCTCCA[C/G]CAGTGCTCC |
| ss158145690 | chrVII-Pitx 1 | -87996 | A/G | N/A | N | N | N | AAAGACATT[A/G]TAACTTCTG |
| ss158145691 | chrVII-Pitx 1 | -85185 | C/G | G | Y | N | N | GATCTCTGT[C/G]TGTGGCTCA |
| ss158145692 | chrVII-Pitx | -83383 | A/G | N/A | N | N | N | GAAATTCAC[A/G]CCAACGTTT |
| ss158145693 | chrVII-Pitx | -76733 | A/T | A | Y | N | N | GTAAATATT[A/T]AACACAAGT |
| ss158145694 | chrVII-Pitx | -74467 | A/C | C | Y | Y | N | ATATAATTA[A/C]TTTGACTCC |
| ss158145695 | chrVII-Pitx | -72879 | A/C | A | Y | N | N | GCTAGCTGC[A/C]GCTCGCTTT |
| ss158145696 | chrVII-Pitx | -69944 | A/C | N/A | N | Y | N | TGGCATGCG[A/C]GCACACACA |
| ss158145697 | chrVII-Pitx 1 | -67201 | A/G | G | Y | Y | N | CAGGAGACA[A/G]AAGGTCTGT |
| ss158145698 | chrVII-Pitx | -65636 | A/G | N/A | N | N | N | AAAGCAGAC[A/G]GACGTTCAC |


| ss158145699 | chrVII-Pitxl | -64004 | A/G | G | Y | N | N | ACATTAAAG[A/G]TGGATTCAT |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| ss158145700 | chrVII-Pitxl | -57072 | C/G | N/A | N | N | N | GCATCAGGA[C/G]CTCAAACCA |
| ss158145701 | chrVII-Pitx | -52419 | A/G | G | Y | N | N | TAATCGATT[A/G]GTTGAACAA |
| ss158145702 | chrVII-Pitx | -49974 | A/C | C | Y | N | N | TCTAACGGG[A/C]CTTTTTGTT |
| ss158145703 | chrVII-Pitx | -48246 | A/C | C | Y | N | N | AATCAATAA[A/C]GTACGTTTT |
| ss158145704 | chrVII-Pitxl | -42886 | A/G | N/A | N | N | N | GGCGAGGAA[A/G]ACGCATTTA |
| ss158145705 | chrVII-Pitx | -38724 | A/G | A | Y | N | N | GTCTTTATG[A/G]GGAGGAAGT |
| ss158145706 | chrVII-Pitx 1 | -38648 | A/G | N/A | N | N | N | GGATGAGGA[A/G]ATGAGATAT |
| ss158145707 | chrVII-Pitx | -38322 | C/G | N/A | N | N | N | ACATTTACA[C/G]GTATACTGT |
| ss158145708 | chrVII-Pitx 1 | -38056 | A/G | A | Y | Y | N | CTCAATTAA[A/G]CCTGGACGT |
| ss158145709 | chrVII-Pitx 1 | -37729 | A/G | A | Y | N | Y | CAACTAGAG[A/G]GCTCAAAAA |
