

## ORIGINAL ARTICLE

# Transcriptome-wide signature of hybrid breakdown associated with intrinsic reproductive isolation in lake whitefish species pairs (*Coregonus* spp. Salmonidae)

S Renaut and L Bernatchez

Département de Biologie, IBIS (Institut de Biologie Intégrative et des Systèmes), Université Laval, Québec, Canada

Genetic analyses of speciation have focused nearly exclusively on retrospective analyses of reproductive isolation between highly divergent species. Yet, a full understanding of the speciation process must encompass analysis of the consequences of genomic divergence in young lineages still capable of exchanging genes under natural conditions. The accumulation of conditionally neutral genetic variation may lead to the evolution of divergent gene networks. In a hybrid background, such mutations may no longer compensate one another, resulting in the appearance of selectively disadvantageous traits, including disruption of gene expression regulation. Here, we documented genome-wide patterns of gene expression divergence between young lineages of normal and dwarf lake whitefish and their backcross hybrids for which strong, yet incomplete post-zygotic isolation barriers exist. A significant proportion (33%) of backcross hybrids showed developmental abnormalities not seen in parental forms and eventually leading to death. Although the

transcriptome of parental forms was nearly identical during embryonic development, suggesting a role for stabilizing selection, all hybrids displayed strongly divergent patterns of gene expression. By comparing healthy, surviving hybrids against moribund ones showing abnormal development, we observed that over 2000 genes were misregulated in these abnormal embryos. In particular, misregulation was significantly biased toward essential developmental genes, which were severely underexpressed. Furthermore, genes previously documented to be highly transgressive (exaggerated inter-individual variance) were almost invariably underexpressed in hybrids. Our results thus clearly showed a transcriptome-wide signature of hybrid breakdown in young, incipient species and demonstrated a persuasive link between misexpression of essential developmental genes and post-zygotic isolation.

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## Introduction

Despite ongoing efforts, the nature of the genomic changes underlying reproductive isolation and ultimately leading to speciation remain elusive (Coyne and Orr, 2004; Schluter, 2009; Presgraves, 2010). On the basis of our current comprehension of post-zygotic isolation, hybrid sterility and/or inviability arises as a consequence of incompatible allele combinations in hybrids (Dobzhansky–Muller incompatibilities) that have diverged between pure species or populations (Dobzhansky, 1937; Muller, 1942). These alleles, whether neutral or positively selected within their own genetic background interact negatively when brought together in inter-specific hybrid genomes. The break-up of co-adapted gene complexes will therefore generate mosaic chromosomes composed of the two diverging genomes and is expected to be more deleterious in later generation hybrids compared with

F1 generations (that is, hybrid breakdown; Rieseberg *et al.*, 1999; Coyne and Orr, 2004; Burton *et al.*, 2006). It is believed that the most significant point in the speciation process is the initial development of reproductive isolation (Dettman *et al.*, 2007; Via, 2009). By studying diverging lineages, which are only partially reproductively isolated, early barriers to gene flow and their underlying genetic basis can be identified before they become confounded with other differences that accumulate after speciation. Yet, studies of the effect of genomic incompatibilities on gene regulation have focused nearly exclusively on retrospective analyses of reproductive isolation between highly divergent species in which gene flow does not occur in natural conditions (Michalak and Noor, 2003; Ranz *et al.*, 2004; Noor and Feder, 2006; Landry *et al.*, 2007; Rottschmidt and Harr, 2007).

At the gene transcription level, genomic incompatibilities can lead to the disruption of the transcriptional machinery and the appearance of novel, unforeseen patterns of gene expression in hybrid lineages (Landry *et al.*, 2007). This is often hypothesized to be a major factor underlying hybrid breakdown. Analyses of gene expression profiles using microarrays have shown that the combination of divergent regulatory elements within

Correspondence: Dr S Renaut, Department of Botany, University of British Columbia, 6270 University Boulevard, Vancouver, British Columbia, Canada V6T 1Z4.

E-mail: sebastien.renaut.1@ulaval.ca

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a common genetic background often results in the disruption of gene expression (reviewed by Landry *et al.*, 2007; Ortiz-Barrientos *et al.*, 2007). For example, the global expression profile of *Drosophila melanogaster* and *Drosophila simulans* are more closely related to each other than to their hybrid progeny (Ranz *et al.*, 2004). Similarly, in sympatric but ecologically divergent populations of brook charr (*Salvelinus fontinalis*), dramatic breakdown of gene expression patterns in hybrids compared with their parental relatives was observed (Mavarez *et al.*, 2009).

