# Evolution of the immune system influences speciation rates in teleost fishes

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Teleost fishes constitute the most species-rich vertebrate clade and exhibit extensive genetic and phenotypic variation, including diverse immune defense strategies. The genomic basis of a particularly aberrant strategy is exemplified by Atlantic cod, in which a loss of major histocompatibility complex (MHC) II functionality coincides with a marked expansion of MHC I genes. Through low-coverage genome sequencing (9–39×), assembly and comparative analyses for 66 teleost species, we show here that MHC I is missing in the entire Gadiformes lineage and thus was lost once in their common ancestor. In contrast, we find that MHC I gene expansions have occurred multiple times, both inside and outside this clade. Moreover, we identify an association between high MHC I copy number and elevated speciation rates using trait-dependent diversification models. Our results extend current understanding of the plasticity of the adaptive immune system and suggest an important role for immune-related genes in animal diversification.

With over 32,000 extant species<sup>1</sup>, teleost fishes comprise the majority of vertebrate species. Their taxonomic diversity is matched by extensive genetic and phenotypic variation, including novel immunological strategies. Although the functionality of the adaptive immune system has been considered to be conserved since its emergence in the ancestor of all jawed vertebrates<sup>2,3</sup>, fundamental modifications of the immune gene repertoire have recently been reported in teleosts<sup>4–7</sup>. One of the most dramatic changes has occurred in Atlantic cod (Gadus morhua), involving complete loss of the MHC II pathway that is otherwise responsible for the detection of bacterial pathogens in vertebrates<sup>4</sup>. Moreover, this loss is accompanied by a substantially enlarged repertoire of MHC I genes, which normally encode molecules for protection against viral pathogens. It has thus been hypothesized that the expanded MHC I repertoire of cod evolved as a compensatory mechanism, whereby broader MHC I functionality makes up for the initial loss of MHC II (refs. 4,6). However, the questions of how and when MHC II was lost relative to the MHC I expansion, and whether these genomic modifications are causally related, have so far remained unresolved.

As key components of the vertebrate adaptive immune system, the complex MHC pathways and their functionality are now well characterized<sup>8–10</sup>, but less is known about the causes of MHC copy number variation, which poses an immunological tradeoff<sup>11,12</sup>. Although an increase in the number of MHC genes facilitates pathogen detection, it will also decrease the number of circulating T cells<sup>13–16</sup>, resulting

in an immune system that can detect a large number of pathogens at the expense of being less efficient in removing them. The evolution of MHC copy numbers is therefore likely driven toward intermediate optima determined by a tradeoff between detection and elimination of pathogens-as suggested by selection for 5-10 copies inferred in case studies of fish<sup>17,18</sup> and birds<sup>19</sup>. Because pathogen load and the associated selective pressures vary between habitats, the optimal number of MHC copies depends on the environment<sup>20-22</sup>. As a result, interbreeding between different locally adapted populations is expected to produce hybrids with excess (above optimal) MHC diversity that are characterized by T cell deprivation and low fitness. This process would introduce postzygotic reproductive isolation and promote reinforcement of premating isolation between the populations. Consequently, MHC genes have been suggested to have an important role in speciation<sup>22,23</sup>, but, to our knowledge, this role has never been tested comparatively in a macroevolutionary context.

Here we report comparative analyses of 76 teleost species, of which 66 were sequenced to produce partial draft genome assemblies, including 27 representatives of cod-like fishes within the order Gadiformes. First, we use phylogenomic analysis to resolve standing controversy regarding early-teleost divergences and to firmly establish the relationships of gadiform fishes. Second, by analyzing the presence and absence of key genes in the vertebrate adaptive immune system, we place the loss of MHC II functionality in the common ancestor of all Gadiformes; by time calibrating the teleost

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phylogeny on the basis of fossil sampling rate estimation, we further show that this loss occurred 105–85 million years ago. Third, we demonstrate that expansions of the MHC I gene repertoire are not limited to Gadiformes but are also found in three other clades, including the exceptionally species-rich group of percomorph fishes (Percomorphaceae). By applying phylogenetic comparative methods and ancestral state reconstruction, we trace the evolutionary history of MHC I gene expansions and infer distinct copy number optima in each of them. Finally, using trait-dependent diversification models, we identify a positive association between MHC I gene copy number and speciation rate. Our results highlight the plasticity of the vertebrate adaptive immune system and support the role of MHC genes as 'speciation genes', promoting rapid diversification in teleost fishes.

