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Lineage/species-specific expansion of the Mx gene family in teleosts: Differential expression and modulation of nine Mx genes in rainbow trout Oncorhynchus mykiss

Tingyu Wang, Fuguo Liu, Guangming Tian, Christopher J. Secombes, Tiehui Wang

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9	Tingyu Wang ¹ , Fuguo Liu ¹ , Guangming Tian ¹ , Christopher J Secombes ¹ *, Tiehui Wang ¹ *
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13 14	^a Scottish Fish Immunology Research Centre, School of Biological Sciences, University of Aberdeen, Aberdeen AB24 2TZ, UK
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18	*Corresponding authors:
19	Dr. Tiehui Wang, E-mail: <u>t.h.wang@abdn.ac.uk</u>
20	Dr. Christopher J. Secombes, E-mail: <u>c.secombes@abdn.ac.uk</u>
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22 Abstract

 Myxovirus resistance (Mx) proteins are interferon (IFN)-inducible Dynamin-like GTPases, which play an important role in antiviral immunity. Three Mx genes (Mx1-3) have been cloned previously in rainbow trout. In this study, an additional six Mx genes were cloned that reside in four chromosomal loci. Further bioinformatics analysis suggests the presence of three teleost Mx groups (TMG) each with a characteristic gene organisation. Salmonid Mx belong to TMG1 and TMG2. The increased salmonid Mx gene copies are due mainly to local gene duplications that happened before and after salmonid speciation, in a lineage/species specific manner. Trout Mx molecules have been diversified in the loop 1 and 4 regions, and in the nuclear localisation signal in loop 4. The trout Mx genes were shown to be differentially expressed in tissues, with high levels of expression of TMG1 (Mx1-4) in blood and TMG2 (Mx5-9) in intestine. The expression of the majority of the trout Mx genes was induced by poly IC *in vitro* and *in vivo*, and increased during development. In addition, induction by antiviral (IFN) and proinflammatory cytokines was studied, and showed that type I IFN, IFNγ and IL-1β can induce Mx gene expression in an Mx gene-, cytokine- and cell line-dependent manner. These results show that salmonids possess a large number Mx genes as well as complex regulatory pathways, which may contribute to their success in an anadromous life style.

Key words: Rainbow trout, Mx, anti-viral defence, evolution, gene expression, modulation, type I interferon,

40 IFN γ , IL-1 β

1. Introduction

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- 44 Mx (myxovirus resistance) proteins are interferon (IFN)-inducible Dynamin-like GTPases, with an important
- 45 role in antiviral immunity [1-2]. They are members of a family of large GTPases, and share an N-terminal
- 46 GTPase domain, a middle domain (MD), and a C-terminal GTPase effector domain (GED). The GTPase
- domain is the most conserved part that consists of a tripartite GTP-binding motif (GDXXSGKS, DLPG, and
- 48 TKPD) and a dynamin signature (LPRXXGXXTR). The MD is important for oligomerization and viral target
- 49 recognition, whilst the GED has a conserved C-terminal leucine zipper that folds back to join the N-terminal
- 50 GTP-binding domain to establish the enzymatically active center of Mx proteins [1-3]. Mx proteins form
- tetramers in solution that oligomerize further into large filaments and rings [3], with both GTPase activity and
- oligomerization required for antiviral immunity.
- A prototype Mx gene has been found in amphioxus, containing the N-terminal GTPase domain [4]. Typical
- Mx genes are found in all vertebrate groups. The first evidence of Mx genes in fish started with the isolation
- of an Mx genomic DNA fragment in perch (*Perca fluviatilis*) in 1989 [5]. The first full-length characterisation
- of Mx genes was reported in rainbow trout *Oncorhynchus mykiss*, that has three Mx genes (Mx1-3) [6-7].
- 57 Subsequently Mx genes have been characterised in many fish species, with 1-9 genes present [4, 8-21].
- However, some fish species such as the Gadiformes have lost their Mx genes [22]. The role of fish Mx
- 59 proteins in antiviral defence has been established in a few species, such as Japanese flounder *Paralichthys*
- 60 olivaceus, Atlantic salmon Salmo salar and grass carp Ctenopharyngodon Idella [23-26].
- The multiple copies of mammalian Mx are closely linked and arise from local gene duplications [2]. How
- multiple fish Mx genes have evolved is currently unclear [4,16]. A recent publication has shown that there are
- 63 nine Mx genes present on three chromosomes (Ch) in Atlantic salmon with Mx1-3 on Ch12, Mx4-8 on Ch25,
- and Mx9 on Ch9 [27]. The origin of multiple copies of Mx genes on the same chromosome, that are linked
- closely and share high sequence identities, is likely to also be via local gene duplications. However, due to the
- the state of the same angle sequences, as the same see the section and the same seems and the same seems and the same seems are same seems are same seems and the same seems are same see
- 67 copy in mammals are present as four copies on four chromosomes in salmonids [28]. Thus it is possible there

third teleost-wide whole genome duplication (3R WGD) and the salmonid 4R WGD, many genes with single

- 68 could be a fourth chromosome that harbours Mx genes in salmonids, and if discovered this may shed light on
- 69 how the different Mx-bearing chromosomes evolved in salmonids.
- 70 Mammalian Mx gene expression is induced by type I and type III IFNs but not by type II IFNy or other
- 71 proinflammatory cytokines [29-31]. Interestingly, the diversified repertoire of Atlantic salmon Mx genes
- appear to show some differential responsiveness to type I and II IFNs, with those on Ch12 being highly
- 73 induced by type I IFNs and those on Chr25 being more strongly induced by IFNy than by type I IFN [27].
- 74 This finding is very interesting and raises the question as to whether a diversified Mx repertoire may also be
- 75 responsive to other cytokines released during innate antiviral defence, and remains to be examined. Hence, in
- 76 this study we aimed to shed light on Mx gene evolution in actinopterygian fish, in an attempt to establish a

better model of their evolution, and to establish whether the increased Mx copy number in salmonids has allowed neo-functionalisation giving a broader responsiveness to a variety of cytokines. We first identified and cloned an additional six Mx genes in rainbow trout, and found that all salmonids with a mapped genome have four chromosomes harbouring Mx genes. We identified three groups of Mx genes present in teleosts in a lineage-specific manner, with some (Ostariophysi) having all three groups, some having two groups (Protacanthopterygii, including salmonids) but the percomorphs possessing only a single group. We next investigated the expression of the nine trout Mx family members individually. We found that the trout Mx genes are differentially expressed constitutively in tissues, that they increase during development, are induced *in vivo* by poly IC, and are modulated *in vitro* by type I and type II IFNs, and by other proinflammatory cytokine in a gene-, cytokine- and cell line-specific manner.

2. Materials and methods

2.1. Rainbow trout

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- 90 Healthy rainbow trout (~40 g) were purchased from the Mill of Elrich Trout Fishery (Aberdeenshire,
- 91 Scotland, UK). The fish were fed twice a day with a commercial diet (EWOS) and maintained in 1-m-
- 92 diameter fibreglass tanks with recirculating freshwater at 14°C at the Scottish Fish Immunology Research
- 93 Centre, University of Aberdeen, UK. Head kidney (HK) swabs were taken routinely and showed no bacterial
- 94 presence [32]. Fish were given at least two weeks for acclimation prior to use and ranged in size from 100-140
- 95 g when experiments were performed. All the experiments described comply with the Guidelines of the
- 96 European Union Council (2010/63/EU) for the use of laboratory animals, and were carried out under UK
- 97 Home Office project licence PPL 60/4013, approved by the ethics committee at the University of Aberdeen.

2.2. Identification, cloning and sequence analysis of Mx cDNA in rainbow trout

- Three Mx genes (Mx1-3) are known in rainbow trout [6-7]. To identify additional Mx genes in this
- species, we searched the recently released rainbow trout reference genome (GCF_002163495.1)
- using TBLASTN [33] with the known trout Mx genes as query, resulting in the identification of four
- genomic loci (Chromosomes (Ch)3, 11, 17 and 24) that harbour Mx genes. The Mx genes were then
- predicted as described previously [34-35]. In addition, potential exons in untranslated regions (UTR)
- were predicted by using trout RNA-seq datasets (SRP108798) through aligning to the reference
- genome. Primers (supplementary **Table S1**) were subsequently designed in the predicted 5'- and 3'-
- 106 UTR for PCR cloning of the complete coding region of each predicted Mx gene. The general cloning
- and sequence analysis was as described previously [34-35]. The nucleotide sequences generated
- were assembled and analysed with the AlignIR programme (LI-COR, Inc.). Homology search was
- performed using the BLAST program (http://blast.ncbi.nlm.nih.gov/Blast.cgi) [33] and the gene
- organization was predicted using the Spidey program at NCBI. Protein prediction was undertaken
- using software at the ExPASy Molecular Biology Server (http://www.expasy.org/tools) [36].
- Multiple sequence alignments were generated using CLUSTALW [37]. Amino acid sequence
- identity/similarity comparison was performed using the scoring matrix BLOSUM62 within the
- MatGAT program, with a gap open penalty of 10 and gap extension penalty of 1 [38].

2.3. Analysis of Mx genes in other salmonids

- The Mx genes in other salmonids were predicted/analysed using recently released genomes of
- Atlantic salmon (Salmo salar, Atlantic, acc. no. GCF_000233375.1), chinook salmon (Oncorhynchus
- tshawytscha, Chinook, acc. no. GCF_002872995.1), coho salmon (Oncorhynchus kisutch, Coho, acc.

- no. GCF_002021735.1), and Arctic charr (*Salvelinus alpinus*, Charr, acc. no. GCF_002910315.2).
- Each Mx aa and nucleotide sequence was mapped to chromosomes/scaffolds. Similarly, Mx genes
- were analysed in the pike (*Esox lucius*) reference genome (acc. no. GCF_000721915.3), the closest
- relative of salmonids that has not undergone the salmonid 4R WGD and that has a sequenced
- genome. The aa sequences were used for phylogenetic tree analysis using MEGA7.0 software [39]
- based on an multiple alignments generated by CLUSTALW. The evolutionary distances were
- 125 computed using the JTT matrix-based method. A neighbour-joining phylogenetic tree was
- constructed using pair-wise deletion option.

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2.4. Evolutionary analysis of teleost Mx family

- Mx genes/proteins were analysis at NCBI from selected teleost fish, including species known to
- possess multiple Mx genes. The naming of Mx genes/proteins followed those already published [4,
- 130 16, 40-41] or simple Mx with an acc. no. For phylogenetic tree analysis, Mx protein sequences were
- extracted from one holostean species, spotted gar (*Lepisosteus oculatus*, Lepisosteiformes) that is an
- early actinopterygian fish species without the 3R WGD, twenty-one teleosts and three mammals
- 133 (human Homo sapiens, mouse Mus musculus and cow Bos taurus) as an outgroup. The teleost
- species included an elopomorph, European eel (*Anguilla anguilla*, Anguilliformes), five Ostariophysi
- 135 (Otophysi) species including three Cypriniformes fish (common carp Cyprinus carpio, goldfish
- 136 Carassius auratus and zebrafish Danio rerio), channel catfish (Ictalurus punctatus, Siluriformes),
- and Mexican tetra or blind cave fish (Astyanax mexicanus, Characiformes), five protacanthopterygii
- 138 (the salmonids and pike described above), and ten percomorphs including two Pleuronectiformes
- 139 (turbot Scophthalmus maximus, and olive flounder Paralichthys olivaceus), Atlantic killifish
- 140 (Fundulus heteroclitus, Cyprinodontiformes), stickleback (Gasterosteus aculeatus,
- 141 Gasterosteiformes), Medaka (Oryzias latipes, Beloniformes), fugu (Takifugu rubripes,
- 142 Tetraodontiformes), Nile tilapia (*Oreochromis niloticus*, Cichliformes) and three Perciforme fish
- 143 (gilt-head sea bream *Sparus aurata*, orange-spotted grouper *Epinephelus coioides*, and the Asian sea
- bass Lates calcarifer). A neighbour joining phylogenetic tree was constructed as above. Synteny
- analysis was performed using the Genomicus program [42] or with information extracted from
- reference genome sequence at NCBI.

2.5. Real-time PCR analysis of gene expression

- 148 Specific primers for each Mx gene were carefully designed based on a multiple cDNA sequence alignment to
- ensure that at least one primer was isoform specific, and one primer crosses an intron to prevent genomic
- DNA amplification. The primers for qPCR analysis of Mx genes and other cytokine genes are detailed in

- Table S1 and S2, respectively. Total RNA preparation, cDNA synthesis and qPCR analysis were as described
- previously [43]. The expression of each gene was first normalized to that of the house keeping gene
- elongation factor-1α (EF-1α). To directly compare the expression level of the different Mx paralogues, a
- reference was constructed using equal molar amounts of PCR product from each gene, including EF-1α.

2.6. Tissue distribution of rainbow trout Mx gene family

- Six healthy rainbow trout (~140 g) were killed and seventeen tissues (blood, thymus, gills, scales,
- skin, muscle, tail fins, adipose fin, brain, adipose tissue, spleen, liver, heart, intestine, gonad, head
- kidney (HK) and caudal kidney) were collected and processed as described previously [34-35]. The
- relative expression level of Mx genes in each sample was normalized against the expression level of
- EF-1α and expressed as arbitrary units (AU) where 1 AU = the expression level of EF-1α/1,000,000.

2.7. Ontogeny of the expression of the Mx gene family

- To investigate if the expression of Mx is correlated to immune capacity in early life, the ontogeny of the
- expression of Mx genes was examined. Archived samples from a previous experiment were used in this study
- as detailed in Wang et al. [44]. Briefly, eyed eggs, immediate post-hatch fry, pre-first feeding (Pre-feeding)
- fry at the stage of full disappearance of the yolk sac, and fry 3 weeks following first feeding were sampled and
- 166 cDNA prepared. Six samples for each developmental stage were prepared. The qPCR quantification of gene
- expression was as described above.

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2.8. Production of recombinant trout type I IFNa

- The cDNA sequence encoding the mature peptide of trout IFNa was amplified from a poly IC
- stimulated cDNA sample using the primers IFNaF (TGTGACTGGATCCGACACCAT) and IFNaR
- 171 (GTACATCTGTGCTGCAAGGATATCC). The amplified product was cloned to a pTriEx vector
- 172 (Novagen) as described previously [45]. Sequence analysis of the construct used for recombinant
- protein production revealed that it encodes a His-tag (MAHHHHHHHHG) at the N-terminus
- followed by the 152 aa mature peptide identical to XP_021480273. Thus, the recombinant trout IFNa
- was 163 aa with a calculated molecular weight of 19.5 kDa and a theoretical pI of 9.17. A sequence
- confirmed plasmid was transformed into BL21 Star (DE3) competent cells (Invitrogen). The protein
- was produced, purified under denaturing conditions, refolded, and quantified as described previously
- 178 [34,43,45]. The refolding buffer was phosphate buffered saline (PBS, pH7.4, Sigma, UK) containing
- 179 10% glycerol, 0.5 M arginine monohydrochloride, and 5 mM 2-mercaptoethanol (2-ME). The
- purified protein was buffer changed using a centrifugal concentrator (10 kDa cutoff, Thermo
- Scientific). The storage buffer was PBS (pH7.4) containing 10% glycerol, 2 mM EDTA, 10 mM

- arginine monohydrochloride, 10 mM glutamine, and 5 mM 2-ME. After sterilization with a 0.2 µm
- filter, the recombinant protein was aliquoted and stored at -80°C ready for bioactivity analysis.