Lake whitefish species pairs represent excellent model species to study the early onset of reproductive isolation and its effect on genomic divergence (Lu and Bernatchez, 1998; Rogers and Bernatchez, 2007; Bernatchez *et al.*, 2010). Following the last glacier retreat (<15 000 years ago), secondary contact of lake whitefish (*Coregonus clupeaformis*) evolutionary lineages isolated during the Pleistocene (100 000–200 000 years ago) has led to the parallel evolution of two morphologically and ecologically divergent sympatric species in several lakes of northeastern North America: benthic normal and limnetic dwarf whitefish (Bernatchez 2004). Furthermore, the ecological specificity of each lake seems to be the main driving isolation factor because, in certain lakes, secondary contact of these same lineages resulted in a hybrid swarm (Lu *et al.*, 2001; Landry *et al.*, 2007). As expected from a recent divergence event, the overall level of genetic differentiation between species pairs is relatively low and hybrids are still present at low frequency in natural populations (Falush *et al.*, 2007). Moreover, early in ontogeny, normal and dwarf fish are phenotypically and ecologically indistinguishable (Chouinard and Bernatchez, 1998; Bernatchez, 2004). At the same time, hybrid whitefish experience striking fitness consequences of genomic incompatibilities and ongoing reproductive isolation (increased mortality, disruption of hatching time and reduced sperm performance, Rogers and Bernatchez, 2006; Whiteley *et al.*, 2009; Bernatchez *et al.*, 2010).

On the basis of this knowledge, we followed the ontogeny of pure dwarf and normal whitefish as well as their hybrids to identify the precise moment at which embryos started to show early signs of hybrid breakdown. By quantifying genome-wide levels of gene expression when developmental defects were the most extreme, we could identify transcriptome changes potentially associated with post-zygotic isolation. Most specifically, we tracked patterns of regulation for genes known to be essential for early fish development (Amsterdam *et al.*, 2004). We predicted that these essential genes would be the most affected in hybrids because their expression is especially critical and has a severe cascading effect on embryonic development and fitness in general.

## Materials and methods

### Strains, crosses and fish maintenance

Eggs were obtained from outbred laboratory strains (normal whitefish originating from Aylmer Lake (45° 54'N, 71° 20'W), dwarf whitefish originating from Témiscouata Lake (47° 41'N, 68° 47'W), as detailed in Lu and Bernatchez, 1998) at the Laboratoire de Recherche en Sciences Aquatiques (LARSA, Université Laval,

Quebec, Canada). We created half-sib families as follows. The dwarf group was created using one lab strain dwarf female crossed to five different dwarf males. The normal whitefish group was created by crossing one lab strain normal female to three separate normal males. Finally, an independent group of backcross individuals was obtained using an F1-hybrid female generated in the laboratory in a previous study (Rogers and Bernatchez, 2007) crossed to five normal males. As such, the backcross individuals are composed of a 75% normal and 25% dwarf genetic background. Fish were anesthetized with 0.001% eugenol solution whereupon eggs and semen were stripped and immediately fertilized *in vitro*. Fertilized eggs were disinfected (0.0001% iodine solution) and incubated on submerged grids in the same flow-through system (4.5–5.5 °C). To avoid contamination by fungi, dead eggs were removed on a daily basis and all samples were treated weekly with malachite green oxalate.

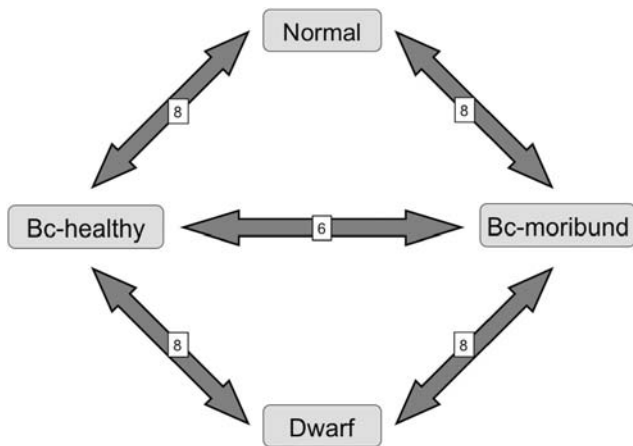
### Sampling

We sampled embryos approximately 60 days post-fertilization (between 280 and 288 degree-days) in normal, dwarf and backcross families. Individually chosen eggs were preserved in RNA later (Ambion, Austin, TX, USA) and frozen at –20 °C for storage. Embryos sampled in the backcross family were separated into normally healthy looking ones, hereafter referred as *backcross-healthy*, and ones that showed developmental problems, hereafter referred as *backcross-moribund* (see results for the description of the developmental phenotype of each group). To avoid sampling dead eggs in which RNA degradation would affect gene expression measurements, 40 independent moribund embryos were observed daily to confirm that all were still alive at least several days following sampling (yolk sac of dead embryos changes from translucent to opaque). At the same time, these 40 moribund embryos as well as nearly all eggs classified as backcross-moribund died in the following weeks, before or just after hatching (at ~100 days).

### Experimental design

Gene expression analysis was performed using the 16 000 (v2.0) Salmon complementary DNA microarray provided by the cGRASP (consortium for Genomic Research on All Salmon Project, von Schalburg *et al.*, 2005). More than 175 complementary DNA libraries constructed from a wide variety of tissues and different developmental stages (Atlantic salmon and rainbow trout) were used to produce the array. Furthermore, the validity of this array has been amply demonstrated in lake whitefish, whereas hybridization signal is the same as for Atlantic salmon (von Schalburg *et al.*, 2005). Sequence divergence between normal and dwarf is also very low and therefore did not affect hybridization kinetics (Derome *et al.*, 2006; Whiteley *et al.*, 2008). Samples of dwarf, normal, backcross-healthy and backcross-moribund were hybridized in a loop design, involving eight biological replicates for the backcross-healthy and backcross-moribund comparison and six for the others. Dye swap was performed between each replicate. As a result, we obtained a final set of 32 microarray slides (Figure 1).