| ss158145710 | chrVII-Pitx 1 | -37554 | A/C | N/A | N | N | Y | ACACCCTCC[A/C]GGAGATATT |
| ss158145711 | chrVII-Pitxl | -37401 | A/C | A | Y | N | Y | GCGCTAAAC[A/C]AACGTACAG |
| ss158145712 | chrVII-Pitx | -36628 | A/T | A | Y | Y | Y | CTTGACTTG[A/T]GGACGACAG |
| ss158145713 | chrVII-Pitxl | -36045 | A/G | N/A | N | N | Y | GGCGCTGGC[A/G]AGCGATGAG |
| ss158145714 | chrVII-Pitx | -34903 | C/G | N/A | N | N | Y | GAGTGAAGC[C/G]CCACAGAAA |
| ss158145715 | chrVII-Pitx | -34497 | A/G | A | Y | Y | Y | AACCGGTCC[A/G]TTTGCAGCT |
| ss158145716 | chrVII-Pitx 1 | -33286 | $\mathrm{A} / \mathrm{T}$ | N/A | N | N | Y | AGAGTTCTA[A/T]AGTGGTCGC |
| ss158145717 | chrVII-Pitx 1 | -33268 | A/T | N/A | N | N | Y | TGCATTAGG[A/T]CCTTGTGCA |
| ss158145718 | chrVII-Pitx 1 | -32530 | A/G | N/A | N | N | Y | TCTGGGGAC[A/G]GCCTGTCTC |
| ss158145719 | chrVII-Pitx 1 | -32520 | A/G | A | Y | N | Y | GTCCCCAGA[A/G]GGAGGTCTG |
| ss158145720 | chrVII-Pitx 1 | -31905 | A/G | G | Y | Y | Y | CAAATCCTC[A/G]ACGGAGGCT |
| ss158145721 | chrVII-Pitx 1 | -31771 | A/G | N/A | N | N | Y | ATGTCATTT[A/G]TCTAATTGT |
| ss158145722 | chrVII-Pitx 1 | -31460 | A/C | N/A | N | N | Y | GGATGAAAT[A/C]AAACAATTA |
| ss158145723 | chrVII-Pitx | -31251 | A/G | N/A | N | N | Y | CCTTAAAGA[A/G]ACACTGAAC |
| ss158145724 | chrVII-Pitx | -31238 | A/C | C | Y | N | N | ATAATAATC[A/C]TTTGTTCAG |
| ss158145725 | chrVII-Pitx | -30351 | A/C | N/A | N | N | N | TGATCGAAT[A/C]CCACGTGAT |
| ss158145726 | chrVII-Pitx 1 | -30297 | A/T | T | Y | N | N | CCAAAAATT[A/T]GTTTAATCG |
| ss158145727 | chrVII-Pitx 1 | -30172 | C/G | N/A | N | N | N | TTTTAAAGT[C/G]TGAAAACAT |
| ss158145728 | chrVII-Pitx 1 | -29968 | A/G | N/A | N | N | N | AACCTAATC[A/G]TTTAAAACG |
| ss158145729 | chrVII-Pitx 1 | -29919 | $\mathrm{A} / \mathrm{T}$ | N/A | N | N | N | TGTTTAAAA[A/T]AAAACGTTT |