2.9. Stimulation of cell lines with PAMPs and recombinant cytokines

- Three trout cell lines, a macrophage-like cell line RTS-11 from spleen [46], a fibroblast-like cell line RTG-2
- from gonad [47], and an epithelial-like cell line RTGill from gills [48] were used for *in vitro* stimulation. All
- the cells were maintained at 20°C in Leibovitz medium (L-15) supplemented with 100 U/ml penicillin and
- 188 100 μg/ml streptomycin (P/S), and 10% (for RTG-2 and RTGill cell lines) or 30% (for RTS-11 cells) foetal
- bovine serum (FBS). The cells were seeded at 1x10⁶ cells/ml (RTS-11) or 0.5x10⁶ cells/ml (RTG-2 and
- 190 RTGill) in L-15 containing 10% FCS at 2 ml/well in 12-well cell culture plates overnight before stimulation.
- 191 RTS-11 cells were first stimulated with pathogen-associated molecular patterns (PAMPs), the bacterial cell
- wall component lipopolysaccharide (LPS, from E. coli strain 055:B5, Sigma) and the viral dsRNA mimic
- 193 polyinosinic: polycytidylic acid (poly IC, Sigma). The stimulants were added to the cells at 25 μg/ml for LPS
- and 50 µg/ml for poly IC, or medium alone as control. The treatments were terminated by dissolving the cells
- in TRI reagent (Sigma, UK) 4 h, 8 h and 24 h post-stimulation. Total RNA isolation and gene expression
- analysis was as described above.
- 197 The RTS-11 cells were then stimulated with five trout recombinant cytokines, IFNγ (20 ng/ml) [49], IFNα (25
- 198 ng/ml) prepared above, IL-1β (25 ng/ml) [50], IL-6 (100 ng/ml) [51] and TNFα (50 ng/ml) [52], or medium
- alone as control. The treatments were terminated at 4 h, 8 h and 24 h and gene expression analysed as above.
- 200 Finally, RTG-2 and RTGill were stimulated with IFNγ (20 ng/ml) and IFNα (25 ng/ml) for 4 h and gene
- 201 expression analysed as above.

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2.10. Modulation of Mx gene expression in vivo by poly IC

- 203 Poly I:C (Sigma, UK) was dissolved at 10 mg/ml in sterile cell culture-grade water, stored at -80 °C and
- 204 diluted to 5 mg/ml in PBS before intraperitoneal (ip) injection. Trout (~100 g, N=24) were injected
- intraperitoneally (ip) with 1 mg poly IC in 0.2 ml of PBS, or the same amount of PBS as control. Six fish from
- each group were killed at 6 h and 24 h post injection, and spleen, HK, gills and intestine were collected for
- gene expression analysis as described previously [53]. The time points chosen were based on past studies of
- 208 the rapid PAMP response in vivo in rainbow trout [54]. The expression was expressed as AU after
- normalisation with EF-1 α , where 1 AU = the average expression level in control fish at 6 h in each tissue.

210 2.11. Statistical analysis

- The data were statistically analyzed using the SPSS Statistics package 24 (SPSS Inc., Chicago, Illinois). The
- analysis of real-time PCR data was as described previously (43). To improve the normality of data, real-time

213	quantitative PCR measurements were scaled, with the lowest expression level in a data set defined as 1, and
214	log2 transformed. One way-analysis of variance (ANOVA) and the LSD post hoc test were used to analyse
215	the gene expression data, with $P \le 0.05$ between treatment and control groups considered significant.

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218 **3. Results**

- 3.1. Identification, cloning and sequence analysis of Mx gene family in rainbow trout
- In addition to the known Mx1-3 in rainbow trout, six additional Mx genes (Mx4-9) have been identified and
- cloned in this study (Supplementary Figs. S1-S6, acc. nos. MK301134-MK301139). Mx4, as with Mx1-3, was
- located on Ch17 and was located between Mx2 and Mx3. Mx5-6, Mx7-8 and Mx9 were located on Ch3, Ch11
- and Ch24, respectively (**Table 1**).
- Each trout Mx cDNA sequence had a complete open reading frame that encoded for 635, 614, 606, 613, 608,
- and 640 aa for Mx4-9, respectively. Each trout Mx had a N-terminal dynamin GTPase domain, and a C
- terminal GTPase effector domain, that were well conserved as shown in a multiple alignment of the 9 trout
- and two human Mx proteins (Fig. 1). The tripartite GTP-binding motif (GDXXSGKS, DLPG, and TKPD) in
- 228 all trout Mx were identical to human MxA and MxB. The dynamin signature (LPRXXGXXTR), and the
- leucine residues that form leucine zipper folds in the GTPase effector domain, were also conserved (Fig. 1).
- The middle domain and the GTPase effector domain of Mx fold into a four-helical bundle that constitutes a
- stalk that mediates oligomerization and transmits conformational changes from the G domain to the target
- structure [55]. The regions forming the helix, and loops L2 and L3 were all conserved. However, relatively
- 233 large differences were present in loops L1 that connects the N and C-terminal of the helix α1 and introduces a
- kink, and L4 that connects the helix $\alpha 3$ and $\alpha 4$ (Fig. 1). Potential nuclear localisation signals (KKRKR) are
- present in trout Mx2 and Mx4 in L4, where a lysine motif (KKKK) is also present in human MxA that
- contribute to membrane association of MxA [2].

3.2. Sequence analysis of Mx family in salmonids

- Nine Mx genes (Mx1-9) have been described recently in Atlantic salmon [27] that map to three chromosomes
- 239 (Ch9, 12 and 25, **Table 1**). In addition, a partial sequence for Atlantic Mx10 (XP_013998960) has been
- mapped to Ch15 (**Table 1**). At least 6 Mx genes each in chinook salmon and coho salmon, and 10 Mx genes
- in Arctic charr could be identified and mapped to chromosomes or scaffolds (**Table 1**). Partial sequences for
- three pike Mx genes were also found, with Mx1 on Ch17 and Mx2-3 on Ch13 (**Table 1**).

- A phylogenetic tree constructed using all the known salmonid Mx and the three pike Mx protein sequences 243 244 showed that the salmonid Mx family clustered into four separate clades (Fig. 2A). Moreover, salmonid Mx 245 genes are located on four cognate chromosomes, at least in rainbow trout, Atlantic, chinook and coho salmon, 246 in which their genome sequences have been mapped to chromosomes. The Mx genes on the same 247 chromosome are grouped together (Fig. 2B), as seen also in Atlantic salmon [27], suggesting that multiple 248 genes on the same chromosome originate from local gene duplication events in each species. Thus there are 249 four salmonid Mx groups (SMG, Fig. 2). Pike Mx1 was grouped with SMG1, whilst pike Mx2 and Mx3, 250 which are linked on Ch13, were grouped with SMG2. SMG1 consisted of Mx1-4 of trout, Mx1-3 of Atlantic, chinook and coho salmon, and Mx1-2 of charr. SMG2 consisted of trout Mx5-6, Atlantic salmon Mx4-8, 251 252 chinook and coho salmon Mx4, and charr Mx3-8. SMG3 contained trout Mx7-8, Atlantic salmon Mx10, chinook and coho salmon Mx5 and char Mx9-10. SMG4 had trout and Atlantic salmon Mx9, and chinook and 253 254 coho salmon Mx6 (Fig. 2).
- It is notable that trout Mx1/3 and Mx2/4, along with their cognate salmonid Mx molecules formed two separate branches with high bootstrap support in SMG1 group (**Fig. 2A**), suggesting that the existence of these genes, or their ancestral gene predates salmonid speciation. A similar situation was also observed with trout Mx7 and Mx8 in SMG3 group (**Fig. 2A**). Although more Mx genes might still be found, the data for salmonids with an advanced (sequenced) genome suggests that the distinct numbers of Mx genes in SMG1-3 are due to species-specific independent local gene duplication or deletion events after salmonid speciation.
- 261 In agreement with four SMGs in the phylogenetic tree, the Mx aa sequences within each SMG share high aa 262 identities (**Table 2**). In SMG1, trout Mx1-4 share high as sequence identities between each other (86.3-98.4%) 263 in similar range to SMG1 Mx from different salmonids (83.4-98.2%), but have relatively low identities to Mx from SMG2 (43.8-47.0%), SMG3 (41.5-48.1%) and SMG4 (42.1-48.1) (**Table 2**). Similarly, Mx sequences 264 265 share high identities within SMG2 (82.7-93.6), SMG3 (57.8-93.8%) and SMG4 (80.2-97.0%). However, the identities of Mx between SMGs are similarly low (41.5-51.6%) with the exception of Mx molecules in SMG2 266 267 and SMG3 that share moderate 53.4-71.2% as identities (Table 2). Furthermore, the Mx bearing 268 chromosomes in rainbow trout (Ch3, 11, 17 and 24) and Atlantic salmon (Ch9, 12, 15 and 25) do not share 269 syntenic origins [56-57]. These data suggest that the four Mx-bearing chromosomes do not appear to originate 270 from the salmonid 4R WGD.

3.3. Phylogenetic tree analysis of Mx in vertebrates

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To understand how the four SMGs evolved, we analysed the Mx gene family in other vertebrates with a focus on teleost Mx genes. Three Mx genes, Mx1-2 on Ch17 and Mx3 on Ch3, are present in spotted gar, an early Actinopterygian (Holostei) that has not undergone the 3R WGD that may represent an ancestral state [4, 27]. A neighbour-joining phylogenetic tree was constructed based on a multiple alignment of Mx proteins from selected mammalian and teleost fish species. In agreement with previous studies, mammalian Mx form an

independent group separate from all teleost Mx molecules (Fig. 3). Three teleost Mx groups (TMG) can be 277 278 identified, with a gar Mx at the root of each clade. TMG1 contained gar Mx1, salmonid SMG1 and Mx from 279 all the major teleost groups (Fig. 3). TMG2 contained gar Mx3, salmonid SMG2, SMG3 and SMG4, and Mx 280 molecules from European eel, zebrafish, goldfish, Mexican tetra and catfish. TMG3 contained gar Mx2, and 281 Mx from Cypriniformes (zebrafish, common carp and goldfish) and Characiformes (tetra) (Fig. 3). This 282 phylogenetic tree may suggest that the 3R WGD duplicated 6 Mx genes (from the 3 ancestral Mx genes 283 present in gar) that have subsequently undergone lineage specific deletion, with Cypriniformes and Characiformes species retaining a copy of each of the duplicates, protacanthopteryggii such as the salmonids 284 and pike retained two whilst in the majority of teleosts, the percomorphs only one is present. Within a species 285 286 the numbers of Mx genes might be increased again by local gene duplication.

3.4. Synteny analysis of Mx locus in vertebrates

- Despite much analysis, the evolutionary relationship of Mx genes in different vertebrates is still unclear [4,16].
- In the present study we performed a synteny analysis using the most advanced genomes available. Pike Ch17
- 290 (Mx1) and trout Ch17 (Mx1-4), and pike Ch13 (Mx2-3) and trout Ch11 (Mx7-8) share a considerable syntenic
- relationship (Fig. 4). However, trout Ch3 (Mx5-6) and Ch24 (Mx9) share no clear syntenic relationships to
- pike Mx loci, but have a good relationship instead with gar Ch17 (Mx1-2) and Ch3 (Mx3), respectively (Fig.
- **293 4**).

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- Interestingly, the Gar Mx3 (Ch3) locus also has considerable synteny with zebrafish Ch15 (MxF) and Ch25
- 295 (MxD, G1 and G2), in addition to the trout Mx9 locus, and all the Mx residing in these loci belong to TMG2.
- Furthermore, zebrafish Ch15 and 25 combined share a perfect match syntenically to gar Ch3, suggesting a
- break of the ancestral gar-like derived chromosome in zebrafish. The gar Mx3 locus also shares synteny with
- the tetrapod Mx locus, eg. human Ch21 (MxA and MxB) (Fig. 4), as also reported by Robertsen et al. [27].
- 299 This suggests that a gar Mx3-like ancestral locus gave rise to the teleost Mx group loci in zebrafish and
- 300 salmonids, and led to the tetrapod Mx locus.
- The gar Mx1-2 locus shares synteny to tetra Ch12 (Mx1-7) and the combined tetra Ch19 (Mx8) and scaffold
- 302 NW_019172839 (Mx9) of tetra, indicating the retention of two 3R derived Mx loci in this species. Similarly,
- the gar Mx1-2 locus shares synteny to both zebrafish Ch1 (MxA/B) and Ch9 (MxC/E) (Fig. 4). It is notable
- that the tetra Ch12 Mx locus has Mx genes that belong to TMG1 (Mx1) as well as TMG2 (Mx2-7), whilst the
- zebrafish TMG2 resides in Ch15 and 25 derived from gar Ch3. Taken as a whole, two ancestral gar-like Mx
- loci gave rise to the current teleost Mx loci in a lineage-specific manner.

3.5. Gene organisation analysis of Mx genes in vertebrates

- To shed more light on the evolution of the three teleost Mx groups, we analysed the gene organisation of all
- trout Mx genes in comparison with Mx genes from other teleosts and humans. All exon-intron boundaries of

- 310 trout Mx genes conformed to the consensus sequences (GT/AG). In TMG1, trout Mx1-4 genes all had a 12
- 311 coding exon/11 intron structure with identical intron phase. The coding region of exons were identical with
- 312 the exception of exon 8 and 11 (Fig. 5). A similar gene organisation of the coding region was observed with
- other TMG1 genes, eg. zebrafish MxA and MxB that are on the same chromosome, tetra Mx1 and gar Mx1,
- with the exception of a non-coding exon in the 5'UTR of the gar and tetra Mx gene in this group (**Fig. 5**).
- 315 Trout Mx5-9 belong to TMG2. They all had 13 coding exons with the coding regions of exons well conserved,
- with the exception of the first and the last two exons (**Fig. 5**). Compared to TMG1 genes, the last eleven intron
- 317 phases were identical to that of TMG1 genes. The main difference in gene organisation was an extra N-
- 318 terminal coding exon that brought a phase II intron in Mx5-9 that was missing in Mx1-4. This gene
- organisation was conserved in other TMG2 Mx genes except gar Mx3, that shared the same gene organisation
- with TMG1 Mx genes (**Fig. 5**).
- Human MxA and MxB also had the same 13 coding exon structure as TMG2 but with the first intron in phase
- 322 0 (Fig. 5). Interestingly, some TMG3 Mx genes had the same gene organisation as in humans (zebrafish MxE
- and gar Mx2), and others (eg. zebrafish MxC and tetra Mx8) had the same as in TMG1 (Fig. 5). Some of
- 324 TMG3 Mx genes have lost the last third exon.
- In general, the exon size and intron phase in the regions encoding for the N-terminal GTPase domain, the
- 326 middle domain and the C-terminal GTPase effector domain are well conserved. The noticeable variations in
- size were the 5th last exon that encodes the L1 loop, and the second last exon that encodes for the L4 loop (**Fig.**
- 328 **5**).

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3.6. The expression of rainbow trout Mx family in tissues and cell lines

- The expression of each trout Mx gene was comparatively studied using gene specific primers and
- serial dilutions of references, and expressed as arbitrary units (AU) relative to EF-1 a expression.
- Thus the AU of the relative expression is on an equal molar basis. The expression of paralogues on
- the same chromosome was grouped together, and the tissues were ordered according to the average
- expression level of Mx1 (Fig. 6). The expression level of Mx1-4 on Ch7 was medium (AU = 100 to
- 1,000) to high (AU > 1,000) across tissues (**Fig. 6A**). Mx5 and Mx6 expression was detectable in all
- the seventeen tissues but at low levels (AU <100). The exceptions were Mx5 in intestine that was at
- a high level, and Mx5 in thymus, gills, adipose fin, tail fins, spleen, scales and gonad, and Mx6 in
- intestine and gills that was at medium expression levels (Fig. 6B). Mx8 expression was also
- detectable in all tissues examined but at low levels except for high level expression in intestine and
- medium level expression in thymus, brain and gonad. Mx7 expression was undetectable in head/
- caudal kidney and tail fins, but had high level expression in intestine, medium level in thymus, with

- low levels in other tissues (Fig. 6C). Mx9 expression was also high in intestine but low or
- undetectable in other tissues (**Fig. 6D**).
- Each Mx gene was differentially expressed across different tissues. In the same tissue, the majority
- of Mx genes had varying expression levels, as shown by the ratio of different genes and paired-
- samples T tests (**Table S3**). In general, the expression of the Mx1-4 and Mx5-9 was more similar
- within each group than between groups. It is noteworthy that blood expressed highest levels of Mx1-
- 4 but low levels of Mx5-9. In contrast, intestine expressed highest levels of Mx5-9 genes amongst
- the tissues examined (**Fig. 6**).
- 350 The constitutive expression of the trout Mx gene family was also examined in three trout cell lines.
- 351 The macrophage-like cell line RTS-11 expressed all the Mx genes at low level (Fig. 6). The
- 352 fibroblast-like cell line RTG-2 and epithelial-like cell line RTGill expressed medium levels of Mx2
- and Mx3, and low levels of other Mx genes but Mx5 and Mx8 in RTG-2 and Mx9 in RTGill were
- not detectable (AU < 1).