Total RNA was extracted using the Trizol Reagent protocol (Invitrogen, Carlsbad, CA, USA). A pool of five



**Figure 1** Microarray design with the four experimental groups. Each double-headed arrow represents one microarray slide. Numbers written on arrows refer to the number of comparisons done between each group for a total of 32 comparison and 28 separate biological samples.

whole embryos preserved in RNA later was homogenized using a bead mill (Qiagen, Germantown, MD, USA) for each sample hybridized on the array. RNA pooling is a common practice when quantity of material is limiting and inference on gene expression patterns for most genes is not affected by this (Kendzioriski *et al.*, 2005). Crude total RNA was treated with DNase I for 15 min at room temperature (1 unit/ug, Invitrogen) and further cleaned by ultra filtration using microcon YM30 spin columns (Millipore, Billerica, MA, USA). Total RNA was eluted in pure water supplemented with Superase-In RNase Inhibitor (Ambion), quantified and quality checked using Experion RNA StdSens Analysis Kit (Bio-Rad, Hercules, CA, USA), then stored at  $-80^{\circ}\text{C}$ . Reverse transcription reactions were performed using Superscript II Kit (Invitrogen). Following this, Genisphere (Hatfield, PA, USA) 50Array Expression Array Detection Kit (Cy3/cy5) was used following the vendor's protocol ( $\sim 20\text{ug}$  of complementary DNA material hybridized). Microarrays were scanned using a ScanArray Express scanner (Packard Bioscience, Wellesley, MA, USA) and spots were located and quantified using the histogram method in QuantArray 3.0 (Perkin Elmer, Waltham, USA). Gene expression data were deposited at Gene Expression Omnibus ([www.ncbi.nlm.nih.gov/geo](http://www.ncbi.nlm.nih.gov/geo), series accession GSE23095).

#### Data analysis

Unless stated otherwise, all analyses were performed in R (v.2.9.0. The R Foundation for Statistical Computing, 2009, 3-900051-07-0). First, local background was subtracted from spots. Only spots above the mean of empty (blank) spots plus 2 s.d., in at least one channel, were kept for further analysis. Missing data were imputed using the K-nearest neighbors (20 neighbors) and data were  $\log_2$  transformed and normalized using Lowess algorithm. Data were fitted into a mixed model analysis of variance (R/maanova package v1.14, Kerr *et al.*, 2000) to identify differential expression. The mixed model included the following terms as fixed sources of

variance: group (dwarf, normal, backcross-healthy and backcross-moribund) and dye (two fluorescent dyes). Sample (28 biological samples) and array (32 individual microarrays) were included as random sources of variance. Finally, a surrogate variable analysis (Leek and Storey, 2007) enabled the identification of an eigengene explaining 68% of the residual variance and corresponding to a strong block effect (eight slides of the backcross-healthy/backcross-moribund comparisons done in August 2007, the rest of the slides done in June 2008). As a result, this block effect was also included in the mixed model as a random term. A permutation-based F-test was performed to test for statistically significant divergence in gene expression and restricted maximum likelihood was used to solve the mixed model equations ( $P$ -value  $< 0.05$ , 1000 permutations, Fs-test). Following this,  $P$ -values were corrected for multiple hypotheses testing using  $q$ -value correction ( $q$ -value  $< 0.05$ , Storey, 2002). To test for specific pairwise differences between groups (N, D, BC-healthy, BC-moribund), permutation-based  $t$ -tests between the four groups for all genes identified as differentially expressed in the analysis of variance ( $q$ -value  $< 0.05$ , 1000 permutations, Fs-test) were performed. Finally, fold changes for the group effect ( $\log_2$ ) were also extracted from the analysis for all significant transcripts. In order to remain conservative in interpreting the number of significant features, transcripts with  $< 1\%$  sequence divergence over 95% of the sequence printed on the array were also compressed into a single gene.

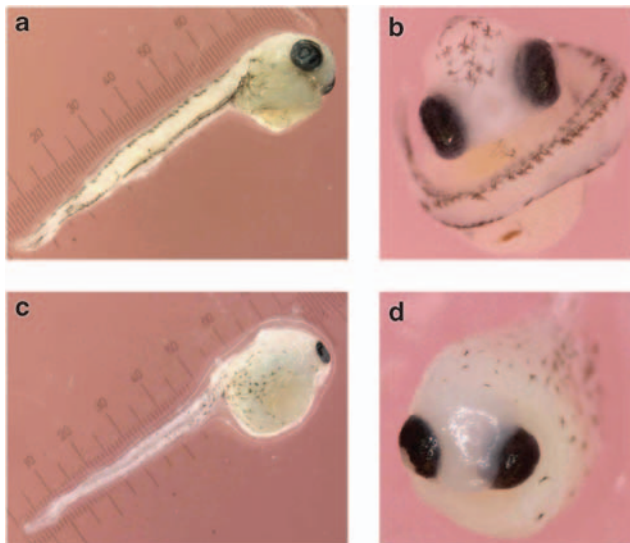
Best linear unbiased estimates of the dye, array and block effects (that is, technical variation) were subtracted from the normalized fitted values of gene expression data for each gene. Following this, each gene expression value was divided by the mean of its channel such that values should only represent the group and sample effects (that is, biological variation). Finally, as the same biological sample is sometimes represented on several arrays, we calculated mean gene expression for each of the 28 different samples used in the study. These values were then used to perform a hierarchical clustering analysis to construct a heatmap, with pairwise gene and sample distance matrices estimated from Pearson correlation coefficients.