| ss158145730 | chrVII-Pitx 1 | -29900 | A/T | T | Y | N | N | AGAGAATTT[A/T]AAAAGCAAA |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| ss158145731 | chrVII-Pitx 1 | -29796 | C/G | G | Y | N | N | TGTTTCTAT[C/G]CATGTTTAA |
| ss158145732 | chrVII-Pitx ${ }^{\text {l }}$ | -29795 | C/G | N/A | N | N | N | TTTAAACAT[C/G]GATAGAAAC |
| ss158145733 | chrVII-Pitx ${ }^{\text {l }}$ | -29755 | A/G | G | Y | N | N | AAACCATCT[A/G]AAGAATTTT |
| ss158145734 | chrVII-Pitx ${ }^{\text {l }}$ | -28795 | A/G | N/A | N | N | N | GAATATAAT[A/G]AACGTGTTG |
| ss158145735 | chrVII-Pitx 1 | -28703 | A/C | N/A | N | N | N | TATACAATA[A/C]ATTGTAAAT |
| ss158145736 | chrVII-Pitx ${ }^{\text {l }}$ | -28665 | A/G | A | Y | N | N | GTACATAAT[A/G]CTTACACAT |
| ss158145737 | chrVII-Pitx 1 | -28652 | A/C | C | Y | N | N | ATGTACACA[A/C]TTTCCATTT |
| ss158145738 | chrVII-Pitx 1 | -26659 | A/C | A | Y | Y | N | CAATTATGA[A/C]TTCCCTGAT |
| ss158145739 | chrVII-Pitx 1 | -24803 | A/G | N/A | N | N | N | CTGACTGAC[A/G]ATGTATACG |
| ss158145740 | chrVII-Pitx ${ }^{\text {l }}$ | -24377 | A/G | N/A | N | N | N | ACCAGCACC[A/G]CGGCCCTGA |
| ss158145741 | chrVII-Pitx ${ }^{\text {l }}$ | -24237 | C/G | N/A | N | N | N | ACAACATTT[C/G]TGACAATTT |
| ss158145742 | chrVII-Pitx 1 | -24124 | A/C | N/A | N | N | N | TTGGTCAGA[A/C]GCACAAACG |
| ss158145743 | chrVII-Pitx ${ }^{\text {l }}$ | -23975 | A/G | N/A | N | N | N | TGGAGGCCG[A/G]CAGGTTGAT |
| ss158145744 | chrVII-Pitx ${ }^{\text {l }}$ | -23894 | A/G | N/A | N | N | N | CTCTGGAAA[A/G]GATTAACTA |
| ss158145745 | chrVII-Pitx 1 | -23372 | A/C | C | Y | N | N | TGTAATTAA[A/C]CAGTTCCGT |
| ss158145746 | chrVII-Pitx 1 | -23320 | A/G | N/A | N | N | N | ATTAAAACC[A/G]GCACATGCA |
| ss158145747 | chrVII-Pitx 1 | -22821 | A/T | N/A | N | N | N | TATCAGATT[A/T]TGCTCTCCA |
| ss158145748 | chrVII-Pitx | -22780 | A/C | A | Y | N | N | CATCGTACG[A/C]TCAATCTTA |
| ss158145749 | chrVII-Pitx ${ }^{\text {l }}$ | -22662 | A/G | G | Y | N | N | AGCTAACAC[A/G]GTGACACAC |
| ss158145750 | chrVII-Pitx 1 | -21285 | A/C | N/A | N | N | N | ATGAATATC[A/C]GCCTCGTTA |
| ss158145751 | chrVII-Pitx ${ }^{\text {l }}$ | -19464 | A/G | N/A | N | N | N | GTtTATGGA[A/G]CAAATCTCT |
| ss158145752 | chrVII-Pitx ${ }^{\text {l }}$ | -18485 | A/G | G | Y | Y | N | AATCATTCC[A/G]CGTGCCGGA |
| ss158145753 | chrVII-Pitx 1 | -17958 | A/G | A | Y | N | N | AATGACAGT[A/G]ACAGTGACC |
| ss158145754 | chrVII-Pitx ${ }^{\text {l }}$ | -17357 | A/G | G | Y | N | N | CCAGTTAAA[A/G]TGTGAATAA |
| ss158145755 | chrVII-Pitx 1 | -17014 | A/G | G | Y | Y | N | CACGACAAC[A/G]GTGATTGAC |
| ss158145756 | chrVII-Pitx ${ }^{\text {l }}$ | -16878 | A/G | G | Y | N | N | GGATTAGAA[A/G]CTCATGGAC |
| ss158145757 | chrVII-Pitx 1 | -16661 | A/G | A | Y | N | N | GAGTTAAAG[A/G]CCTCCAAAG |
| ss158145758 | chrVII-Pitx 1 | -15801 | A/G | N/A | N | N | N | TGGTGTCCC[A/G]GGTGAGCTG |
| ss158145759 | chrVII-Pitx 1 | -15479 | C/G | C | Y | N | N | GACCGGCTT[C/G]AGACGCCTG |
| ss158145760 | chrVII-Pitx ${ }^{\text {I }}$ | -15044 | A/G | ? | Y | N | N | GTGCAGAGA[A/G]TCGTCTCCA |