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3.7. Transcript expression of Mx gene family during developmental stages

- 356 The high levels of expression of Mx gene family members in blood and intestine suggest an important role in
- immune defence. We next examined the expression of these genes in eyed-eggs, immediately post-hatch fry,
- 358 pre-first feeding fry or fry 3 weeks after first feeding, which represent a critical period when the fish encounter
- potential pathogens from the environment and food [44]. The expression levels of all Mx genes were
- maintained from eyed-eggs till post-hatch. The expression of Mx1, Mx8 and Mx9 was increased in pre-
- feeding fry and maintained at the same levels in post-feeding fry (Fig. 7). Mx5 expression was low in eyed-
- eggs and post-hatch fry but increased significantly in pre-feeding fry and increased further in post-feeding fry.
- 363 The expression of Mx2 and Mx5 was only increased in post-feeding fry whilst that of Mx3, Mx4 and Mx6
- was unchanged across the different developmental stages (**Fig. 7**).

3.8. Modulation of the expression of trout Mx and proinflammatory cytokine genes in vivo by poly IC

- 366 Poly IC, a known strong inducer of Mx expression, was used to investigate its ability to modulate Mx
- expression in vivo. The expression of Mx genes was examined in two major systemic lymphoid tissues, the
- spleen and HK, and two mucosa-associated lymphoid tissues, the gills and intestine. The expression of Mx1-4
- was induced in all four tissues at both 6 h and 24 h post poly IC injection, with the exception of Mx1 and Mx3
- in intestine at 6 h (**Fig. 8A-D**). As seen *in vitro*, poly IC did not increase Mx9 expression *in vivo* (**Fig. 8I**). The
- induction of other Mx (5-8) genes was time- and tissue-dependent (Fig. 8E-H). In the spleen, poly IC
- increased Mx5 and Mx7 expression at 24 h and Mx6 and Mx8 expression at both time points. In the HK, poly
- 373 IC increased Mx5 and Mx8 expression at both time points, and Mx6 expression at 24 h, but had no effect on

- 374 Mx7 expression. In the gills, poly IC increased the expression of Mx6-8 at 24 h, and Mx5 at both time points.
- 375 In the intestine that has high constitutive expression of Mx5-8, poly IC increased Mx5 expression at both time
- points and Mx7 expression at 6 h, but had no effects on Mx6 and Mx8 (**Fig. 8E-H**). In summary, poly IC was
- also a strong inducer of Mx gene expression *in vivo* with highest induction seen at 24 h post-injection (**Fig. 8**).
- 378 In addition to inducing Mx gene expression, poly IC also induced the expression of many pro-inflammatory
- 379 cytokines, including IL-1β1-2, IL-6, IL-8, TNFα1-2, IFNa1, IFNγ and CXCL11, at least at one time point, in
- all the four tissues examined (Fig. 9). In contrast to the later (at 24 h) peak induction of Mx gene expression,
- poly IC induced an early (6 h) induction of the majority of the proinflammatory cytokines studied (eg IL-1β1-
- 382 2, IL-6, IL-8, TNF α 2 and IFNa1) (**Fig. 9**).

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3.9. Modulation of trout Mx expression in RTS-11 cells by PAMPs

- The expression of all trout Mx genes was detectable in the macrophage cell line RTS-11 (Fig. 6). Thus we
- examined the modulation of trout Mx gene family members first in this cell line using poly IC and LPS,
- 386 classical viral and bacterial PAMPs. Poly IC was a strong inducer of Mx gene expression. It significantly
- induced the expression of Mx2, Mx3 and Mx4 from 4 h to 24 h, that of Mx1, Mx5 and Mx6 from 8 h to 24 h,
- and that of Mx7 and Mx8 at 8 h, but had little effect on Mx9 expression (Fig. 10). As expected, LPS had only
- minor effects on Mx gene expression; it induced a small upregulation of Mx4 at 8 h and Mx5 at 24 h, and a
- small downregulation of Mx1 at 24 h and Mx2 at 4 h (**Fig. 10**).

3.10. Modulation of Mx expression by proinflammatory cytokines in RTS-11 cells

392 The early peak induction of proinflammatory cytokine expression and late peak induction of Mx genes may 393 suggest that poly IC can induce Mx indirectly via proinflammatory cytokines as well as by virus sensing 394 pathways. Indeed, IFNy has been shown recently to modulate some of the Mx isoforms in Atlantic salmon 395 [27]. Hence, the possibility of modulation of Mx gene expression by IFNa1, IFNγ, IL-1β, IL-6 and TNFα was studied using RTS-11 cells. Mx9 expression was refractory to all the cytokines (Fig. 11I). However, the 396 397 expression modulation of the other Mx genes was cytokine-specific. IFNa induced the expression of Mx1-4 and Mx6 from 4 h to 24 h, Mx5 at 4 h and 24 h, and Mx7 at 8 h, but had no effects on Mx8 (Fig. 11). IFNy 398 399 induced the expression of Mx2-6 from 4 h to 24 h, Mx7-8 at 24 h, but decreased Mx1 expression at 24 h (Fig. 11A-H). IL-18 induced the expression of Mx3-4 and Mx8 from 4 h to 24 h, Mx2 and Mx6 at 4 h and 8 h, Mx5 400 401 at 24 h, but had no effects on the expression of Mx1 and Mx7 (Fig. 11A-H). IL-6 increased the expression of 402 Mx3 at 8 h and 24 h, Mx5 at 24 h, and Mx6 at 4 h, but decreased the expression of Mx 1 and Mx4 at 24 h, and 403 Mx8 at 8 h and 24 h. It had no effects on Mx2 and Mx7 (Fig. 11A-H). TNFα induced the expression of Mx2 and Mx4 at 8 h, Mx3 from 4 h to 24 h, Mx5 at 24 h, but decreased Mx1 expression at 24 h and Mx5 404 405 expression at 8 h. It had no effects on Mx6-8 (Fig. 11A-H). It is noteworthy that IFNa is a strong inducer of 406 the expression of Mx1-4 and Mx7, IFNγ is a strong inducer of Mx5-6 expression and IL-1β a strong inducer

- of Mx8 (**Fig. 11A-H**). In conclusion, all the Mx genes except Mx9 can be modulated by multiple antiviral and
- 408 proinflammatory cytokines in an Mx- and cytokine-dependent manner.

3.11. Modulation of Mx expression by type I and type II IFNs in RTG-2 and RTGill cell lines

- The cytokine-dependent Mx modulation may also be cell-type dependent. Thus, the IFN modulated Mx gene
- expression was further studied in the fibroblast-like cell line RTG-2 and the epithelial like cell line RTGill. In
- RTG-2 cells, both IFNy and IFNa induced the expression of Mx1-6, with similar potency for Mx5 and Mx6.
- 413 However, IFNa was more potent for Mx1-4 (**Fig. 12**). In RTGill, both IFNγ and IFNa induced the expression
- of Mx2 and Mx3, with IFNy more potent for Mx2, but had no effects on Mx1 and Mx4 (Fig. 12A-D). Only
- 415 IFNγ but not IFNa induced the expression of Mx5 and Mx6 in RTGill cells (Fig. 12E-F). The expression of
- 416 Mx7-9 was low and refractory (data not shown).

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4. Discussion

- This study reveals that at least 9 active Mx genes are present in the rainbow trout genome, the same number as
- 420 reported recently in Atlantic salmon [40]. However, in this study we show that there are in fact 4 Mx loci
- present in salmonids and that the number of Mx genes at each locus differs between these two species at 3 of
- these loci. Multiple Mx genes are also present in other salmonids at four chromosomal loci. The salmonid Mx
- genes at the same genomic locus share high sequence identities within and between species, suggesting they
- arose from local gene duplication events. It seems that local Mx gene duplication/gene loss is common with
- some duplication events likely to have happened before salmonid speciation, eg. duplication of Mx1/3 and
- 426 Mx2/4 in SMG1, and Mx7 and Mx8 in SMG3, but others after salmonid speciation, eg. Atlantic salmon Mx4-
- 8 in SMG2. The four Mx bearing chromosomal loci could have arisen from the 3R and 4R WGDs as seen
- with other genes when mammals have one and salmonids have 4 [28]. However, sequence homology, synteny
- and phylogenetic tree analysis do not clearly support this, and past models [4] do not adequately explain their
- evolutionary path in bony fish.
- 431 Multiple Mx genes (up to 10) can be found in many teleost species. Our phylogenetic tree analysis indicates
- that three TMG exist. TMG1 are present in different teleost lineages, but TMG2 and TMG3 are found in only
- more basal teleosts [57]. Each TMG has a unique gene organisation in terms of coding exon number and the
- first intron phase. For example, whilst TMG1 has a 12 coding exon structure with the first intron in phase I,
- 435 TMG2 has 13 coding exons with the first intron in phase II, and TMG3 has either 13 coding exons with the
- 436 first intron in phase 0 (as seen with mammalian Mx genes) or the same organisation as in TMG1. Interestingly,
- the spotted gar possesses three Mx genes, with one present in each TMG.

Although the four cognate Mx chromosomal loci between salmonids are well conserved, no clear syntenic conservation have been observed in trout and other salmonid species between the four Mx loci. However, a syntenic relationship between the two gar Mx loci and those in zebrafish/ tetra is apparent. For example, the Gar Mx1-2 locus and tetra Mx loci on Ch12 and Ch19, that harbour Mx genes in all the three teleost Mx groups. However, the zebrafish cognate Mx loci of gar Mx1-2 only have Mx genes that belong to TMG1 and TMG3, and the zebrafish TMG2 locus shares synteny to the gar Mx3 locus. The two gar Mx loci also share apparent synteny with two of the trout Mx loci whilst the other two show a syntenic relationship with the two pike Mx loci. This complex syntenic relationship may suggest that the current Mx genes in 3R or 4R teleosts may have arisen from the three Mx genes present at two chromosomal loci as seen in spotted gar, with the 3R duplicated Mx loci retained/lost in a lineage specific manner (Fig. 13). This model differs from that in Qi et al. [4] in taking into account the number of loci present in actinopterygians as well as Mx copy number.

The increased copy number of Mx genes seen in many teleosts may confer increased expression level and hence heightened antiviral defence. The duplicated copies may also acquire novel sequence properties that confer anti-viral specificity and efficiency. The nine trout Mx genes have considerable variation in the nucleotide sequence coding for the L1 and L4 loops in the stalk, as seen in the multiple aa alignments and their gene organisation. Both L1 and L4 are at the surface of the stalk [3] that can interact with surrounding proteins and may be involved in interaction with viral components. L4 of mammalian Mx is a critical determinant of viral substrate specificity [58-59]. The diversification of these regions might have been driven by past virus exposure and life history traits of different species. For example, zebrafish and tetra have short life cycles but live in diverse changing water environments. Their survival depends heavily on innate immunity against viral pathogens. Salmonids survive successfully in both fresh and marine waters, and may encounter a larger virus repertoire compared to species living in only fresh water or marine water. Hence, the increase of copy number and types of Mx genes in these species may confer a fitness advantage.

Mx antiviral effects depend on where the Mx protein is present. Thus, the mouse Mx1 protein which is localized in the nucleus mainly inhibits orthomyxoviruses that replicate in the nucleus, whereas mouse Mx2 is confined to the cytoplasm and inhibits viruses with an exclusively cytoplasmic replication phase [60]. There is a potential NLS in the L4 of some salmonid Mx proteins eg. trout Mx2, and Mx4, but not Mx1 and Mx3. This NLS may indeed be functional as trout Mx2 is found in the nucleus, and Mx1/ Mx3 in the cytoplasm [7]. This suggests the nuclear presence of trout Mx4. Taken as a whole, salmonids, such as rainbow trout are equipped with a battery of diversified Mx genes with their protein products present in the cytoplasm and nucleus to protect themselves from viral attack during their life cycle.

Investigation of Mx isoform expression will help understand their functional roles. Although multiple Mx genes have been identified in several teleost species, a comparative expression study in healthy fish at the individual gene level is lacking [17, 27]. Our results show that the nine trout Mx genes were differentially expressed across different tissues and cell lines, as outlined below, suggesting a level of neofunctionalisation

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of trout Mx paralogues through variation in expression patterns. The high levels of Mx1-4 transcript in blood and Mx5-9 in intestine is of particular interest. Many different viruses can infect hosts via the intestine to cause acute infectious gastroenteritis, or get access to the blood by physical breaches (wounds) or during viremia [61]. So preventing their spread at these sites is a good antiviral strategy. The differential expression of Mx genes in the cell lines may suggest that specific cell types preferentially express a particular Mx gene or a set of Mx genes to defend against potential cell type-tropic viruses. However, the three cell lines examined expressed relatively low levels of Mx genes compared to the tissues analysed, perhaps due to the need for humoral factors present *in vivo* to maintain high level Mx gene expression.

The expression of Mx genes was also studied during development, and several were increased in pre-feeding and post-feeding fry (eg Mx1, Mx2, Mx5, Mx7-9). First feeding is a critical stage in the life of a fish, when potential food borne viruses are met directly for the first time and when the adaptive immune system has not fully developed. Indeed, it was the genes preferentially expressed in the intestine in adults that were increased in the post-hatch fry.

Next we studied whether the Mx genes could be modulated by PAMPS or cytokines. In agreement with Mx induction in other species, poly IC was a strong inducer of trout Mx1-8 gene expression in vitro and in vivo, with Mx9 more refractory. Although the induction patterns in vivo were gene- and tissue-dependent, highest expression was seen at 24 h with most of the Mx genes. Injection of poly IC also induced the expression of proinflammatory cytokines, such as IL-1β, IL-6 and TNFα as well as type I and type II IFNs. In contrast to the late peak of induction of Mx gene expression, poly IC caused an early peak of expression in the cytokines studied. Therefore, the later peak in Mx expression could be influenced by such molecules. To test this hypothesis, we stimulated RTS-11 cells with these cytokines to see if they could modulate Mx expression. Seven of the nine Mx genes were induced by type I IFNa and type II IFNγ, and six were induced by IL-1β. In contrast, IL-6 and TNFa had only minor effects on Mx expression. This cytokine mediated induction was gene-dependent. IFNa was a strong inducer of Mx1-4 and Mx6-7. Past studies have shown Mx1-3 to be modulated by type I IFNs, and so it was no surprise that Mx4 as an additional SMG1/TMG1 member was also induced. Mx6 and Mx7 on the other hand are TMG2 genes. Studies with two other cell lines confirmed the induction of Mx6 by type I IFN as well as a small induction of Mx5 (as seen in RTS11 cells) but Mx7 was not expressed in these cells. IFNy was a strong inducer of Mx5-6, although some induction of Mx1-4 was also seen in the different cell lines. Trout Mx5 and Mx6 are on the same locus as Mx4-8 in Atlantic salmon, that were also responsive to IFNy [27], and are SMG2/TMG2 genes. The SMG3 (Mx7-8) and SMG4 (Mx9) genes did not show this responsiveness. IL-1β was able to induce Mx8 (SMG3) in RTS-11 cells although some induction of Mx2-4 and Mx5-6 was also seen, suggesting a broader responsiveness across SMGs. In common with salmon, no induction of Mx9 was found with these PAMPS/cytokines and its role, if any, in antiviral defence remains to be elucidated.