Functional classification and assessment of significant differential representation of functional classes were performed using DAVID bioinformatics resources (v6.7, Huang *et al.*, 2009). First, Unigen clusters were obtained from cGRASP for all annotated transcripts printed on the microarray. Then, overrepresented gene ontology (GO) classes (molecular function or biological process) among differentially expressed genes were identified (modified Fisher's test (*Ease score*) corrected for multiple hypothesis testing (*Benjamini correction*)  $< 0.05$ ). Finally, previously published data (Amsterdam *et al.*, 2004) were used to identify patterns of gene expression for essential embryonic development genes in fish. Briefly, Amsterdam *et al.* (2004) identified a suite of 315 genes in early embryonic development in which knockdown mutations are lethal in zebrafish (*Danio rerio*). Furthermore, their study also confirmed that the propensity of being an essential gene is highly conserved throughout evolution. We then used BLASTn algorithm (Altschul *et al.*, 1997) to match these essential genes to our own sequences printed on the array ( $e$ -value  $< 1e-15$ ).

## Results

### Development

Approximately 60 days post-fertilization (between 280 and 288 degree-days) embryos were sampled in normal, dwarf and backcross families. At this stage, embryos have developed into the pharyngula period, which corresponds to the phylotypic stage (*cf. Danio rerio* developmental staging in Slack *et al.*, 1993; Kimmel *et al.*, 1995). Nearly all normal and dwarf embryos showed the same developmental phenotype, and only six embryos out of 104 (5.8%) assessed for developmental characteristics were abnormally developing in these pure crosses. It is noteworthy that these six embryos had major developmental defects that were substantially different (no axial body plan, 'cyclops') from the backcross-moribund group described below. Normally developing pure and backcross embryos were phenotypically similar and characterized by: differentiated pigment cells, a beating heart pumping blood throughout the circulatory system, eyes pigmented and a fully mobile tail, detached from the yolk sac (Figures 2a and b). In the backcross family, a significant proportion of embryos (33%, 52 out of 156 embryos visually inspected (Fisher's exact test,  $P$ -value < 0.0001)) showed either evident lag in development or atypical phenotype (small head, small eyes, deformed eye lens, heart not beating, reduced cell pigmentation, deformities of the tail) and were hereafter referred to as backcross-moribund embryos (Figures 2c and d). Thus, while not all backcross-moribund possessed exactly the same phenotype, they undoubtedly represented a discrete phenotype different from all other pure and healthy backcross embryos.



**Figure 2** (a, b) General developmental characteristics of normal, dwarf and backcross-healthy embryos (differentiated pigment cells, beating heart pumping blood throughout the circulatory system, eyes pigmented, eye lens completely formed and a fully mobile tail, detached from the yolk sac). (c, d) General phenotype of backcross-moribund embryos (small head, small eyes, deformed eye lens, reduced cell pigmentation, deformities of the tail). Note that egg chorion was removed (a, d) and embryos were manually unfolded with dissecting needles in a and c. Units on scale are 0.1 mm.