| ss158145761 | chrVII-PitxI | -13498 | A/G | A | Y | N | N | GAGTTCAGC[A/G]CGTTACGCT |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| ss158145762 | chrVII-Pitx 1 | -12328 | A/T | T | Y | N | N | AAGCACAGA[A/T]CTGTGAGCT |
| ss158145763 | chrVII-Pitx ${ }^{\text {l }}$ | -12124 | A/C | N/A | N | N | N | AGGACCGAA[A/C]AATGCTGCT |
| ss158145764 | chrVII-Pitx ${ }^{\text {l }}$ | -12056 | A/G | G | Y | N | N | TACACCTGC[A/G]GCGAGGAAC |
| ss158145765 | chrVII-Pitx ${ }^{\text {I }}$ | -11709 | A/G | N/A | N | N | N | GCTTCAATA[A/G]ATAAACTGT |
| ss158145766 | chrVII-Pitx 1 | -10768 | A/G | G | Y | N | N | GAGCTTCCT[A/G]CCCGCCGCC |
| ss158145767 | chrVII-Pitx 1 | -8984 | A/C | C | Y | N | N | TTCACGTGA[A/C]GTTTATTAG |
| ss158145768 | chrVII-Pitx 1 | -7742 | C/G | C | Y | N | N | TTCCAATCA[C/G]AGCATGTTA |
| ss158145769 | chrVII-Pitx 1 | -5241 | A/C | N/A | N | N | N | AGACGGAGA[A/C]GATCGGCCG |
| ss158145770 | chrVII-Pitx ${ }^{\text {l }}$ | 767 | A/T | T | Y | N | N | TGTTTTATT[A/T]ATTGGACCG |
| ss158145771 | chrVII-Pitx 1 | 2203 | A/C | A | Y | N | N | GAGTTAACA[A/C]AGTGACTGC |
| ss158145772 | chrVII-Pitx 1 | 4435 | A/G | G | Y | N | N | ACTGTCCAC[A/G]CGGACGTGA |
| ss158145773 | chrVII-Pitx ${ }^{\text {l }}$ | 7632 | A/C | N/A | N | N | N | CTTTTATGT[A/C]ATTGGGGTC |
| ss158145774 | chrVII-Pitx ${ }^{\text {l }}$ | 9336 | A/C | C | Y | N | N | TGTGGTTCA[A/C]GGTTCCATC |
| ss158145775 | chrVII-Pitx ${ }^{\text {l }}$ | 11255 | A/G | G | Y | N | N | CACCCTCTC[A/G]TGTTACAAG |
| ss158145776 | chrVII-Pitx 1 | 19740 | A/G | A | Y | N | N | TAAAAAAAC[A/G]TGACCTCAC |
| ss158145777 | chrVII-Pitx ${ }^{\text {l }}$ | 21779 | A/G | ? | Y | N | N | AGTACATCG[A/G]TGCAGGTGA |
| ss158145778 | chrVII-Pitx 1 | 25121 | A/G | G | Y | N | N | AAACCGGTA[A/G]GTAGTGTCC |
| ss158145779 | chrVII-Pitx 1 | 26504 | A/T | A | Y | N | N | AAATAGTTT[A/T]CGGTCCACG |
| ss158145780 | chrVII-Pitx 1 | 29448 | A/C | N/A | N | N | N | ATCTGTGAG[A/C]CAAAGAAAA |
| ss158145781 | chrVII-Pitx ${ }^{\text {l }}$ | 32012 | C/G | N/A | N | N | N | CAAGATCAA[C/G]CGCACCAGT |
| ss158145782 | chrVII-Pitx 1 | 36350 | C/G | C | Y | Y | N | TGATCTATA[C/G]CCGGTAATC |
| ss158145783 | chrVII-Pitx 1 | 40204 | A/G | A | Y | N | N | TTCATCCTC[A/G]TCTTCCAAA |
| ss158145784 | chrVII-Pitx ${ }^{\text {l }}$ | 42215 | A/G | G | Y | Y | N | TGTTCTATC[A/G]ATGAATGAA |
| ss158145785 | chrVII-Pitx ${ }^{\text {l }}$ | 46864 | A/G | G | Y | N | N | ACTTTAGGA[A/G]GAGATTGTT |
| ss158145786 | chrVII-Pitx 1 | 51203 | A/G | A | Y | N | N | TTGGATCAC[A/G]TATGCGGTT |
| ss158145787 | chrVII-Pitx 1 | 58571 | A/T | N/A | N | N | N | ACATTAGTA[A/T]CCAGTAAGC |
| ss158145788 | chrVII-Pitx 1 | 59124 | A/G | N/A | N | N | N | GTTGGATTG[A/G]AGATAAAAT |
| ss158145789 | chrVII-Pitx | 61420 | A/C | C | Y | N | N | CTTGAGCTC[A/C]TGAAAGGCT |
| ss158145790 | chrVII-Pitx 1 | 65085 | C/G | N/A | N | N | N | TTTTAAATC[C/G]CTGGCTGAT |
| ss158145791 | chrVII-Pitx 1 | 66469 | A/G | N/A | N | N | N | TGTGGAACT[A/G]TCGGTGGAA |