Such findings contrast with mammalian Mx genes that are strictly induced by type I and type III IFNs but are not induced by IFN γ or other proinflammatory cytokines [29-31]. Salmonids possess multiple type I (IFNa-f) and type II (IFN γ 1-2 and IFN γ rels) IFNs, but type III IFN has not been identified in any fish species [62]. In addition to the induction of Mx genes by type I and type II IFNs, this study confirms that some proinflammatory cytokines also influence Mx expression in fish. IL-1 β in particular has a clear impact on Mx gene expression in trout and was the only cytokine that induced Mx8 expression. Thus, it is apparent that cytokines other than IFNs can have a role in antiviral defence.

Conclusions:

- Up to 10 Mx genes are present in salmonids that reside in four chromosomal loci. Three teleost Mx groups (TMG) can be identified with characteristic gene organisations, each with a spotted gar Mx gene at the root in the phylogenetic tree. Synteny analysis suggests that the current Mx genes in 3R or 4R teleosts may be evolved from the three Mx genes present at two chromosomal loci in spotted gar, with the 3R duplicated Mx loci retained/lost in a lineage specific manner. Salmonid Mx belong to TMG1 and TMG2. The increased salmonid Mx gene copies are due to local gene duplications that have happened before and after salmonid speciation in a lineage/species specific manner. Salmonids are equipped with a diversified battery of Mx genes, with their protein products present in both cytoplasmic and nuclear locations to protect against viral attack during their life in freshwater and seawater.
- Trout Mx genes are differentially expressed in tissues with high levels of expression of TMG1 (Mx1-4) in blood and TMG2 (Mx5-9) in the intestine. The expression of most of the trout Mx genes was induced by poly IC (*in vitro* and *in vivo*), and increased during early developmental stages. In addition to induction by type I IFN, IFNγ and IL-1β also induced Mx expression in rainbow trout and are cytokines that are highly modulated by viral infection. These results show that salmonids possess a large number Mx genes as well as complex regulatory pathways to induce Mx gene expression for antiviral defence, which may contribute to their success in an anadromous life style.

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- **6. References**
- 540 [1] Haller O, Staeheli P, Schwemmle M, Kochs G. Mx GTPases: dynamin-like antiviral machines of innate
- 541 immunity. Trends Microbiol. 2015;23:154-63.
- 542 [2] Verhelst J, Hulpiau P, Saelens X. Mx proteins: antiviral gatekeepers that restrain the uninvited. Microbiol
- 543 Mol Biol Rev. 2013;77:551-66.
- 544 [3] Haller O, Gao S, von der Malsburg A, Daumke O, Kochs G. Dynamin-like MxA GTPase: structural
- insights into oligomerization and implications for antiviral activity. J Biol Chem. 2010;285:28419-24.
- 546 [4] Qi F, Yang A, Ambreen S, Bai X, Hou Y, Lu X. Birth and death of Mx genes and the presence/absence of
- 547 genes regulating Mx transcription are correlated with the diversity of anti-pathogenicity in vertebrate species.
- 548 Mol Genet Genomics. 2019;294:121-133.
- [5] Staeheli P, Yu YX, Grob R, Haller O. 1989. A double-stranded RNA-inducible fish gene homologous to
- the murine influenza virus resistance gene Mx. Mol. Cell. Biol. 9:3117–21.
- 551 [6] Trobridge GD, Leong JA. Characterization of a rainbow trout Mx gene. J Interferon Cytokine Res.
- 552 1995;15:691-702.
- 553 [7] Trobridge GD, Chiou PP, Leong JA. Cloning of the rainbow trout (*Oncorhynchus mykiss*) Mx2 and Mx3
- 554 cDNAs and characterization of trout Mx protein expression in salmon cells. J Virol. 1997;71:5304-11.
- [8] Robertsen B, Trobridge G, Leong JA. Molecular cloning of double-stranded RNA inducible Mx genes
- from Atlantic salmon (*Salmo salar* L.). Dev Comp Immunol. 1997;21:397-412.
- [9] Jensen V, Robertsen B. Cloning of an Mx cDNA from Atlantic halibut (*Hippoglossus hippoglossus*) and
- 558 characterization of Mx mRNA expression in response to double-stranded RNA or infectious pancreatic
- necrosis virus. J Interferon Cytokine Res. 2000;20:701-10.
- 560 [10] Lee JY, Hirono I, Aoki T. Cloning and analysis of expression of Mx cDNA in Japanese flounder,
- 561 Paralichthys olivaceus. Dev Comp Immunol. 2000;24:407-15.
- 562 [11] Altmann SM, Mellon MT, Johnson MC, Paw BH, Trede NS, Zon LI, Kim CH. Cloning and
- characterization of an Mx gene and its corresponding promoter from the zebrafish, *Danio rerio*. Dev Comp
- 564 Immunol. 2004;28:295-306.
- 565 [12] Zhang YB, Li Q, Gui JF. Differential expression of two Carassius auratus Mx genes in cultured CAB
- cells induced by grass carp hemorrhage virus and interferon. Immunogenetics. 2004;56:68-75.
- 567 [13] Plant KP, Thune RL. Cloning and characterisation of a channel catfish (*Ictalurus punctatus*) Mx gene.
- 568 Fish Shellfish Immunol, 2004;16:391-405.
- 569 [14] Wu YC, Chi SC. Cloning and analysis of antiviral activity of a barramundi (*Lates calcarifer*) Mx gene.
- 570 Fish Shellfish Immunol. 2007;23:97-108.
- 571 [15] Zenke K, Kim KH. Molecular cloning and expression analysis of three Mx isoforms of rock bream,
- 572 *Oplegnathus fasciatus*. Fish Shellfish Immunol. 2009;26:599-605.
- 573 [16] Li G, Zhang J, Sun Y, Wang H, Wang Y. The evolutionarily dynamic IFN-inducible GTPase proteins
- play conserved immune functions in vertebrates and cephalochordates. Mol Biol Evol. 2009;26:1619-30.
- 575 [17] Huang B, Huang WS, Nie P. Characterization of four Mx isoforms in the European eel, *Anguilla anguilla*.
- 576 Fish Shellfish Immunol. 2013;35:1048-54.

- 577 [18] Fernández-Trujillo MA, García-Rosado E, Alonso MC, Castro D, Álvarez MC, Béjar J. Mx1, Mx2 and
- 578 Mx3 proteins from the gilthead seabream (Sparus aurata) show in vitro antiviral activity against RNA and
- 579 DNA viruses. Mol Immunol. 2013;56:630-6.
- 580 [19] Alvarez-Torres D, Bejar J, Collet B, Alonso MC, Garcia-Rosado E. Structural and functional
- 581 characterization of the Senegalese sole (Solea senegalensis) Mx promoter. Fish Shellfish Immunol.
- 582 2013;35:1642-8.
- 583 [20] Novel P, Fernández-Trujillo MA, Gallardo-Gálvez JB, Cano I, Manchado M, Buonocore F, Randelli E,
- Scapigliati G, Alvarez MC, Béjar J. Two Mx genes identified in European sea bass (*Dicentrarchus labrax*)
- respond differently to VNNV infection. Vet Immunol Immunopathol. 2013;153:240-8.
- 586 [21] Saxena A, Belwal K, Chauhan A, Pande A. Interferon induced Mx protein from Indian snow trout
- 587 Schizothorax richardsonii (Gray) lacks critical functional features unlike its mammalian homologues. Comput
- 588 Biol Chem. 2018;73:31-40.
- 589 [22] Solbakken MH, Rise ML, Jakobsen KS, Jentoft S. Successive Losses of Central Immune Genes
- 590 Characterize the Gadiformes' Alternate Immunity. Genome Biol Evol. 2016;8:3508-3515.
- 591 [23] Caipang CM, Hirono I, Aoki T. In vitro inhibition of fish rhabdoviruses by Japanese flounder,
- 592 Paralichthys olivaceus Mx. Virology. 2003; 317:373–82
- 593 [24] Larsen R, Rokenes TP, Robertsen B. Inhibition of infectious pancreatic necrosis virus replication by
- 594 Atlantic salmon Mx1 protein. J. Virol. 2004; 78:7938–44
- 595 [25] Su J, Yang C, Zhu Z, Wang Y, Jang S, Liao L. Enhanced grass carp reovirus resistance of Mx-transgenic
- rare minnow (*Gobiocypris rarus*). Fish Shellfish Immunol. 2009;26:828-35.
- 597 [26] Peng L, Yang C, Su J. Protective roles of grass carp *Ctenopharyngodon Idella* Mx isoforms against grass
- 598 carp reovirus. PLoS One. 2012;7:e52142.
- 599 [27] Robertsen B, Greiner-Tollersrud L, Jørgensen LG. Analysis of the Atlantic salmon genome reveals a
- 600 cluster of Mx genes that respond more strongly to IFN gamma than to type I IFN. Dev Comp Immunol.
- 601 2019;90:80-89.
- 602 [28] Husain M, Bird S, van Zwieten R, Secombes CJ, Wang T. Cloning of the IL-1β3 gene and IL-1β4
- 603 pseudogene in salmonids uncovers a second type of IL-1β gene in teleost fish. Dev Comp Immunol.
- 604 2012;38:431-46.
- 605 [29] MordsteinM, Neugebauer E, Ditt V, Jessen B, Rieger T, et al. 2010. Lambda interferon renders epithelial
- 606 cells of the respiratory and gastrointestinal tracts resistant to viral infections. J. Virol. 84:5670–77
- 607 [30] Holzinger D, Jorns C, Stertz S, Boisson-Dupuis S, Thimme R, et al. 2007. Induction of MxA gene
- expression by influenza A virus requires type I or type III interferon signaling. J. Virol. 81:7776–85
- 609 [31] Simon A, Fah J, Haller O, Staeheli P. 1991. Interferon-regulated Mx genes are not responsive to
- 610 interleukin-1, tumor necrosis factor, and other cytokines. J. Virol. 65:968–71
- 611 [32] Wangkahart E, Secombes CJ, Wang T. Dissecting the immune pathways stimulated following injection
- of vaccination of rainbow trout (*Oncorhynchus mykiss*) against enteric redmouth disease (ERM). Fish Shellfish
- 613 Immunol. 2019;85:18-30.
- 614 [33] Altschul SF, Gish W, Miller W, Myers EW, Lipman DJ. Basic local alignment search tool. J Mol Biol
- 615 1990; 215:403–10.

- 616 [34] Wang T, Hu Y, Wangkahart E, Liu F, Wang A, Zahran E, Maisey KR, Liu M, Xu Q, Imarai M,
- 617 Secombes CJ. Interleukin (IL)-2 Is a Key Regulator of T Helper 1 and T Helper 2 Cytokine Expression in Fish:
- Functional Characterization of Two Divergent IL2 Paralogs in Salmonids. Front Immunol. 2018;9:1683.
- 619 [35] Wang B, Wangkahart E, Secombes CJ, Wang T. Insights into the evolution of the suppressors of cytokine
- signalling (SOCS) gene family in vertebrates. Mol Biol Evol. 2019 36:393-411.
- 621 [36] Gasteiger E, Hoogland C, Gattiker A, Duvaud S, Wilkins MR, Appel RD, et al. Protein identification and
- analysis tools on the ExPASy server. In: Walker JM, editor. The Proteomics Protocols Handbook. New York:
- 623 Humana Press; (2005). p. 571–607.
- 624 [37] Chenna R, Sugawara H, Koike T, Lopez R, Gibson TJ, Higgins DG, Thompson JD. Multiple sequence
- alignment with the Clustal series of programs. Nucleic Acids Res. 2003; 31:3497-500,
- 626 [38] Campanella JJ, Bitincka L, Smalley J. MatGAT: an application that generates similarity/identity matrices
- using protein or DNA sequences. BMC Bioinformatics. 2003; 4:29.
- 628 [39] Kumar S, Stecher G, Tamura K. MEGA7: Molecular Evolutionary Genetics Analysis Version 7.0 for
- 629 Bigger Datasets. Mol Biol Evol. 2016; 33:1870-4.
- 630 [40] Staeheli P, Haller O, Boll W, Lindenmann J, Weissmann C. 1986. Mx protein: constitutive expression in
- 3T3 cells transformed with cloned Mx cDNA confers selective resistance to influenza virus. Cell 44:147–58
- [41] Staeheli P, Sutcliffe JG. Identification of a second interferon-regulated murine Mx gene. Mol. Cell. Biol.
- 633 1988; 8:4524–28
- 634 [42] Louis A, Muffato M, Roest Crollius H. Genomicus: five genome browsers for comparative genomics in
- eukaryota. Nucleic Acids Res. 2013;41(Database issue):D700-5.
- 636 [43] Wang T, Diaz-Rosales P, Costa MM, Campbell S, Snow M, Collet B, Martin SA, Secombes CJ.
- Functional characterization of a nonmammalian IL-21: rainbow trout *Oncorhynchus mykiss* IL-21 upregulates
- the expression of the Th cell signature cytokines IFN-gamma, IL-10, and IL-22. J Immunol. 2011;186:708-21.
- 639 [44] Wang T, Monte MM, Huang W, Boudinot P, Martin SA, Secombes CJ. Identification of two FoxP3
- 640 genes in rainbow trout (Oncorhynchus mykiss) with differential induction patterns. Mol Immunol.
- 641 2010;47:2563-74.
- 642 [45] Wangkahart E, Scott C, Secombes CJ, Wang T. Re-examination of the rainbow trout (Oncorhynchus
- 643 mykiss) immune response to flagellin: Yersinia ruckeri flagellin is a potent activator of acute phase proteins,
- anti-microbial peptides and pro-inflammatory cytokines in vitro. Dev Comp Immunol. 2016;57:75-87.
- 645 [46] Ganassin RC, Bols NC. Development of a monocyte/macrophage-like cell line, RTS11, from rainbow
- trout spleen. Fish Shellfish Immunol. 1998; 8:457–476
- 647 [47] Wolf K, Quimby MC. Established eurythermic lines of fish cells in vitro. Science 1962; 135:1065
- 648 [48] Schirmer K, Chan AG, Greenberg BM, Dixon DG, Bols NC. Ability of 16 priority PAHs to be
- photocytotoxic to a cell line from the rainbow trout gill. Toxicology. 1998; 127:143–155
- 650 [49] Wang T, Huang W, Costa MM, Martin SA, Secombes CJ. Two copies of the genes encoding the subunits
- of putative interleukin (IL)-4/IL-13 receptors, IL-4Rα, IL-13Rα1 and IL-13Rα2, have been identified in
- 652 rainbow trout (Oncorhynchus mykiss) and have complex patterns of expression and modulation.
- 653 Immunogenetics. 2011;63:235-53.
- 654 [50] Hong S, Zou J, Crampe M, Peddie S, Scapigliati G, Bols N, Cunningham C, Secombes CJ. The
- production and bioactivity of rainbow trout (Oncorhynchus mykiss) recombinant IL-1 beta. Vet Immunol
- 656 Immunopathol. 2001;81:1-14.