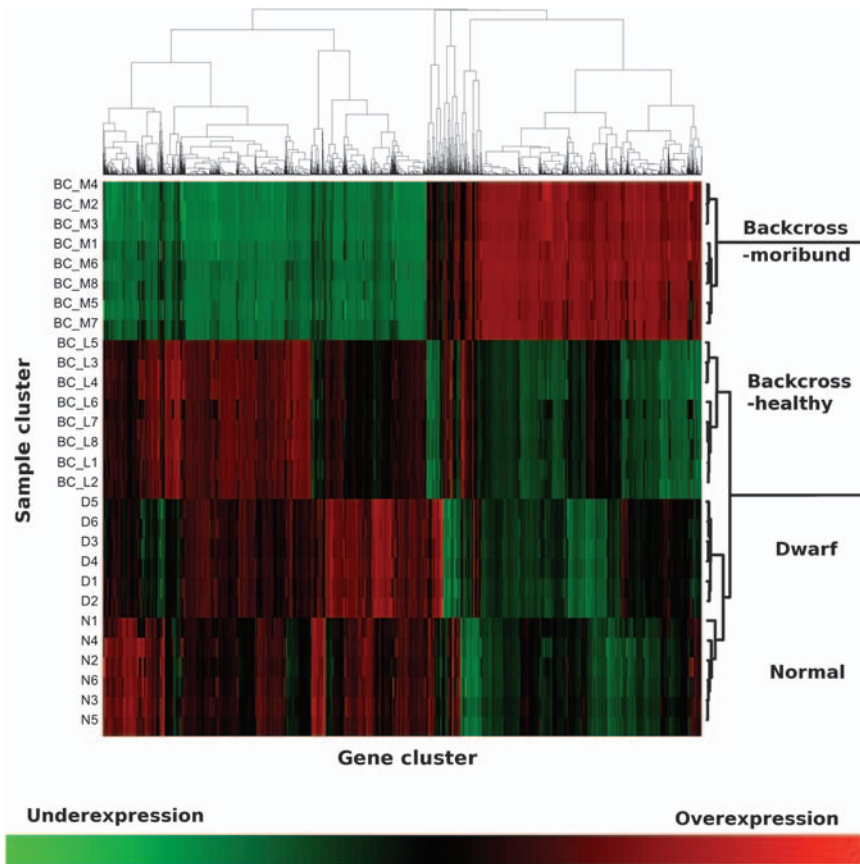
### Gene expression regulation

Very little difference in gene expression was found between normal and dwarf whitefish, with only two transcripts differentially expressed (60s ribosomal protein L23a, hemoglobin subunit beta) after correction for multiple hypotheses testing and replicate spotting on the array, 0.1% of all transcripts expressed (false discovery rate  $q$ -value < 0.05). This was in stark contrast with backcross-healthy hybrids for which 162 and 77 (3.0 and 1.5%) genes were differentially expressed compared with normal and dwarf, respectively. Even more strikingly, 2214 (39%) transcripts were differentially expressed between backcross-healthy and backcross-moribund, 1993 (35%) between normal and backcross-moribund and 1964 (35%) between dwarf and backcross-moribund (Figure 3, Supplementary Table 1).

### Functional classification

Functional analyses using DAVID bioinformatic resources revealed that several biological processes and molecular functions were overrepresented among the lists of differentially expressed genes. First, translation and generation of precursor metabolites and energy GO terms were significantly overrepresented in all comparisons involving backcross-moribund individuals (Table 1). Comparatively, glycolysis, alcohol and carbohydrate catabolic processes were overrepresented in the list of genes differentially expressed between backcross-healthy and dwarf, while no GO terms were overrepresented in the backcross-healthy/normal or normal/dwarf categories. Transcripts differentially expressed in backcross-moribund were further divided into two categories: transcripts underexpressed and overexpressed compared with the average of all groups. In overexpressed genes, a total of 15 different GO terms, which were further grouped into 7 general categories, were overrepresented. These were mostly related to transport and energy metabolism functions. In underexpressed genes, 11 different GO terms, which were further grouped into six general categories, were overrepresented. These were related to completely distinct functional categories; mostly macromolecule metabolism and regulation of mRNA translation.

Of particular interest were 450 (3.2% of all transcripts printed on the array) transcripts, which matched (BLASTn  $e$ -value <  $1e^{-15}$ ) to genes essential for early fish development (Figure 4). Essential genes refer to those genes for which knockdown mutations are embryonic lethal in *Danio rerio* (Amsterdam *et al.*, 2004). Among these essential transcripts, 336 were expressed and 204 differentially expressed. Both of these values were significantly higher compared with the proportion of all transcripts expressed and differentially expressed, respectively ( $\chi^2$  test,  $P$ -value < 0.01). Moreover, 81% (166 transcripts) of these essential transcripts were underexpressed in backcross-moribund compared with the average of the four groups and this was highly significant compared with the proportion of all transcripts underexpressed in these embryos (56% or 1445) ( $\chi^2$  test,  $P$ -value < 0.0001, Figure 4). We also identified several GO terms that were overrepresented among the lists of essential genes differentially expressed against all genes differentially expressed. These were related to different functional categories but mostly,



**Figure 3** Heatmap representing hierarchical clustering based on gene distance matrix (all genes differentially expressed between groups as identified by analysis of variance (ANOVA,  $q$ -value  $<0.05$ ) and sample distance matrix (28 biological samples).

macromolecule metabolism and regulation of mRNA translation (Table 1).

Finally, in a previous study looking at gene expression differences in 30-day-old whitefish embryos, a suite of seven genes that showed highly transgressive patterns of expression in backcross hybrids and which also comprised three essential developmental genes was identified (Renaut *et al.*, 2009). These genes did not differ in mean expression in 30 days old backcross embryos, but showed exaggerated inter-individual variance extending outside the range of both parents. In this study, 18 transcripts matched to these 7 transgressive genes and all except 1, were underexpressed in both backcross groups compared with the parents. Moreover, this effect was even more pronounced for backcross-moribund (Table 2).

## Discussion

Our main objective was to quantify genome-wide levels of gene expression when developmental defects were the most extreme and thus potentially associate changes in expression to post-zygotic isolation. Here, we discuss the implications and inherent limitations of these results in light of our understanding of post-zygotic isolation in incipient species of lake whitefish and of the genetics of speciation in general.