| ss158145792 | chrVII-Pitxl | 71346 | A/G | N/A | N | N | N | ACGCCTCGT[A/G]AGAGAAGGC |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| ss158145793 | chrVII-Pitx | 74302 | A/G | N/A | N | N | N | TATTTATCT[A/G]CAGCAGATA |
| ss158145794 | chrVII-Pitx 1 | 84162 | A/G | N/A | N | N | N | GAGTTCATG[A/G]GTTTAGTTT |
| ss158145795 | chrVII-Pitx | 86984 | A/G | G | Y | Y | N | CTGAATATT[A/G]CTCATTATT |
| ss158145796 | chrVII-Pitx 1 | 89813 | A/G | G | Y | N | N | TTTCACGCG[A/G]CATCAGAAT |
| ss158145797 | chrVII-Pitx 1 | 91956 | A/T | A | Y | N | N | TCTGTGGAG[A/T]CCAGTTTTC |
| ss158145798 | chrVII-Pitx 1 | 97195 | A/C | N/A | N | N | N | AAAGAACAT[A/C]ATGTCGTGG |
| ss158145799 | chrVII-Pitx 1 | 98750 | C/G | G | Y | Y | N | AGTACATTA[C/G]AAGATTCTT |
| ss158145800 | chrVII-Pitx 1 | 101274 | A/C | C | Y | N | N | ACCTGCAGG[A/C]GGAGAGACA |
| ss158145801 | chrVII-Pitx 1 | 104731 | A/T | N/A | N | N | N | AACAATAAA[A/T]TCACCCCTA |
| ss158145802 | chrVII-Pitx | 106793 | A/G | A | Y | Y | N | AAAGCCGTT[A/G]TAGTCTTTT |
| ss158145803 | chrVII-Pitx | 109545 | A/T | A | Y | N | N | ACAATGTGC[A/T]GCTCAGGTT |
| ss158145804 | chrVII-Pitx 1 | 113259 | A/G | N/A | N | N | N | AAACTTCTA[A/G]GTTGTAAGA |
| ss158145805 | chrVII-Pitx 1 | 116974 | A/C | C | Y | Y | N | ACCAAGATC[A/C]GAGGTTCAA |
| ss158145806 | chrVII-Pitx 1 | 120540 | A/C | N/A | N | N | N | TCCTGGTTC[A/C]GTTAAAAAG |
| ss158145807 | chrVII-Pitx 1 | 121789 | A/G | N/A | N | N | N | ACTGGAAAT[A/G]AAACGCTGC |
| ss158145808 | chrVII-Pitx | 124456 | C/G | N/A | N | N | N | TTGAGTTTG[C/G]CGATTTTGG |
| ss158145809 | chrVII-Pitx | 126604 | A/G | N/A | N | N | N | CCTTTAGAC[A/G]TTTATTCAG |
| ss158145810 | chrVII-Pitx | 129893 | A/G | G | Y | N | N | CCTGTGAGG[A/G]AACGTTTTA |
| ss158145811 | chrVII-Pitx 1 | 132309 | A/G | N/A | N | N | N | TTTAACGTC[A/G]TTGATGTGA |
| ss158145812 | chrVII-Pitx | 134152 | A/T | N/A | N | N | N | CCTTTGCAT[A/T]TGGCTGTTA |
| ss158145813 | chrVII-Pitx | 141805 | A/C | C | Y | Y | N | CTCATTCTA[A/C]GATTACCAC |
| ss158145814 | chrVII-Pitx | 145217 | A/C | C | Y | Y | N | GTCTTGGTG[A/C]TAAATGAAT |
| ss158145815 | chrVII-Pitx | 148679 | A/G | ? | Y | Y | N | GTGTTTGGT[A/G]TTACTGCCG |
| ss158145816 | chrVII-Pitx | 149112 | A/C | C | Y | Y | N | CCTCCGGTG[A/C]TGAAGTCGC |
| ss158145817 | chrVII-Pitx | 152310 | A/G | N/A | N | Y | N | ACTGTCAGG[A/G]CGCTTATCT |
| ss158145818 | chrVII-Pitx | 154762 | A/G | A | Y | N | N | TCGTGTTCC[A/G]CCATCTTTA |
| ss158145819 | chrVII-Pitx | 163377 | A/G | G | Y | Y | N | TGAGAAGTC[A/G]GTCCAGATG |
| ss158145820 | chrVII-Pitx 1 | 165511 | A/G | N/A | N | Y | N | TTAACAAGT[A/G]TAAAATTGC |
| ss158145821 | chrVII-Pitx 1 | 166416 | A/G | N/A | N | Y | N | TTCTCGTTT[A/G]TTTACGCTA |
| ss158145822 | chrVII-Pitx 1 | 168923 | C/G | N/A | N | N | N | AGCATGAAA[C/G]CAGGATACG |