- 657 [51] Costa MM, Maehr T, Diaz-Rosales P, Secombes CJ, Wang T. Bioactivity studies of rainbow trout
- 658 (Oncorhynchus mykiss) interleukin-6: effects on macrophage growth and antimicrobial peptide gene
- expression. Mol Immunol. 2011;48:1903-16.
- 660 [52] Hong S, Li R, Xu Q, Secombes CJ, Wang T. Two types of TNF-α exist in teleost fish: phylogeny,
- 661 expression, and bioactivity analysis of type-II TNF-α3 in rainbow trout *Oncorhynchus mykiss*. J Immunol.
- 662 2013;191:5959-72.
- 663 [53] Husain M, Martin SA, Wang T. Identification and characterisation of the IL-27 p28 subunits in fish:
- 664 Cloning and comparative expression analysis of two p28 paralogues in Atlantic salmon Salmo salar. Fish
- 665 Shellfish Immunol. 2014. 41:102-12.
- 666 [54] Wangkahart E, Secombes CJ, Wang T. Studies on the Use of Flagellin as an Immunostimulant and
- Vaccine Adjuvant in Fish Aquaculture. Front Immunol. 2019. 9:3054.
- 668 [55] Gao S, von der Malsburg A, Paeschke S, Behlke J, Haller O, Kochs G, Daumke O. Structural basis of
- oligomerization in the stalk region of dynamin-like MxA. Nature. 2010. 465(7297):502-6.
- 670 [56] Berthelot C, Brunet F, Chalopin D, Juanchich A, Bernard M, Noël B, Bento P, Da Silva C, Labadie K,
- Alberti A, Aury JM, Louis A, Dehais P, Bardou P, Montfort J, Klopp C, Cabau C, Gaspin C, Thorgaard GH,
- Boussaha M, Quillet E, Guyomard R, Galiana D, Bobe J, Volff JN, Genêt C, Wincker P, Jaillon O, Roest
- 673 Crollius H, Guiguen Y. The rainbow trout genome provides novel insights into evolution after whole-genome
- duplication in vertebrates. Nat Commun. 2014. 5:3657.
- [57] Lien S, Koop BF, Sandve SR, Miller JR, Kent MP, Nome T, Hvidsten TR, Leong JS, Minkley DR, Zimin
- A, Grammes F, Grove H, Gjuvsland A, Walenz B, Hermansen RA, von Schalburg K, Rondeau EB, Di
- 677 Genova A, Samy JK, Olav Vik J, Vigeland MD, Caler L, Grimholt U, Jentoft S, Våge DI, de Jong P, Moen T,
- Baranski M, Palti Y, Smith DR, Yorke JA, Nederbragt AJ, Tooming-Klunderud A, Jakobsen KS, Jiang X, Fan
- D, Hu Y, Liberles DA, Vidal R, Iturra P, Jones SJ, Jonassen I, Maass A, Omholt SW, Davidson WS. The
- Atlantic salmon genome provides insights into rediploidization. Nature. 2016; 533(7602):200-5.
- 681 [58] Staeheli P, Haller O. Human MX2/MxB: a Potent Interferon-Induced Postentry Inhibitor of
- 682 Herpesviruses and HIV-1. J Virol. 2018; 92(24). pii:e00709-18.
- 683 [59] Goujon C, Moncorgé O, Bauby H, Doyle T, Barclay WS, Malim MH. Transfer of the amino-terminal
- nuclear envelope targeting domain of human MX2 converts MX1 into an HIV-1 resistance factor. J Virol.
- 685 2014; 88:9017-26.

- 686 [60] Zurcher T, Pavlovic J, Staeheli P. Mouse Mx2 protein inhibits vesicular stomatitis virus but not influenza
- 687 virus. Virology. 1992; 187:796–800
- 688 [61] Stuempfig ND, Seroy J. Viral Gastroenteritis. 2019; StatPearls
- [62] Secombes CJ, Zou J. Evolution of Interferons and Interferon Receptors. Front Immunol. 2017; 8:209.

Table 1. Summary of Mx gene family in salmonids and pike

Common name Ger		Chromosome	Location	Genebank accession number (mRNA/protein)			
Rainbow trout	Mx1	NC_035093.1 (Ch17)	54,895,743 -> 54,907,262	NM_001171901.1			
Rainbow trout	Mx2	NC_035093.1 (Ch17)	54,827,385→54,839,851	NM_001124751.1			
Rainbow trout	Mx3	NC_035093.1 (Ch17)	54,879,332 -> 54,883,908	XM_021569609.1			
Rainbow trout	Mx4	NC_035093.1 (Ch17)	54,848,974 -> 54,863,639	MK301134			
Rainbow trout	Mx5	NC_035079.1 (Ch3)	82,015,213 -> 81,992,259	MK301135			
Rainbow trout	Mx6	NC_035079.1 (Ch3)	82,029,332 82,045,373	MK301136			
Rainbow trout	Mx7	NC_035087.1 (Ch11)	76,228,384→76,215,926	MK301137			
Rainbow trout	Mx8	NC_035087.1 (Ch11)	76,240,419→76,420,666	MK301138			
Rainbow trout	Mx9	NC_035100.1 (Ch24)	21,286,705 -> 21,268,999	MK301139			
Atlantic salmon	Mx1	NC_027311(Ch12)	66,798,275→66,829,177	NM_001123690/NP_001117162			
Atlantic salmon	Mx2	NC_027311(Ch12)	66,776,028→66,803,979	NM_001139918/NP_001133390			
Atlantic salmon	Mx3	NC_027311(Ch12)	66,816,288→66,829,177	NM_001123675/NP_001117147			
Atlantic salmon	Mx4	NC_027324(Ch25)	47,088,993→47,121,652	XM_014174614/XP_014030089			
Atlantic salmon	Mx5	NC_027324(Ch25)	47,228,437→47,217,827	XM_014174615/XP_014030090			
Atlantic salmon	Mx6	NC_027324(Ch25)	47,161,992→47,139,132	XM_014174616/XP_014030091			
Atlantic salmon	Mx7	NC_027324(Ch25)	47,193,272→47,175,785	XM_014174617/XP_014030092			
Atlantic salmon	Mx8	NC_027324(Ch25)	47,243,602→47,262,616	XM_014174618/XP_014030093			
Atlantic salmon	Mx9	NC_027308(Ch9)	117,838,750→117,853,816	XM_014214722/XP_014070197			
Atlantic salmon	Mx10	NC_027314(Ch15)	5,299,091 -> 5,292,439	XM_014143485/XP_013998960			
Chinook salmon	Mx1	NW_020142590	72,571→83,518	XM_024415949/XP_024271717			
Chinook salmon	Mx2	NW_020142590	17,032→35,480	XM_024415950/XP_024271718			
Chinook salmon	Mx3	NW_020142590	56,817→83,518	XM_024415946/XP_024271714			
Chinook salmon	Mx4	NW_020133776	172→6,566	XM_024410424/XP_024266192			
Chinook salmon	Mx5	NC_037108(Ch12)	2,001,377 -> 2,013,759	XM_024438118/XP_024293886			
Chinook salmon	Mx6	NC_037110(Ch14)	41,090,466 -> 41,103,747	XM_024445373/XP_024301141			
Coho salmon	Mx1	NC_034174(Ch1)	46,664,816→46,675,808	LOC109896993			
Coho salmon	Mx2	NC_034174(Ch1)	46,587,607→46,621,678	XM_020468497/XP_020324086			
Coho salmon	Mx3	NC_034174(Ch1)	46,651,400→46,656,202	XM_020491468/XP_020347057			
Coho salmon	Mx4	NW_018090236	57,121→68,549	GDQG01031501/ /Q6PW23			
Coho salmon	Mx5	NC_034181(Ch8)	66,940,031 → 66,941,209	XM_020488627/XP_020344216			
Coho salmon	Mx6	NC_034191(Ch18)	53,794,477 -> 53,822,277	XM_020508491/XP_020364080			
Arctic charr	Mx1	NC_036838(Ch1)	44,797,136 44,802,044	XM_023993827/XP_023849595			
Arctic charr	Mx2	NC_036838(Ch1)	44,762,649 44,772,138	XM_023993825/XP_023849593			
Arctic charr	Mx3	NW_019943275	202,142-225,309	XM_024139811/XP_023995579			
Arctic charr	Mx4	NW_019943275	93,732→108,405	XM_024139809/XP_023995577			
Arctic charr	Mx5	NW_019943275	87,444→170,634	XM_024139810/XP_023995578			
Arctic charr	Mx6	NW_019945020	48,231→54,383	XM_024143207/XP_023998975			
Arctic charr	Mx7	NW_019945020	11,050→28,797	XM_024143206/XP_023998974			
Arctic charr	Mx8	NW_019946381	2,678→17,616	XM_024144359/XP_024000127			
Arctic charr	Mx9	NW_019942645	359,971→369,223	XM_024136430/XP_023992198			
Arctic charr	Mx10	NW_019942645	378,229→388,058	XM_024136431/XP_023992199			
Pike	Mx1	NC_025984(Ch17)	25,113,053 -> 25,132,843	XM_013138351/XP_012993805			
Pike	Mx2	NC_025980(Ch13)	25,725,708—25,740,882	ENSELUG00000023570/ENSELUT 00000043341			
Pike	Mx3	NC_025980(Ch13)	25,694,138→25,702,004	ENSELUG00000023626/ENSELUT 00000036437			

Table 2. Comparison of trout Mx aa sequence identities to Mx from other salmonids, spotted gar and mammals. The amino acid number for each sequence is also shown. Only full-length aa sequences were included in the analysis.

			SMG1				SMG2		SMG3		SMG4
		No. of aa	Trout Mx1	Trout Mx2	Trout Mx3	Trout Mx4	Trout Mx5	Trout Mx6	Trout Mx7	Frout Mx8	Trout Mx9
	Trout Mx1	621	100.0	86.5	96.3	86.5	46.2	46.3	47.9	46.4	47.7
	Trout-Mx2	635	86.5	100.0	86.3	98.4	45.6	45.0	47.0	45.1	47.3
	Trout Mx3	623	96.3	86.3	100.0	86.0	45.9	46.0	48.1	46.6	47.9
	Trout Mx4	635	86.5	98.4	86.0	100.0	45.8	45.5	47.0	45.2	47.0
	Atlantic Mx1	623	95.3	85.8	97.3	85.4	45.7	45.9	47.9	46.4	47.4
	Atlantic Mx2	638	85.4	94.0	85.4	93.9	45.4	44.8	46.4	44.7	46.7
	Atlantic Mx3	623	96.3	86.5	95.8	86.6	46.5	46.0	48.1	46.7	48.3
	Chinook Mx1	621	98.2	86.8	96.0	86.5	46.2	46.2	47.9	46.7	47.9
	Chinook Mx2	633	83.7	95.0	83.4	95.1	43.6	43.5	45.4	43.5	45.6
	Chinook Mx3	623	95.2	86.0	97.8	86.2	46.2	46.2	48.1	46.3	47.8
	Coho-Mx2	648	84.3	95.2	84.1	95.7	44.7	44.3	45.8	44.2	45.9
	Coho Mx3	623	96.8	85.5	97.8	85.4	46.0	46.2	47.9	46.3	47.8
<u>G</u>	Charr Mx1	623	96.0	86.6	97.9	86.2	45.7	46.0	48.1	46.1	47.6
SMG1	Charr Mx2	638	85.6	94.8	85.6	95.1	45.4	44.9	47.0	45.3	47.3
	Trout Mx5	614	46.2	45.6	45.9	45.8	100.0	93.6	61.2	69.4	51.4
	Trout Mx6	606	46.3	45.0	46.0	45.5	93.6	100.0	61.4	69.7	50.8
	Atlantic Mx4	606	45.6	45.4	45.2	45.6	86.0	88.3	62.2	70.1	49.7
	Atlantic Mx5	608	45.4	44.7	45.5	44.8	90.1	88.8	62.7	70.8	50.2
	Atlantic Mx6	627	45.3	44.4	45.0	44.7	83.9	84.4	61.3	67.9	49.5
	Atlantic Mx7	603	46.4	45.0	46.1	45.5	86.6	87.1	61.8	69.9	49.8
	Atlantic Mx8	607	47.0	45.5	46.7	45.6	87.0	90.0	62.5	71.2	50.6
	Coho Mx4	614	46.0	45.2	45.7	45.3	95.1	91.7	61.1	69.9	51.6
G2	Charr Mx4	607	46.5	45.9	46.2	46.1	90.9	89.8	61.2	69.3	50.2
SMG2	Charr Mx5	612	45.1	43.8	45.2	44.2	82.7	85.0	60.6	68.2	49.7
	Trout Mx7	613	47.9	47.0	48.1	47.0	61.2	61.4	100.0	68.6	49.1
	Trout Mx8	608	46.4	45.1	46.6	45.2	69.4	69.7	68.6	100.0	49.8
G3	Charr Mx9	549	42.3	41.7	42.4	41.5	53.6	53.4	80.7	57.8	44.5
SMG3	Charr Mx10	608	46.1	45.2	46.4	45.4	69.3	69.6	68.7	93.8	49.8
	Trout Mx9	640	47.7	47.3	47.9	47.0	51.4	50.8	49.1	49.8	100.0
	Atlantic Mx9	642	48.1	47.6	48.2	47.1	51.6	50.8	49.5	49.5	94.9
25	Chinook Mx6	638	47.4	47.2	47.8	46.9	51.1	50.5	49.3	49.4	97.0
SMG4	Coho Mx6	646	43.2	42.3	43.2	42.1	45.2	44.8	45.4	43.6	80.2
	Spotted gar Mx1	619	73.9	71.3	74.8	71.3	46.6	47.1	50.8	47.0	48.1
,.	Spotted gar Mx2	684	48.5	47.8	48.5	47.8	38.7	38.6	41.2	39.2	39.4
Gar	Spotted gar Mx3	616	47.6	44.3	47.0	44.4	41.3	40.8	41.1	40.8	37.3
als	Human MxA	670	52.6	52.1	52.4	52.0	40.5	40.4	42.1	41.2	43.1
l mm	Mouse-Mx1	631	51.8	51.2	52.0	51.6	40.4	40.4	42.1	40.9	43.3
Mammals	Cow Mx1	648	51.8	52.0	52.1	51.7	39.5	40.7	42.7	41.5	42.8

702 Figure legend

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- **Fig. 1. Amino acid multiple alignment of rainbow trout Mx family.** The multiple alignment was produced using ClustalW, and conserved amino acids shaded using BOXSHADE (version 3.21). Human MxA and MxB were included in the alignment for comparison. The N-terminal GTPase domain and C-terminal GTPase effector domain are indicated above the alignment. The conserved tripartite GTP-binding motif (GDXXSGKS, DLPG, and TKPD) and a dynamin signature (LPRXXGXXTR) in the GTPase domain, and leucine residues that form leucine zipper folds in the GTPase effector domain are in red. The four α-helices and the four loops connecting them are shown under the alignment as defined by Gao et al. [55]. The potential nuclear localisation signal (KKRKR) in trout Mx2 and 4, the four lysine residues of human MxA in L4 are in blue.
- Fig. 2 Phylogenetic tree (A) and chromosome localisation (B) of salmonid Mx genes. A. The phylogenetic tree was constructed using a multiple alignment of salmonid and pike Mx aa sequences and the neighbour-joining method within the MEGA7.0 program. The evolutionary distances were computed using the JTT matrix-based method with all ambiguous positions removed for each sequence pair. The percentage (>50%) of replicate trees in which the associated taxa clustered together in the bootstrap test (10,000 replicates) is shown next to the branches. The accession number for each sequence is given in Table 2. Four salmonid Mx groups (SMG)1-4 are indicated on the right. B. the chromosome localisation of Mx genes in salmonids and pike.
- 719 Fig. 3 Phylogenetic tree analysis of bony fish Mx. The phylogenetic tree was constructed using amino acid 720 multiple alignments of Mx from selected teleosts and mammals, and the neighbour-joining method within the 721 MEGA7.0 program. The evolutionary distances were computed using the JTT matrix-based method with all 722 ambiguous positions removed for each sequence pair. Node values represent percent bootstrap confidence derived from 10,000 replications. The accession number for each sequence is given after the species and 723 724 molecule names. The salmonid clades are highlighted and condensed under the name of SMG (salmonid Mx 725 group)1-4, which share the same topologies as in Fig. 2. The root bootstrap values of mammalian Mx and 726 teleost Mx group 1-3 are highlighted in red with the tentative groupings indicated on the right of the tree.
- Fig. 4. Synteny analysis of Mx loci in bony fish and human. The synteny was predicted using the Genomicus program [42] or information extracted from recently released reference genomes at NCBI.
- Fig. 5. Comparison of gene organisation of the Mx gene family in rainbow trout, other bony fish and humans. Boxes represent exons, and lines between exons represent introns. The black and white boxes represent non-coding and amino acid (aa) coding regions, respectively. The sizes (bp) of each exon are numbered in the boxes. The gene organization of rainbow trout Mx genes was predicted using the Splign program based on the sequence information from Table 1 and Figures S1–S6 in Supplementary Material. The information of other species was extracted from recent released reference genomes at NCBI.