A significant proportion of backcross embryos showed strong developmental defects. These developmental problems leading to a mortality rate of at least 33%

clearly represent a severe fitness cost of producing hybrids between dwarf and normal whitefish in natural conditions and therefore undeniably act as a strong post-zygotic isolation mechanism. Admittedly, as these observations are drawn from five half-sib backcross families derived from a single hybrid female crossed to five different normal males, they should be interpreted cautiously. However, the fertilization success of those backcross half-sib families eggs was similar to the one observed for pure crosses, as also previously reported (Lu and Bernatchez, 1998; Rogers and Bernatchez, 2006). In addition, the distinct abnormal embryonic development phenotype observed in backcross embryos was not observed in any of the pure families. Moreover, our observations indicating that strong post-zygotic isolation barriers exist between normal and dwarf lake whitefish is corroborated by several lines of evidence previously documented. An elevated mortality rate around the same developmental time (Rogers and Bernatchez, 2006) as well as a reduced sperm performance (Whiteley *et al.*, 2009) has been identified in independent backcross families. Elevated, albeit lower than for the backcross, mortality was also observed in F1 hybrids at the same developmental time (Lu and Bernatchez, 1998). Thus, increased mortality in hybrids has now been observed in three independent studies. Furthermore, strong segregation distortion for over 30% of mapped genetic markers reflected differential survival rates among backcross hybrids genotypes (Rogers and Bernatchez, 2007). For all these reasons, we propose that the developmental and

**Table 1** Significant over-representation of GO categories (BP and MF) among genes, which showed different transcription levels between all genes expressed on the microarray and the different experimental groups

Comparison	Category	Description <sup>a</sup>	Fold <sup>b</sup>	Count (%) <sup>c</sup>	P-value <sup>d</sup>	Number of GO terms <sup>a</sup>
Backcross-moribund/backcross-healthy	BP	Generation of precursor metabolites and energy	1.74	67 (10.2%)	0.000	1
	BP	Translational elongation	1.91	28 (4.2%)	0.066	1
	MF	Structural molecule activity	1.49	65 (9.8%)	0.022	1
Backcross-moribund/dwarf	MF	Hydrogen ion transporter activity	1.86	28 (4.2%)	0.047	1
	BP	Generation of precursor metabolites and energy	1.72	60 (10.1%)	0.000	1
	BP	Translation	1.84	70 (11.7%)	0.022	2
Backcross-moribund/normal	MF	Structural molecule activity	1.58	62 (10.4%)	0.006	1
	BP	Generation of precursor metabolites and energy	1.78	63 (10.3%)	0.000	1
Backcross-healthy/dwarf	BP	Translational elongation	2.07	28 (4.6%)	0.010	1
	MF	Cytoskeletal protein binding	1.75	38 (6.2%)	0.021	1
	BP	Glycolysis	21.08	6 (22.2%)	0.001	1
	BP	Alcohol catabolic process	15.17	6 (22.2%)	0.009	1
Backcross-healthy/normal	BP	Carbohydrate catabolic process	14.59	6 (22.2%)	0.006	4
	—	—	—	—	—	—
Dwarf/normal	BP	Generation of precursor metabolites and energy	2.09	62 (18.5%)	0.00	1
	BP	Oxidation reduction	1.76	50 (14.9%)	0.00	1
	BP	Electron transport chain	2.12	30 (8.9%)	0.00	1
	BP	Phosphorylation	1.82	29 (8.6%)	0.05	1
	MF	Catalytic activity	1.24	148 (44%)	0.00	—
	MF	Magnesium ion binding	2.05	21 (6.3%)	0.01	—
All genes differentially expressed and backcross-moribund overexpressed genes	MF	Ion transporter activity	1.80	28 (8.2%)	0.02	9
	BP	Translation	1.41	69 (15.4%)	0.001	1
	BP	Gene expression	1.29	136 (30.3%)	0.000	1
	BP	Macromolecule metabolic process	1.26	182 (40.4%)	0.00	4
	BP	RNA metabolic process	1.45	48 (10.7%)	0.033	2
	MF	Structural constituent of ribosome	1.48	8.9 (40%)	0.029	1
All genes differentially expressed and backcross-moribund underexpressed genes	MF	Nucleic acid binding	1.32	86 (19.2%)	0.016	2
	BP	Translation	3.91	22 (48.9%)	0.000	1
	BP	Gene expression	2.64	32 (71.1%)	0.000	1
	BP	Macromolecule biosynthetic process	2.08	30 (66.7%)	0.002	6
	BP	Protein metabolic process	1.97	32 (71.9%)	0.001	3
	MF	Structural constituent of ribosome	5.20	16 (35.6%)	0.000	1
All genes differentially expressed and essential genes differentially expressed	MF	Structural molecule activity	3.25	16 (35.6%)	0.002	1

Abbreviations: BP, biological process; GO, gene ontology; MF, molecular function.

<sup>a</sup>Similar GO terms were grouped together as one category using the most inclusive GO term provided (for example, *macromolecule metabolic process* grouped within *macromolecule biosynthetic process*). Number of original GO terms in each category was written in *number of GO terms* column.

<sup>b</sup>Fold enrichment values given by DAVID as compared to the background group (either genes expressed or all genes differentially expressed).

<sup>c</sup>Number of unigenes in GO category and percentage of total.

<sup>d</sup>Note that when more than one GO term were included in the same category, the *P*-value (Ease score (Bonferroni correction implemented by Huang *et al.*, 2009)) was the mean *P*-values of all the GO terms.

expression differences observed here are direct consequences of hybridization between diverging lineages.