| ss158145823 | chrVII-Pitxl | 172127 | A/C | N/A | N | Y | N | TGTCCAACA[A/C]ATGGATGTA |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| ss158145824 | chrVII-Pitxl | 177714 | A/G | G | Y | Y | N | CTCTCATAC[A/G]CGCTGCAGA |
| ss158145825 | chrVII-Pitx 1 | 179019 | A/G | G | Y | Y | N | AGCAAAACA[A/G]CACCAAATA |
| ss158145826 | chrVII-Pitxl | 183425 | A/C | A | Y | Y | N | TGCTGTTTG[A/C]ATTTAAAAT |
| ss158145827 | chrVII-Pitxl | 184436 | C/G | N/A | N | Y | N | ATTTACAAT[C/G]AGACAAGAA |
| ss158145828 | chrVII-Pitxl | 186351 | A/G | N/A | N | N | N | AATCATCAC[A/G]TGAAGATAA |
| ss158145829 | chrVII-Pitx 1 | 189178 | A/G | G | Y | Y | N | AGAATATAA[A/G]GAAAAACAC |
| ss158145830 | chrVII-Pitx 1 | 196364 | A/G | A | Y | Y | N | TGTTGGCAA[A/G]CCAAACTTT |
| ss158145831 | chrVII-Pitxl | 204135 | A/C | N/A | N | N | N | ATACTTTAC[A/C]GTTACATGC |
| ss158145832 | chrVII-Pitxl | 206784 | A/C | A | Y | Y | N | AGAAGACTC[A/C]CAACAAGCT |
| ss158145833 | chrVII-Pitxl | 214543 | A/C | A | Y | Y | N | CTGAACATA[A/C]TATATGTGT |
| ss158145834 | chrXI | 1669455 | A/G | n.d. | N | Y | N/A | CATTTGGCT[A/G]AATTTGTGA |
| ss158145835 | chrXI | 1675661 | A/C | n.d. | N | Y | N/A | GCTTTCAAA[A/C]GTGGAACTG |
| ss158145836 | chrXI | 1683557 | A/C | n.d. | N | Y | N/A | CTATATATT[A/C]CATCCAACC |
| ss158145837 | chrXI | 1699906 | $\mathrm{A} / \mathrm{T}$ | n.d. | N | Y | N/A | CGTATTGTA[A/T]AGTGGGTGG |
| ss158145838 | chrXI | 1705579 | A/C | n.d. | N | Y | N/A | CATAGAGAG[A/C]GCGGGCTCT |
| ss158145839 | chrXI | 1710629 | A/G | n.d. | N | Y | N/A | TTCACCGCA[A/G]GACCACATT |
| ss158145840 | chrXI | 1712239 | A/G | n.d. | N | Y | N/A | TTTGGTTGC[A/G]AGCCATGAA |
| ss158145841 | chrXI | 1713968 | A/T | n.d. | N | Y | N/A | GTACAAGGA[A/T]GTAAAAGAG |
| ss158145842 | chrXI | 1725724 | A/C | n.d. | N | Y | N/A | CAGAGAAAA[A/C]ATTTGGGTA |
| ss158145843 | chrXI | 1727375 | A/G | n.d. | N | Y | N/A | TGACTCTGT[A/G]TCTTTGGAG |
| ss158145844 | chrXI | 1739125 | A/G | n.d. | N | Y | N/A | TGATGTGTT[A/G]CTATTATTT |
| ss158145845 | chrXI | 1744258 | A/C | n.d. | N | Y | N/A | AAACTGAAT[A/C]AGAAAAACA |
| ss158145846 | chrXI | 1780060 | A/T | n.d. | N | Y | N/A | TTTTACATG[A/T]GTCATTTTC |
| ss158145847 | chrXI | 1828565 | C/G | n.d. | N | Y | N/A | AAATATTTG[C/G]CGAGGCATA |
| ss158145848 | chrXI | 1829666 | A/C | n.d. | N | Y | N/A | GGTATGAAT[A/C]ATATTTTTC |
| ss158145849 | chrXI | 1831726 | A/G | n.d. | N | Y | N/A | TTTTTTCAG[A/G]TTATTATGG |