- 735 Figure 6. Transcript expression of rainbow trout Mx gene family in tissues and cell lines. The expression
- level of Mx1-4 (A), Mx5-6 (B), Mx7-8 (C) and Mx9 (D) was determined by RT-qPCR in 17 tissues from six
- 737 fish and four replicates of each cell line. The transcript level was calculated using a serial dilution of
- 738 references that contained equal molar amounts of the probes for each gene and was normalized against the
- expression level of EF-1 α . The results are presented as the average +SEM.
- 740 Fig. 7. The expression ontogeny of rainbow trout Mx gene family. cDNA samples were prepared from
- eyed-eggs, immediately post-hatch, pre-first feeding fry or fry 3 weeks after first feeding. Six independent
- samples for each developmental stage were prepared for real-time quantification of gene expression as
- 743 described in Fig. 6. The results are presented as the average + SEM. Different letters over bars indicate
- significant differences ($p \le 0.05$, one way-ANOVA).
- 745 Fig. 8. Modulation of Mx gene expression in vivo by poly IC. Rainbow trout were injected ip with 1 mg
- poly IC in 0.2 ml PBS or 0.2 ml PBS as control. The spleen, head kidney (HK), gills and intestine were taken
- at 6 h and 24 h post injection. The quantification of Mx gene expression was as described in Fig. 6. The
- relative expression is shown, where the average expression level in the control fish at 6 h in each tissue was
- defined as 1. The results are presented as the mean + SEM of six fish. Different letters over bars in the same
- tissue indicate significant differences ($p \le 0.05$, one way-ANOVA).
- 751 Fig. 9. Modulation of proinflammatory cytokine gene expression in vivo by poly IC. Rainbow trout were
- 752 injected ip with 1 mg poly IC in 0.2 ml PBS or 0.2 ml PBS as control. The spleen, head kidney (HK), gills and
- 753 intestine were taken at 6 h and 24 h post injection. The quantification of gene expression was as described in
- Fig. 6. The relative expression is shown, where the average expression level in the control fish at 6 h in each
- 755 tissue was defined as 1. The results are presented as the mean + SEM of six fish. Different letters over bars in
- 756 the same tissue indicate significant differences ($p \le 0.05$, one way-ANOVA).
- 757 Fig. 10. Modulation of Mx gene expression in RTS-11 cells by poly IC and LPS. Overnight culture of
- 758 RTS-11 cells were stimulated with poly IC (50 µg/ml), LPS (25 µg/ml), or medium as control for 4h, 8 h and
- 759 24 h, and the expression of trout Mx genes was quantified by RT-qPCR as described in Fig. 6. The data are
- presented as the mean (+SEM, N=4) fold change calculated as the average expression level of stimulated
- samples divided by that of time-matched controls. The relative significance of a LSD post hoc test after a
- 762 significant one-way ANOVA between the stimulated and time-matched controls is shown above the bars as *
- 763 $p \le 0.05$; ** $p \le 0.01$ and *** $p \le 0.001$.
- 764 Fig. 11. Modulation of trout Mx expression in RTS-11 cells by pro-inflammatory cytokines. Overnight
- culture of RTS-11 cells were stimulated with IFNγ (20 ng/ml), IFN-a (25 ng/ml), IL-1β (25 ng/ml), IL-6 (100
- ng/ml), TNFα (50 ng/ml), or medium as control for 4h, 8 h and 24 h, and the expression of trout Mx genes
- was quantified by RT-qPCR as described in Fig. 6. The data are presented as the mean (+SEM, N=4) fold
- change, calculated as the average expression level of stimulated samples divided by that of time-matched

769	controls. The relative significance of a LSD post hoc test after a significant one-way ANOVA between the
770	stimulated and time-matched controls is shown above the bars as * $p \le 0.05$; ** $p \le 0.01$ and *** $p \le 0.001$.

- Fig. 12. Modulation of trout Mx expression in RTG-2 and RTGill cell lines by IFNs. Overnight cultures of cells were stimulated with IFN γ (20 ng/ml), IFN-a (25 ng/ml), or medium as control for 4h, and the expression of trout Mx genes was quantified by RT-qPCR as described in Fig. 6. The data are presented as the mean (+SEM, N=4) fold change, calculated as the average expression level of stimulated samples divided by that of controls. The relative significance of a LSD post hoc test after a significant one-way ANOVA between the stimulated and controls is shown above the bars as * p \le 0.05; **p \le 0.01 and *** p \le 0.001. The line-connected groups are significantly different.
- **Fig. 13.** Hypothetical evolutionary pathways of teleost Mx gene family. Three Mx loci (Mx1-3) were present on two chromosomes in ancestral 2R actinopterygians. 3R WGD is expected to have produced 6 Mx loci on four chromosomes that were retained in a lineage specific manner to to give rise to the three extant teleost Mx groups. The ancestral vertebrates that evolved into the tetrapod lineage appear to have possessed a cognate Mx locus of gar Ch3. Arrow heads indicate ancestral Mx genes. Representative chromosomal loci are shown.

Table 2. Comparison of trout Mx aa sequence identities to Mx from other salmonids, spotted gar and mammals. The amino acid number for each sequence is also shown. Only full-length aa

			SMG1				SM	G2	SMG3		SMG4
		No. of aa	Trout Mx1	Trout Mx2	Trout Mx3	Trout Mx4	Trout Mx5	Trout Mx6	Trout Mx7	Trout Mx8	Trout Mx9
	Trout Mx1	621	100.0	86.5	96.3	86.5	46.2	46.3	47.9	46.4	47.7
	Trout-Mx2	635	86.5	100.0	86.3	98.4	45.6	45.0	47.0	45.1	47.3
	Trout Mx3	623	96.3	86.3	100.0	86.0	45.9	46.0	48.1	46.6	47.9
	Trout Mx4	635	86.5	98.4	86.0	100.0	45.8	45.5	47.0	45.2	47.0
	Atlantic Mx1	623	95.3	85.8	97.3	85.4	45.7	45.9	47.9	46.4	47.4
	Atlantic Mx2	638	85.4	94.0	85.4	93.9	45.4	44.8	46.4	44.7	46.7
SMG1	Atlantic Mx3	623	96.3	86.5	95.8	86.6	46.5	46.0	48.1	46.7	48.3
$\mathbf{S}\mathbf{M}$	Chinook Mx1	621	98.2	86.8	96.0	86.5	46.2	46.2	47.9	46.7	47.9
	Chinook Mx2	633	83.7	95.0	83.4	95.1	43.6	43.5	45.4	43.5	45.6
	Chinook Mx3	623	95.2	86.0	97.8	86.2	46.2	46.2	48.1	46.3	47.8
	Coho-Mx2	648	84.3	95.2	84.1	95.7	44.7	44.3	45.8	44.2	45.9
	Coho Mx3	623	96.8	85.5	97.8	85.4	46.0	46.2	47.9	46.3	47.8
	Charr Mx1	623	96.0	86.6	97.9	86.2	45.7	46.0	48.1	46.1	47.6
	Charr Mx2	638	85.6	94.8	85.6	95.1	45.4	44.9	47.0	45.3	47.3
	Trout Mx5	614	46.2	45.6	45.9	45.8	100.0	93.6	61.2	69.4	51.4
	Trout Mx6	606	46.3	45.0	46.0	45.5	93.6	100.0	61.4	69.7	50.8
	Atlantic Mx4	606	45.6	45.4	45.2	45.6	86.0	88.3	62.2	70.1	49.7
	Atlantic Mx5	608	45.4	44.7	45.5	44.8	90.1	88.8	62.7	70.8	50.2
SMG2	Atlantic Mx6	627	45.3	44.4	45.0	44.7	83.9	84.4	61.3	67.9	49.5
$\mathbf{S}\mathbf{M}$	Atlantic Mx7	603	46.4	45.0	46.1	45.5	86.6	87.1	61.8	69.9	49.8
	Atlantic Mx8	607	47.0	45.5	46.7	45.6	87.0	90.0	62.5	71.2	50.6
	Coho Mx4	614	46.0	45.2	45.7	45.3	95.1	91.7	61.1	69.9	51.6
	Charr Mx4	607	46.5	45.9	46.2	46.1	90.9	89.8	61.2	69.3	50.2
	Charr Mx5	612	45.1	43.8	45.2	44.2	82.7	85.0	60.6	68.2	49.7
	Trout Mx7	613	47.9	47.0	48.1	47.0	61.2	61.4	100.0	68.6	49.1
SMG3	Trout Mx8	608	46.4	45.1	46.6	45.2	69.4	69.7	68.6	100.0	49.8
$\mathbf{S}\mathbf{M}$	Charr Mx9	549	42.3	41.7	42.4	41.5	53.6	53.4	80.7	57.8	44.5
	Charr Mx10	608	46.1	45.2	46.4	45.4	69.3	69.6	68.7	93.8	49.8
	Trout Mx9	640	47.7	47.3	47.9	47.0	51.4	50.8	49.1	49.8	100.0
SMG4	Atlantic Mx9	642	48.1	47.6	48.2	47.1	51.6	50.8	49.5	49.5	94.9
$\mathbf{S}\mathbf{W}$	Chinook Mx6	638	47.4	47.2	47.8	46.9	51.1	50.5	49.3	49.4	97.0
	Coho Mx6	646	43.2	42.3	43.2	42.1	45.2	44.8	45.4	43.6	80.2
•.	Gar Mx1	619	73.9	71.3	74.8	71.3	46.6	47.1	50.8	47.0	48.1
Gar	Gar Mx2	684	48.5	47.8	48.5	47.8	38.7	38.6	41.2	39.2	39.4
	Gar Mx3	616	47.6	44.3	47.0	44.4	41.3	40.8	41.1	40.8	37.3
ıals	Human MxA	670	52.6	52.1	52.4	52.0	40.5	40.4	42.1	41.2	43.1
Mammals	Mouse-Mx1	631	51.8	51.2	52.0	51.6	40.4	40.4	42.1	40.9	43.3
	Cow Mx1	648	51.8	52.0	52.1	51.7	39.5	40.7	42.7	41.5	42.8