#### The specifics of hybrid breakdown on gene expression regulation

Using complementary DNA microarrays, we observed very little difference in gene expression between pure normal and dwarf whitefish. This lack of divergence between the transcriptome of the parental forms early in ontogeny is in line with the lack of differentiation in developmental time until emergence (Rogers and Bernatchez, 2006) and also corroborates previous findings whereas 30 days old embryos had 14 times less genes ( $n=5$  genes) displaying significant regulatory divergence than 16 weeks old juvenile fish (Nolte *et al.*, 2009; Renaut *et al.*, 2009). In fact, dwarf and normal larval whitefish are nearly indistinguishable and their

phenotypic and ecological divergence is expected to take place at the juvenile stage (Chouinard and Bernatchez, 1998). Embryos sampled here were in the phylotypic developmental stage corresponding to the pharyngula period in *Danio rerio* development (Kimmel *et al.*, 1995). During this highly conserved stage of development, individuals are expected to be more similar to each other than during any earlier or later developmental periods (Slack *et al.*, 1993). This observation has frequently been made even for highly divergent taxa and should also apply for gene expression differences (Irie and Sehara-Fujisawa, 2007).

In stark contrast with parental forms, hybrids showed developmental problems that had far-reaching repercussions on gene expression regulation. Backcross hybrids all had strong divergent patterns of gene expression compared with parental crosses and this was especially true for moribund hybrids. The nature of breakdown in

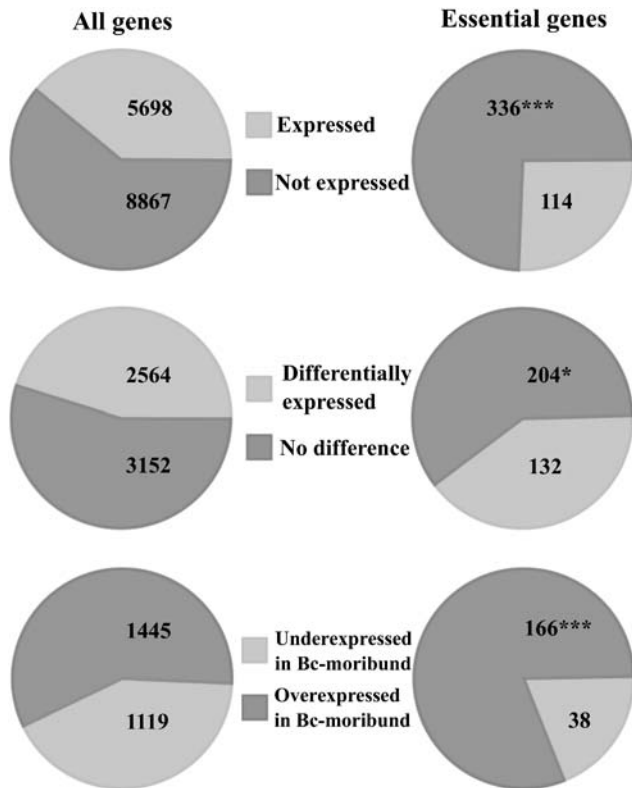
gene regulation observed in whitefish backcross hybrids is more likely a consequence of genomic incompatibilities rather than the mere result of a stochastic RNA degradation process. Genes were affected in a precise way depending on their functionality. This is also confirmed by functional analysis, whereby markedly different GO terms (whether biological process or molecular function) were overrepresented whether a

gene was under or over expressed in the backcross-moribund group (Table 1). Therefore, developmental problems seem to be associated with this breakdown in gene regulation in hybrids. Nevertheless, the challenge will remain to establish whether this association is through causality and this remains an unavoidable limit of all gene expression studies of speciation (Noor and Feder, 2006).

As predicted, essential fish developmental genes were the most affected in hybrids. Their expression is especially critical and has a severe cascading effect on embryonic development and fitness in general. These genes, whose essentiality is highly conserved throughout evolution (Amsterdam *et al.*, 2004), were not only more differentially expressed than expected but also strongly underexpressed in backcross compared with the average of the four groups (81% of those are underexpressed in backcross embryos, Figure 4). The functional analysis of GO terms also revealed that similar categories were overrepresented in essential genes differentially expressed as in backcross-moribund underexpressed genes and these were mostly genes involved in macromolecule metabolism and regulation of mRNA translation (Table 1).

Furthermore, the early signs of gene misexpression previously documented in 30 days old embryos (Nolte *et al.*, 2009; Renaut *et al.*, 2009) culminated in dramatic differences in development and gene expression regulation in the present study in 60 days old hybrid embryos, particularly so for genes known to be essential in early embryonic development (Table 2). For example, in zebrafish, knockdown mutation of Heat Shock Cognate 70 leads after 5 days to: pericardial edema, a necrotic head and a general degeneration of the body; and knockdown of elongation factor 1 alpha leads to a small head and eyes, rounder yolk and increased necrosis (Amsterdam *et al.*, 2004). In lake whitefish backcross hybrids, both of these genes were among the most transgressive of all at 30 days (Renaut *et al.*, 2009), were severely underexpressed in the present study (at 60 days, Table 2), and embryos showed similar general phenotype to zebrafish mutants (Figures 1c and d).