Anc - Ancestral allele as determined by most frequent allele in Pacific Marine populations.
$10 \%$ MAF - At least $10 \%$ minor allele frequency. "Y" indicates that the SNP was used in population genetics analyses Used in $\theta \pi$ - SNPs marked with " $Y$ " indicate that they have similar ascertainment and were used in the quantitative $\theta \pi$ analysis Pel - "SNPs" marked with " Y " indicate that were indel markers and occur within the candidate region containing Pel

## n.d. - Not determined

Position column refers to base pair in stickleback genome assembly (chrIII and chrXI), or relative to the TSS of Pitxl.

Table S4. Statistical analysis at SNP ss158145716. Grouping genotypes by population and then by the indicated grouping methods, the mean frequency of successful genotypes was compared between groups using two-tailed t-tests.

Group Statistics

|  | Grouping methods | N | Mean | Std. <br> Deviation | Std. <br> Error <br> Mean |
| :---: | :---: | :---: | :---: | :---: | :---: |
| $\begin{aligned} & \hline \text { \% Successful } \\ & \text { ss158145716 } \end{aligned}$ | Pelvic-reduced (All) | 13 | 0.3526 | 0.44767 | 0.12416 |
|  | Pelvic-complete (Mar/FW) | 21 | 0.9782 | 0.06126 | 0.01337 |

Equal variances not assumed, $\mathrm{df}=12.279, P<0.0004$

| \% Successful | Pelvic-reduced (Deletion-only) | 9 | 0.0741 | 0.12805 | 0.04268 |
| :--- | :--- | ---: | ---: | ---: | :--- |
| ss158145716 | All others | 25 | 0.9783 | 0.05784 | 0.01157 |

Equal variances not assumed, $\mathrm{df}=9.202, P<1 \times 10^{-8}$

| \% Successful | Pelvic-reduced (Deletion-only) | 9 | 0.0741 | 0.12805 | 0.04268 |
| :--- | :--- | ---: | :--- | :--- | :--- |
| ss158145716 | Pelvic-complete (Mar/FW) | 21 | 0.9782 | 0.06126 | 0.01337 |

Equal variances not assumed, $\mathrm{df}=9.610, P<1 \times 10^{-8}$

Table S5. Genotyping results at ss158145716 and ss158145717, located within the core of Pel.