sequences were included in the analysis.

```
MNNTLNQHYEEKVRPCIDLIDSLRSLGVEKDLALPAIAVIGDQSSGKSSVLEALSGVALPRGSGIVTRCPLELKMKRKKEGEEWHGKISYQDHE
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 Trout Mx2
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                                                  ------MNNTLNQHYEEKVRPCIDLIDSLRSLGVEKDLALPAIAVIGDQSSGKSSVLEALSGVALPRGSGIVTRCPLELKMKRKREGEEWHGKISYQDHE
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-MSHD-DGPRTFQDQLAEKIRPFIDLVDDMRSIGIDKELPLPTIAVVGDQSSGKSSVLETLSGVALPRGTGIVTRCPLLLKLCNDR-TVKWDAVISYGGKI
Trout Mx3
                          1
Trout Mx4
Trout Mx5
Trout Mx6
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Trout Mx7
 Trout Mx8
Trout Mx9
                          1
Human MxA
Human MxB
                      Dynamin-like GTPase Domain
95 EEIEDPSDVEKKIREAQDEMAGVGVGISDDLISLEIGSPDVPDLTLIDLPGIARVAVKGQPENIGEQIKRLIRKFIMKQETISLVVVPCNVDIATTEALKMAQEVDPEGE
Trout Mxl
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EEIEDPSDVEKKIREAQDEMAGVGVGISDDLISLEIGSPDVPDLTLIDLPGIARVAVKGQPENIGEQIKRLIRKFINKQETINLVVVPCNVDIATTEALQMAQEVDPEGE
 Trout Mx2
Trout Mx3
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Trout Mx4
Trout Mx5
Trout Mx6
Trout Mx7
Trout Mx8 100 FEFDDREEVARHVEGAQNELAGGGGCEDLITLKIKSSTVCDLITLDEGTARVPVEGGEDLIGAGIKSLIMKYISKKKTINLVVPCVVDLATTEALKMAGKVDPEGT
Trout Mx8 100 FEFDDREEVARHVEGAQNELAGGGGCEDLITLKIKSSTVCDLSLIDLPGTARVPVFGQPEDLGQIKSLIMKYISKKKTINLVVPCVDLATTEALKMAGKVDPEGA
Human MxA 131 IEISDASEVEKEINKAQNAIAGEGMGISHELITLEISSRDVPDLTLIDLPGTRVAVGNQPADLGYKIKTLIKKYIQRQETISLVVVPSNVDIATTEALSMAQEVDPEGD
Human MxB 179 LELQDPGQVEKEIHKAQNVMAGNGRGISHELISLEITSPEVPDLTIIDLPGTRVAVDNQPRDIGLQIKALIKKYIQRQETINLVVVPCNVDIATTEALSMAHEVDPEGD
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Trout Mx2 205
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Trout Mx5
Trout. Mx6 209
Trout. Mx7
 Trout Mx8 210 RTLAILTKPDLIDKGTEKDVLEIVRNKTLPLNMGYVIVKCRGØKQIDDKMSIAQALEEELDFFQDHEHFKSLLLEERATTKHLATKLTYTLVNHIKKSLPDMSNQIKKQL
                    220 RTLAILTKPDLVDKGAEPDILKIVNGQVVHLNKGYIIVKCRGQNDINQKISLADATRLEMEFFKNHHHFSPLLEQNKVTTQCLATKLTQDLVDHIKTSLPYLTDQIREHL
241 RTIGILTKPDLVDKGTEDKVVDVVRNLVFHLKKGYMIVKCRGQQEIQDQLSLSEALQREKIFFENHPYFRDLLEEGKATVPCLAEKLTSELITHICKSLPLLENQIKETH
 Trout Mx9 220
Human MxB 288 RTIGILTKPDLMDRGTEKSVMNVVRNLTYPLKKGYMIVKCRGQQEITNRLSLAEATKKEITFFQTHPYFRVLLEEGSATVPRLAERLTTELIMHIQKSLPLLEGQIRESH
Trout Mxl 315 SETHAELERYGTGPPEDSAERLYFLIDKVTAFTQDAINLSTGEEMKSGVRLNVFSTLRKEFGKWKLHLERSGEIFNQRIEGEVDDYEKTYRGRELPGFINYKTFEVMVKD
Trout Mx2
                             EETRTALEKCGTGPPEDPKERLYFLIDKVTLFTQDAINLSTGEELKSG-DINVFSTLRTEFGKWKAYVDRSGKNFNKKIEKEVADYEKRYRGRELPGFINYKTFEVIVKD
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                                                                                                                            -VEENLFVLMRKEFTQWMKCLENDKSNYHKVVQQVVDEYDQEHRGSELPGFSNYRVFQHVVQK
-VEENLFELMRKEFTEWMECLKNAKSHYHEVVQQVVDEYDQEHRGSELPGFSNYRVFQHVVQK
-NDENLFINMQNIFAKWFEKLGHSRAGYHKMTQDVVNEFDQKHRGRELPGFNNYTLFESVVQK
Trout Mx5 319 GEVKNSLSKLEGGPPLEPEEKRKYLIQVITDFNEQITQLSKGDII-
Trout Mx6 319 GEVKHSLSKLEGGPPLEPEEKRKYLIQVITDFNEQITQLSKGDII-
Trout Mx7 318 WVYQTELTKYEGGPPVDPVGKRKYLIEVIKQFNYKIDQLCRGELK-
Trout Mx8 320 WNVRKALVECEGGPPSDLAERKEFLIGIITEFNEKITRLSTGDNT
                                                                                                                             -VEENLFVLMRSEFADWMKSLQNAKPNYHEVVQQVVDEYDLKHRGSELPGFTNYMEFKRVVQR
                    330 ETVKTELKKYSTGPPLERKKMGPYLTERLIDFIEKIHELCRIGNS---SEKNLHTCLRPVFQQWDSYLSNTKGSFLNKVAAMIKNYDKEHRGRELMTFSDYCVYEHAVQK
351 QRITEELQKYGVDIPEDENEKMFFLIDKVNAFNQDITALMQGEETVGEEDIRLFTRLRHEFHKWSTIIENNFQEGHKILSRKIQKFENQYRGRELPGFVNYRTFETIVKQ
Human MxB 398 QKATEELRRCGADIPSQEADKMFFLIEKIKMFNQDIEKLVEGEEVVRENETRLYNKIREDFKNWVGILATNTQKVKNIIHEEVEKYEKQYRGKELLGFVNYKTFEIIVHQ
Trout Mx3 425
                            QIKQLEEPAVKKLKEISDAVRKVFLLLAQSSFTGFPNLLKSAKTKIEAIKQVNESTAESMLRTQFKMEMIVYTQDSTYSHSLSERKREEEDD-
 Trout Mx4 424
                            QIKQLEEPAVKKLKELSDAARKAFILLAQNSFTGFPILLKTAKTKIETIKQEKESTAESTLRTQFKMELIVYTQDSTYSSSLKKRKREEEELEEGELVKNTLGSQKGFSV
TTOUL MX4 424 QIRQLEEPAVKLIKELSDAAKRAFILLAQNSFTGFFILLKTAKTKLETIRQEKESTAESTLRTQFKMELIVYTQDSTYSSSLKRKREEEELEEGELV
TTOUL MX5 426 LVAELKRPAMSTLQKIRDMVQKQFDHLSSESFKNYPYLHLVSKKNIETIQEKQSNIVKERIVEQFEMEMQVYTQDEIFNKVMLEAKSHLLEE—
TTOUL MX6 426 LVAELKRPAMSTLQTIRDMVQKQFDHLSSESFKNYPYLHLVSKKNIETIQEKQSNIVKERIVEQFEMEMQVYTQDEIFNKQSRK————
TTOUL MX7 425 LVGELKNPAMDTLQKIKDLVQKHFFVVSKSSFENYPCLQRFSMTNIDDIQKQQLTTVMDRIEEQFEMEM—YTQDEIFARTLTPAQKET————
TTOUL MX8 427 LVAKLREPAMMTLQKIREMVHTQFVNLSKVSFENFPYLQHVSMKNIENIQEWQSNIVMKRIEEQFQMEMQVYTQDEIFFETLNPE————
TTOUL MX9 437 HILGLQEPALDVLKAIGGMVQAEFRNVCEACFKSYPQLRSMALSKIDEIQTKQETKVEKRIKEYINMERLVYTQDSIFIKGLKDHKAQFKEAIEEEH-
HUMBAN MX9 430 VIOLIVEDALAGTIVYDMYRLAFTDVSIKNFEEFFNLHRTAKSKLEDIRAEQEREGEKLIRLHFQMEQIVYCQDQVYRGALQKVREKELEEEKKKK-
Human MxB 508 YIQQLVEPALSMLQKAMEIIQQAFINVAKKHFGEFFNLNQTVQSTIEDIKVKHTAKAENMIQLQFRMEQMVFCQDQIYSVVLKKVREEIFNPLGTPS
                                                                                                                                                                                                                                                    -ONM
                                                                                              L3
α2 helix

GTPase effector domain

Trout Mx1 520 TEIRSTIFSTDNHATLQEMMLHLKSYYWISSQRLADQIPMVIRYLVLQEFASQLQREMLQTLQEKDNIEQLLKEDIDIGSKRAALQSKLKRLMKARSYLVEF
Trout Mx2 534 VSVRSTVNGLDTHATLREMMLHLKSYYHTASQRLADQIPMVIRYLVLQEFASQLQREMLQTLQEKDNIEQLLKEDIDIGSKRAALQSKLKRLMKAHNYLVEF
Trout Mx3 522 PKIRSTIFSTDNHATLQEMMLHLKSYYRISSQRLADQIPMVIRYLVLQEFASQLQREMLQTLQEKDNIEQLLKEDFDIGSKRAALQNKLKRLMKARSYLVEF
Trout Mx3 522 PKLRSTIFSTDNHATLGEMMLHLKSYYRLSSQRLADQLPMVLRYLVLGEFASQLQREMLQTLGEKDNLEQLLKEDFDIGSKRAALQNKLKRLMKARSYLVEF-----
Trout Mx4 534 VSVRSTVNGLDTHATLREMMLHLKSYYHLASQRLADQLPMVLRYLVLGEFASQLQREMLQMLQEKDNLEQLLKEDIDIGSKRAALQNKLKRLMKARDYLVEF-----
Trout Mx5 518 ---GEI-AEDKEQDTRSKYPGLLKAYYEIVVQRLADQVPMLICYFILKQSAKIVCSEMLDLL-HRDDTDNILGEDSEIGQYRAKLQAQADRLILANDKISSL-----
Trout Mx6 510 ---EGT-AEGSDHDTRSKYPGLLKAYYEIVVQRLADQVPMLIRYFILKQSAKIVCSEMLDLL-HSDDTDNILGEDSEIGQYRAKLQAQADRLILANDKISIL-----
Trout Mx7 512 ---PGK-TDCSGYDTRSKYPELLNSYFEIVVQRLADQVPMLIRYFILKESARILSSEMLGLL-NREDLDEMLKEESEIGRKREALRDKVKRLGLANNKISTLWDQSG-
Trout Mx8 512 ---EET-PDCSCYDTRSKYPELLKAYYEIVVQRLADQVPMLIRYFILKESARILCSKMLGLL-NSDDLDEMLTEESEIGRRRSALRSRVERLGLANDKISSL-----
Trout Mx9 541 EDITAT-FNSTTFDSRKLTPDKLGVYYELVYQRLADQVPMLIRYFILKESAKMLCIQIMDER-DGADVVKLLSEDSMEGRRRAGLHQRLDRLKKAQEKLSEF------
Human Mx8 559 WDFGAFQSSSATDSSMEEIFQHLMAYHQEASKRISSHIPLIIQFFMLKEYSAKMLCIQIMDER-DGADVVKLLKERSDTSDKKKFLKERLARLTQARRRALQFPG-----
HUMBAN MXP 600 KINGUPDRAMPCGVGSEPPETGIJU NAVET PRIGNEI AM OTDER TOVENI DER DRAMPCH OF DER PRINDER IN DED CEPTAMPRD IL KEPLYDI MAD ALL CONSCREDE
Human MxB 608 KLNSHFPSNESSVSSFTEIGIHLNAYFLETSKRLANQIPFIIQYFMLRENGDSLQKAMMQILQEKNRYSWLLQEQSETATKRRILKERIYRLTQARHALCQFSSKEIH
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Figure 8

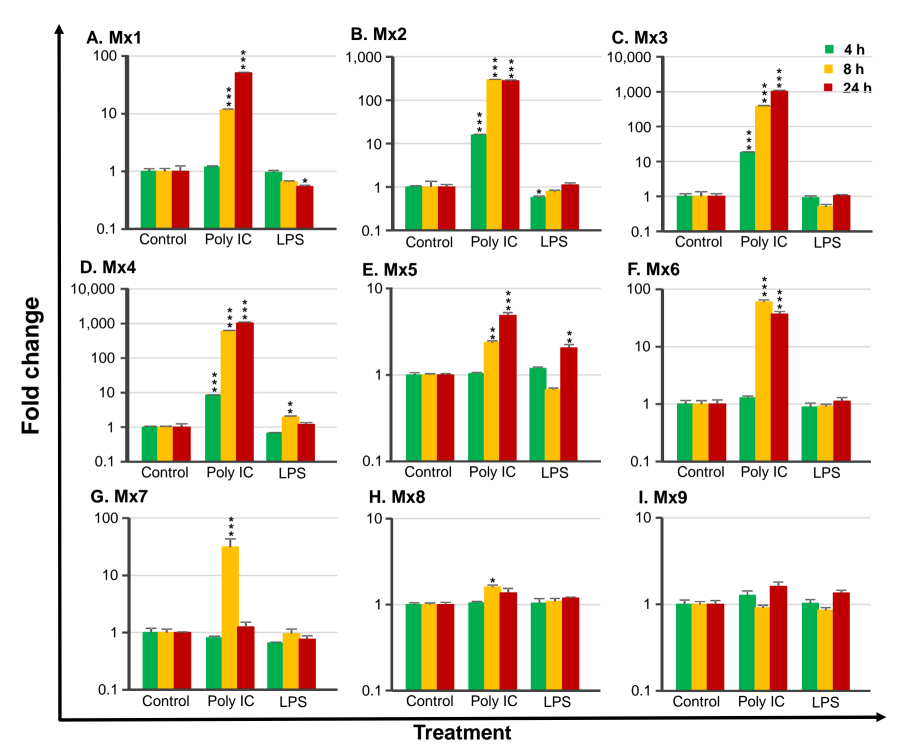
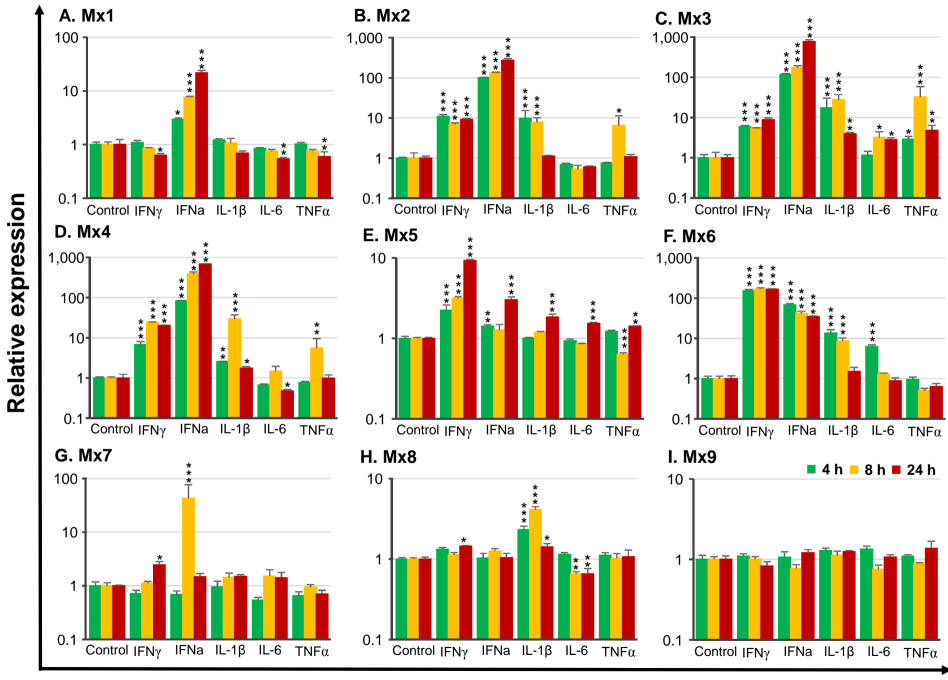
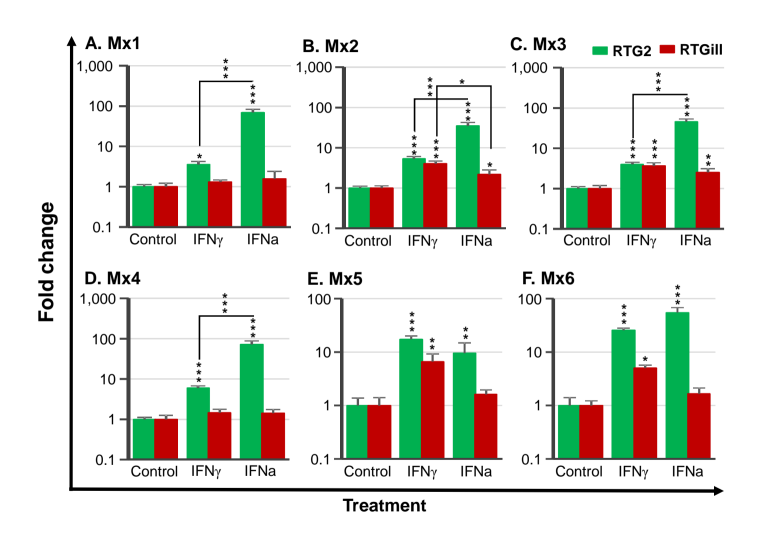


Figure 11

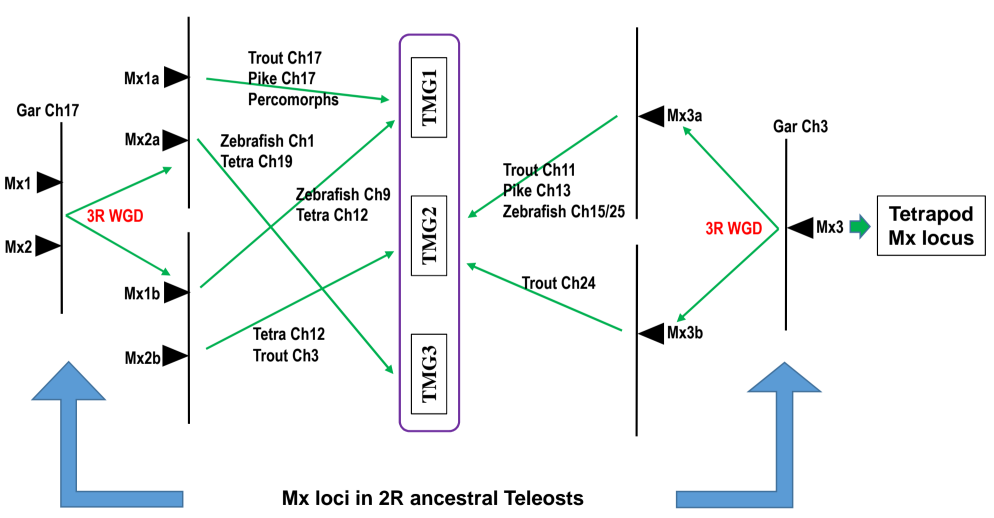


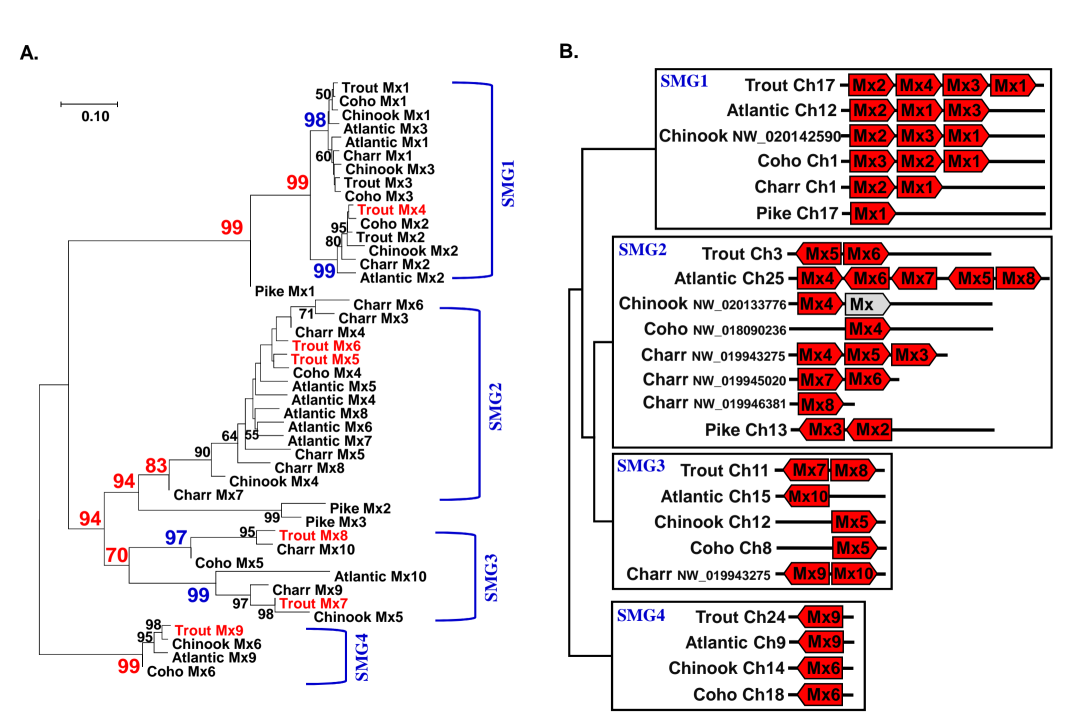
Treatment

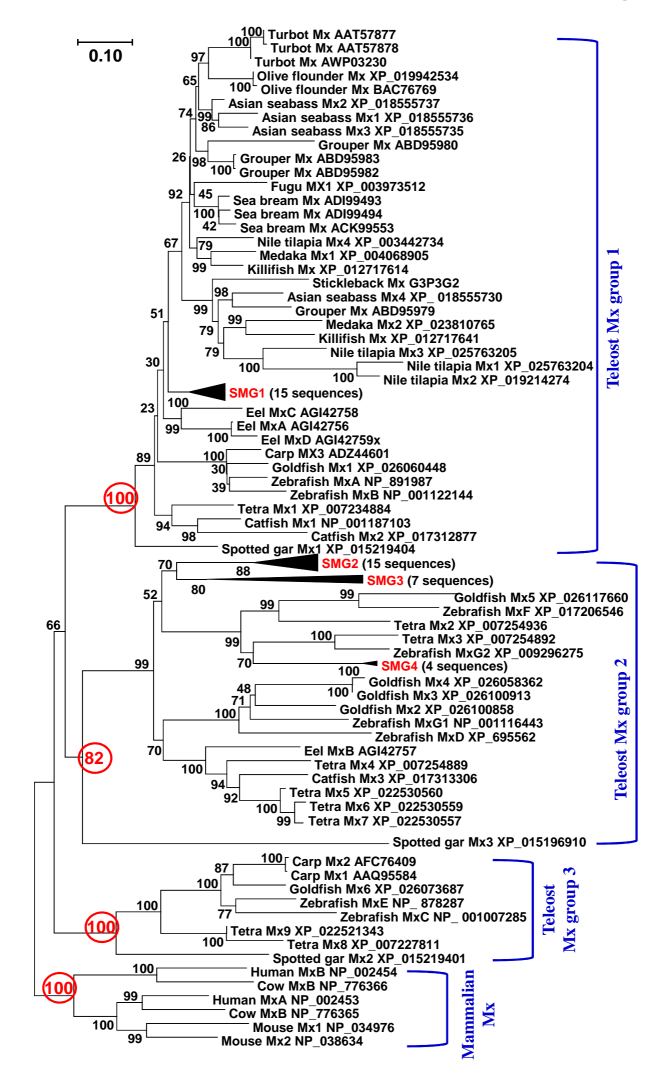
Figure 12

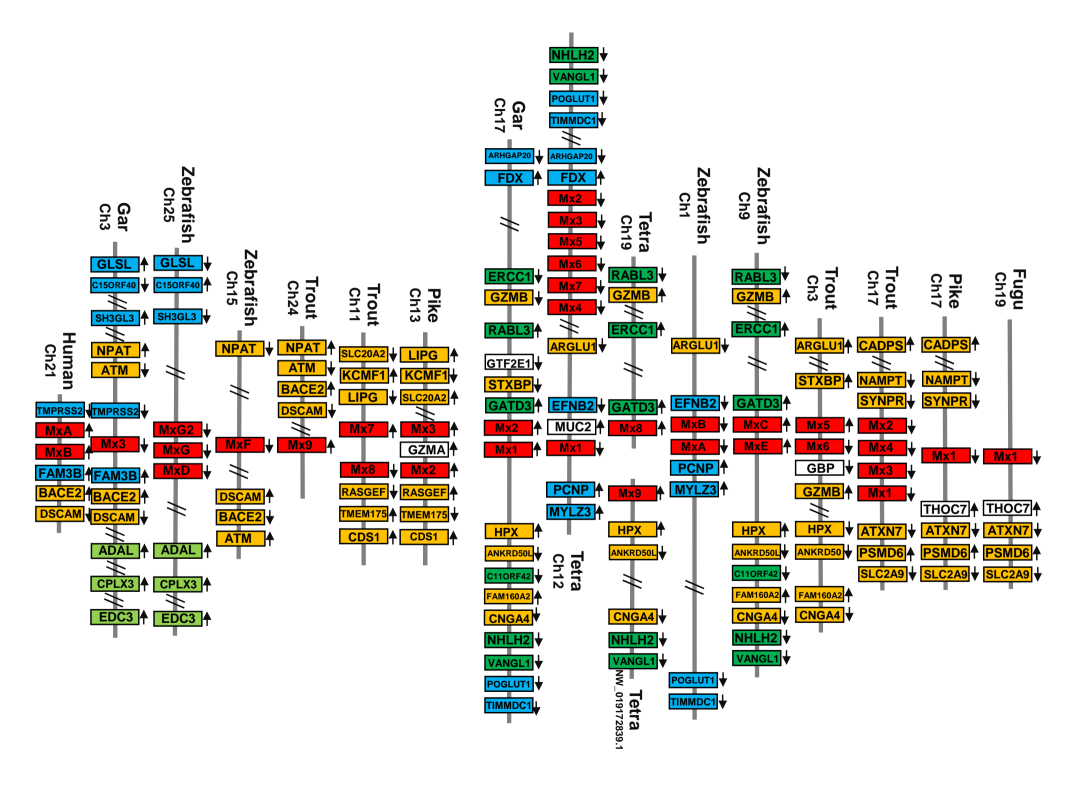










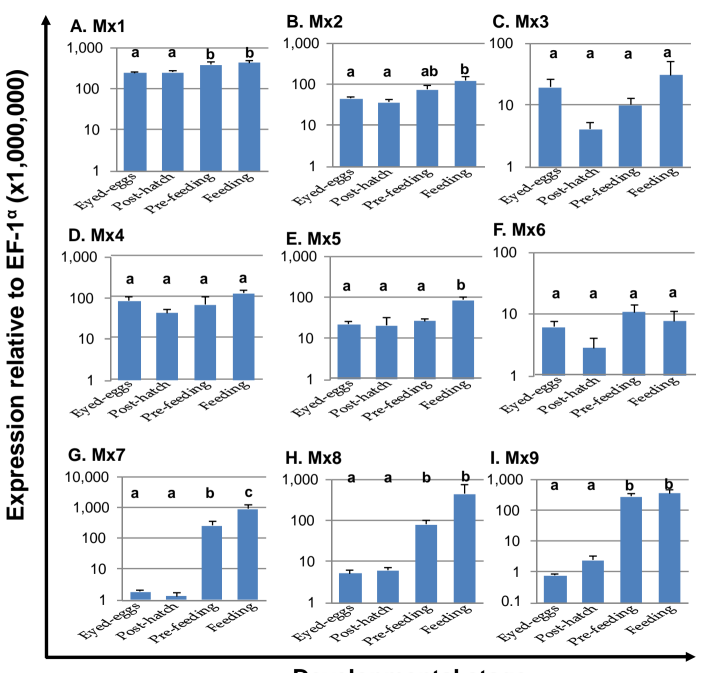


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G Domain
                                                                                                             GED
Teleost Mx group 1
                       Trout Mx1 156 190 1 138 1 155 0 139 1 199 179 123
                       Trout Mx2 53 190 1 138 1 155 0 139 1 199 79 0 123 0 139 1
                       Trout Mx3 85 190^{-1} 138^{-1} 155^{-0} 139^{-1} 199^{-1} 79^{-0} 123
                       Trout Mx4 61 190 \frac{1}{138} \frac{1}{155} \frac{1}{139} \frac{1}{199} \frac{11}{79} \frac{0}{123}
                  Zebrafish MxA 84 199 138 155 139 199 79
                 Zebrafish MxB 118^{199} - 138^{11} - 155^{11} - 139^{11} - 199^{11}
                   Tetra Mx1 165 18 190 138 155 139 199 179
                       Gar Mx1 19 7 190
Teleost Mx group 2
              Trout Mx5 46-12 23 11 182
                                              \frac{1}{135} \frac{1}{155} \frac{1}{139} \frac{1}{199} \frac{1}{179} \frac{1}{123} \frac{1}{133}
                  Trout Mx6 17 23 1182 1 135 1 155 139 1 199 179 123 133 1 159
                Trout Mx7 301 5 11 197
                                                      155^{\circ} 139^{\circ} 199^{\circ} 123^{\circ}
                                                135
                                                                                       133
                 Trout Mx8 27 26 1182
                                                                  1
199
                                                135
                                                             139
                                                                  199<sup>II</sup>
                Trout Mx9 120 38 11 200-
                                                             139
           Zebrafish MxD 178 83 1197
                                                             139 <sup>1</sup> 199 <sup>1</sup>
       Zebrafish MxF 103-33 47 11 209
                                                              139 <sup>1</sup> 199 <sup>1</sup>
           Zebrafish MxG1 9 95 \frac{11}{194}
                                                             139
                                                                    199
         Zebrafish MxG2 225 50 11 197-
                                                138
                                                              139
                                                                    199
            Tetra Mx2 179 8 23 209
                                                132
                                                              139
                                                                    199
            Tetra Mx4 214 16 50 11 197
                                                             139
                      Gar Mx3 ? 7 190
Teleost Mx group 3
                   Zebrafish MxC 12 217 \frac{1}{138} \frac{1}{155} \frac{0}{139}
                                                                   <sup>1</sup>199
            Zebrafish MxE 36 84 \frac{0}{229} \frac{1}{138} \frac{1}{155} \frac{0}{139} \frac{1}{1}
                                                                    199
                                                                  1
199
                  Tetra Mx8 115 18 223 1 138 155 139
              Gar Mx2 \begin{bmatrix} 4 & 4 \\ 8 & 6 \end{bmatrix} 16198 \frac{\sqrt{193}}{193}
 Mammalian Mx
Human MxA[5][9][9][112][7][21][105][0][138][138][155][139][199][79][123][0][142]
           Human MxB 29 71249 193
                                                             139
                                               138
```