A remaining challenge is now to determine how many genomic regions actually contribute to reproductive barriers. One possibility is that this is attributed to



**Figure 4** Number of transcripts matching to essential early developmental genes. Note that transcripts were defined as essential if they matched (BLASTn *e*-value < 1e-15) to the list of 315 genes identified as essential early developmental genes in zebrafish (Amsterdam *et al.*, 2004) ( $\chi^2$  test: \**P*-value < 0.05, \*\*\**P*-value < 0.001).

**Table 2** Relative gene expression for genes previously identified as highly transgressive in 30 days old backcross whitefish embryos (Renaut *et al.*, 2009)

Description	Fold change (N)	Fold change (D)	Fold change (Bc-h)	Fold change (Bc-m)	<i>q</i> -Value (Bc-m/Bc-h)	<i>q</i> -Value (Bc-m/D)	<i>q</i> -Value (Bc-m/N)	<i>q</i> -Value (Bc-h/D)	<i>q</i> -Value (Bc-h/N)	<i>q</i> -Value (N/D)
Asialoglycoprotein receptor 2	0.00	0.03	0.22	0.21	NS	**	**	NS	*	NS
Protein kinase C	0.00	-0.23	-0.27	-0.23	NS	NS	**	NS	*	NS
Guanine nucleotide-binding protein	0.00	-0.29	-0.42	-0.39	NS	NS	***	NS	***	NS
<b>40S ribosomal protein S11 (3 transcripts)</b>	0.00	0.06	-0.03	-0.22	**	***	*	NS	NS	NS
<b>Heat shock 70 kDa protein (10 transcripts)</b>	0.00	-0.21	-0.30	-0.42	NS	NS	***	NS	NS	NS
<b>Elongation factor 1 alpha</b>	0.00	-0.36	-0.39	-0.56	**	**	***	NS	**	NS
Fish-egg lectin	0.00	-0.07	-0.27	-0.95	***	***	***	NS	NS	NS

Abbreviations: Bc-h, backcross-healthy; Bc-m, backcross-moribund; D, dwarf; N, normal; NS, nonsignificant. *q*-values for the respective *t*-tests. \**q*-value < 0.05, \*\**q*-value < 0.01, \*\*\**q*-value < 0.001.

Genes products in bold are also essential developmental genes in fish embryos according to Amsterdam *et al.* (2004). Note that relative gene expression (log<sub>2</sub> values) is expressed relative to the normal group. When several transcripts match to the same gene product, relative gene expression was calculated as the mean of all transcripts matching to that gene product.

the accumulation of conditionally neutral mutations throughout the genome. This is an attractive hypothesis as mutations accumulate throughout the genome at all times in all populations. As such, they provide different architectural starting points, which may subsequently be recruited by adaptive processes and potentially lead to different evolutionary trajectories, whether under stabilizing or divergent selection pressures (Lynch, 2007). Here, as very little difference was observed between normal and dwarf whitefish, post-zygotic reproductive isolation in these young species appears to involve mostly genes under stabilizing selection for gene expression, as it has been previously shown in drosophilids (Haerty and Singh, 2006) and brook charr (Mavarez *et al.*, 2009).

The fact that in whitefish, over 30 % of mapped genetic markers show locus-specific deviations from expected Mendelian segregation (segregation distortion) tends to support the idea that many 'ordinary loci' are associated with reproductive isolation (Rogers and Bernatchez, 2007). On the other hand, we cannot refute that one or a few genetic loci may underlie large-scale disruption of expression, especially so if those are associated with essential developmental genes, in which knockdown mutations are known to be lethal. Another explanation worth mentioning pertains to the fact that large-scale hybrid gene misexpression can also result from a variety of mechanisms related to the disrupting the chromatin integrity or the expression of non coding RNAs (Michalak, 2009). For example, miRNA networks, which have been shown to regulate gene expression in early development in many vertebrate species, can have a large cascading effect if disrupted and thus provide a likely 'major gene' effect (Lee *et al.*, 2007; Michalak, 2009). As such, all other differences in gene expression and phenotype would be a downstream effect of this major factor. Other minor incompatibilities could still exist, which would account for the differential expression of genes observed in the healthy backcross fish versus parental types. Along this line, we previously observed that a least one locus associated with embryonic mortality is also linked to a regulation (expression Quantitative Trait Locus, eQTL) hotspot and may harbor a major regulatory gene with strong pleiotropic effect on the expression of numerous genes (Bernatchez *et al.*, 2010), thus representing a possible mechanism explaining the genome-wide disruption of regulation in hybrids.

## Conclusion

By comparing healthy, surviving hybrids against moribund ones, our results identified a transcriptome-wide signature of hybrid breakdown in young, incipient species and revealed a persuasive link between misexpression of essential developmental genes and post-zygotic isolation. Our analysis of the genomic basis of ongoing speciation in young evolutionary fish lineages helps to bridge the gap between ecological studies of reproductive isolation with limited knowledge of their genetic basis, and genetic studies of speciation with an incomplete ecological perspective. Quite clearly, both are needed toward a truly general theory of the genetics of speciation.

## Conflict of interest

The authors declare no conflict of interest.

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