| Population | Phenotype | Habitat | ss158145716 |  |  |  | ss158145717 |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | Failed | Successful | Total | \% Successful | Failed | Successful | Total | \% Successful |
| Completes | Pelvic-complete | Mixed | 5 | 239 | 244 | 98\% | 3 | 241 | 244 | 99\% |
| Reduced | Pelvic-reduced | Freshwater | 100 | 55 | 155 | 35\% | 102 | 54 | 156 | 35\% |
| Pel-deleted | Pelvic-reduced | Freshwater | 99 | 8 | 107 | 7.5\% | 99 | 9 | 108 | 8.3\% |
| Non-Peldeleted | Pelvic-reduced | Freshwater | 1 | 47 | 48 | 98\% | 3 | 45 | 48 | 94\% |
| BDGB | Pelvic-complete | Marine | 0 | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| LITC | Pelvic-complete | Marine | 0 | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| GJOG | Pelvic-complete | Marine | 0 | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| GORT_4 | Pelvic-complete | Marine | 0 | 12 | 12 | 100\% | 1 | 11 | 12 | 92\% |
| KODK | Pelvic-complete | Marine | 0 | 11 | 11 | 100\% | 0 | 11 | 11 | 100\% |
| NVRO_1 | Pelvic-complete | Marine | 0 | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| RABS | Pelvic-complete | Marine | 0 | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| TYNE_1 | Pelvic-complete | Marine | 0 | 12 | 12 | 100\% | 1 | 11 | 12 | 92\% |
| BIGR_3_63 | Pelvic-complete | Marine | 0 | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| BIGR_20_33 | Pelvic-complete | Freshwater | 0 | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| FLCK_128 | Pelvic-complete | Freshwater | 1 | 7 | 8 | 88\% | 0 | 8 | 8 | 100\% |
| FRIC | Pelvic-complete | Freshwater | 0 | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| FRIL | Pelvic-complete | Freshwater | 0 | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| FTC | Pelvic-complete | Freshwater | 0 | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| OLNY | Pelvic-complete | Freshwater | 0 | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| TYNE_8 | Pelvic-complete | Freshwater | 0 | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| WMSO | Pelvic-complete | Freshwater | 0 | 9 | 9 | 100\% | 0 | 9 | 9 | 100\% |
| PRIL | Pelvic-complete | Freshwater | 0 | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| PRIB | Pelvic-complete | Freshwater | 0 | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| PAXL | Pelvic-complete | Freshwater | 1 | 11 | 12 | 92\% | 0 | 12 | 12 | 100\% |
| WALC | Pelvic-complete | Freshwater | 3 | 9 | 12 | 75\% | 1 | 11 | 12 | 92\% |
| WALR* | Pelvic-reduced | Freshwater | 8 | 4 | 12 | 33\% | 7 | 5 | 12 | 42\% |
| FADA | Pelvic-reduced | Freshwater | 12 | 0 | 12 | 0\% | 12 | 0 | 12 | 0\% |
| VIFI | Pelvic-reduced | Freshwater | 11 | 0 | 11 | 0\% | 12 | 0 | 12 | 0\% |
| MVRO | Pelvic-reduced | Freshwater | 11 | 1 | 12 | 8.3\% | 11 | 1 | 12 | 8.3\% |
| PAXB | Pelvic-reduced | Freshwater | 9 | 3 | 12 | 25\% | 9 | 3 | 12 | 25\% |
| WHAL | Pelvic-reduced | Freshwater | 12 | 0 | 12 | 0\% | 12 | 0 | 12 | 0\% |


| BEPA | Pelvic-reduced | Freshwater | 12 | 0 | 12 | 0\% | 12 | 0 | 12 | 0\% |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| BOOT | Pelvic-reduced | Freshwater | 12 | 0 | 12 | 0\% | 12 | 0 | 12 | 0\% |
| HUMP | Pelvic-reduced | Freshwater | 12 | 0 | 12 | 0\% | 12 | 0 | 12 | 0\% |
| DOLO $^{\dagger}$ | Pelvic-reduced | Freshwater | 0 | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| SCAD | Pelvic-reduced | Freshwater | 0 | 12 | 12 | 100\% | 3 | 9 | 12 | 75\% |
| BOUL $\ddagger$ | Pelvic-reduced | Freshwater | 1 | 11 | 12 | 92\% | 0 | 12 | 12 | 100\% |
| ORPH $\ddagger$ | Pelvic-reduced | Freshwater |  | 12 | 12 | 100\% | 0 | 12 | 12 | 100\% |
| * Represent a mixture of two pelvic-reduced haplotypes, one of which has Pel deletion. <br> $\dagger$ Genetic and morphological information suggest Pitxl-mediated pelvic reduction <br> $\ddagger$ Genetic and morphological information suggest Pitxl-independent pelvic reduction |  |  |  |  |  |  |  |  |  |  |

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## Supporting Online Material

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Materials and Methods
Figs. S1, S2, S3, S4, S5, S6, S7, S8, S9
Tables S1, S2, S3, S4, S5


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