Figure 6 A 100,000 Mx1 10,000 Mx2 1,000 Mx3 100 10 Mx4 Expression relative to EF-1a (x1,000,000) В 10,000 1,000 Mx5 100 Mx6 10 C 100,000 10,000 Mx7 1,000 Mx8 100 10 D 100,000 10,000 1,000 Mx9 100 10 Mus Skin se fin nisde sca Adipose tissue Tailfins Ci read kuri. I dhey Thyrus er Headkidney Gills RISTA RIGT RIGH Arain Blood Heart spleen je scales nad

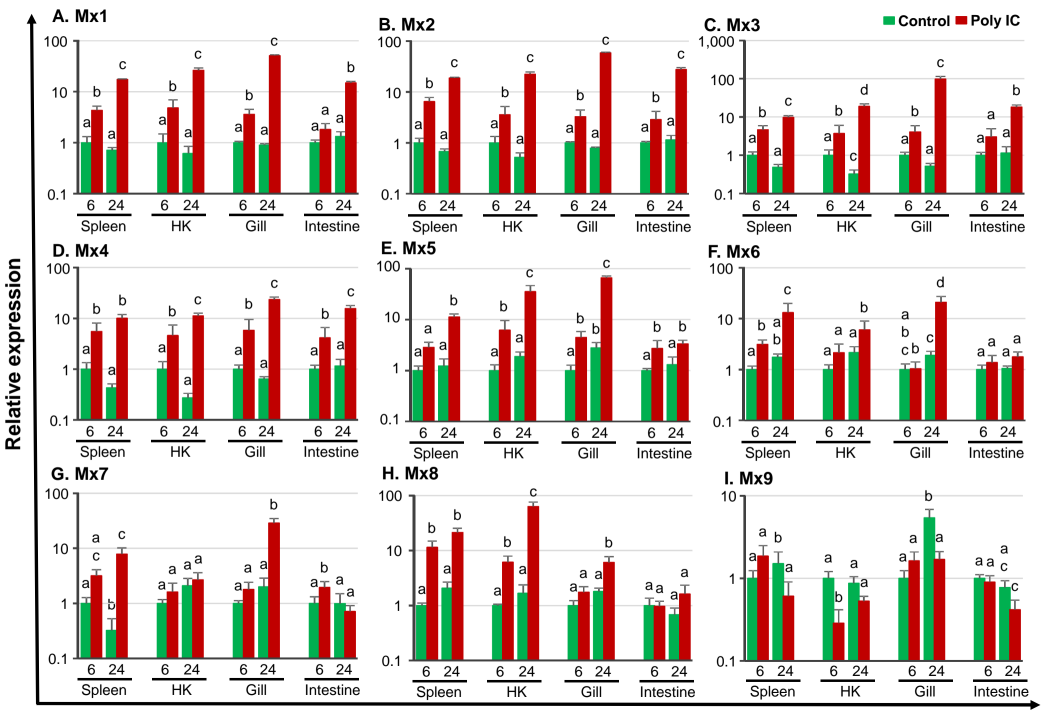
Tissues and cell lines

Figure 7



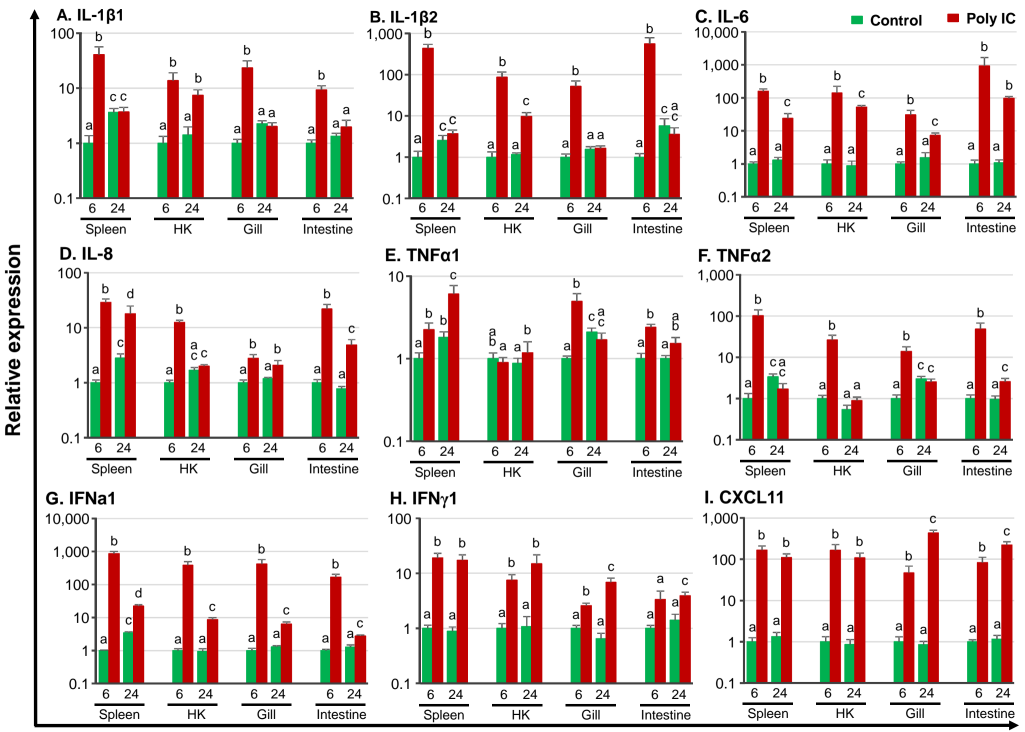
Developmental stage

Figure 9



Time (h) and tissues

Figure 10



Tissue and time (h)

Highlights

In addition to Mx1-3, six novel Mx genes (Mx4-9) have been cloned in rainbow trout Salmonids possesses 4 groups of Mx genes residing at four chromosome loci

Trout Mx1-4 are highly expressed in blood but Mx5-9 are highly expressed in intestine

Trout Mx gene expression can be induced by poly IC, type I and type II IFNs, and IL-1β

The potency of IFN induced Mx expression is gene- and cell line-dependent