# RESULTS

# Sequencing and draft assembly of 66 teleost genomes

To investigate the evolution of the immune gene repertoire of teleost fishes, we selected representatives of all major lineages in the Neoteleostei<sup>24</sup> for low-coverage genome sequencing. As both loss and expansion of MHC genes have been reported in the Gadiformes<sup>4</sup>, we sampled this order more densely with 27 species representing most gadiform families and subfamilies (Supplementary Table 1)<sup>25</sup>. The sequencing strategy for de novo assembly of the 66 genomes was designed on the basis of gene space completeness and assembly statistics of simulated genomes (Supplementary Note). For each species, a single paired-end library was sequenced to 9-39× coverage on the Illumina HiSeq 2000 platform (Supplementary Table 2 and Supplementary Note). The genome sequences were assembled with the Celera Assembler<sup>26</sup>, resulting in N50 scaffold sizes between 3.2 and 71 kb (Supplementary Table 3 and Supplementary Note). For most species, we recovered more than 75% of the conserved eukaryotic genes included in the CEGMA<sup>27</sup> analysis, and on average we recovered 74% of the conserved genes of the Actinopterygii data set included in the BUSCO<sup>28</sup> analysis. Collectively, this is indicative of a sufficiently high degree of gene space completeness for gene detection in our partial draft genome assemblies (Supplementary Table 3 and Supplementary Note).

# A genome-scale phylogeny of teleost fishes

To firmly establish teleost and gadiform relationships, we performed phylogenomic analyses using a stringently filtered data set of 567 exon orthologs from 111 genes, identified in the 66 new draft genome assemblies as well as in 10 previously published genomes. After filtering (**Supplementary Note**), the phylogenomic data set included 71,418 bp, which was used for time calibration and coalescent-based species tree analyses. To establish a timeline of teleost diversification and to determine the timing of immune gene repertoire alterations, we performed Bayesian phylogenetic inference with the concatenated phylogenomic data set, using a relaxed-clock model and 17 fossil constraints (**Supplementary Figs. 1** and **2**, **Supplementary Tables 4** and **5**, and **Supplementary Note**).

The time-calibrated species tree was used to test for incomplete lineage sorting among the species sampled for our phylogeny. Following Jarvis *et al.*<sup>29</sup>, we mapped genomic insertions and deletions on the rooted topology and found no correlation between branch lengths and uniquely mapped indels. This lack of correlation indicates that incomplete lineage sorting did not substantially affect divergences between the teleost lineages included in our phylogeny<sup>29</sup> and supports concatenation as the most appropriate strategy for species phylogeny estimation with our data (**Supplementary Note**). The resulting time tree (**Fig. 1**) supports a monophyletic clade including the orders Gadiformes, Stylephoriformes, Zeiformes, Polymixiiformes and Percopsiformes, thus corroborating the Paracanthopterygii *sensu* Miya *et al.*<sup>30</sup>. We further confirm the placement of *Stylephorus chordatus* as the closest extant relative of the Gadiformes<sup>30,31</sup> and estimate the crown age of this order at approximately 85 million years ago. Within Gadiformes, we find support for a sister group relationship between Bregmacerotidae and all other gadiform families, which began to radiate around 70 million years ago.

# Loss of MHC II pathway genes

To assess the origin and extent of the MHC II pathway gene loss observed in Atlantic cod, we investigated the immune gene repertoires of the teleost genomes through a comparative gene mining pipeline comprising BLAST searches, prediction of ORFs and annotation (**Supplementary Note**). We explicitly investigated the presence of 27 genes chosen for their central role in MHC class I, MHC class II and cross-presentation pathways<sup>8,10</sup>. In addition, three highly conserved genes were included as a control (**Fig. 2**). As query sequences, we used orthologs from zebrafish (*Danio rerio*), medaka (*Oryzias latipes*), spotted green pufferfish (*Tetraodon nigroviridis*), fugu (*Takifugu rubripes*), three-spined stickleback (*Gasterosteus aculeatus*), Nile tilapia (*Oreochromis niloticus*), Amazon molly (*Poecilia formosa*), platyfish (*Xiphophorus maculatus*), cavefish (*Astyanax mexicanus*) and Atlantic cod (*G. morhua*) (**Supplementary Table 6**).

The genome-based phylogeny allowed us to place the immune gene characterization into an evolutionary perspective. Whereas the three control genes were identified in all species, we found that genes associated with the MHC II pathway were consistently missing in all inspected Gadiformes draft genome assemblies, as no orthologs could be identified for invariant chain (also known as cd74), cd4 and the MHC II  $\alpha$  and  $\beta$  chains (Fig. 2a and Supplementary Table 7). This finding reiterates the previous observation in the Atlantic cod genome, where lack of these genes was confirmed through qPCR and synteny analyses<sup>4</sup>. Although individual genes could not be detected in a subset of species outside the Gadiformes, these occurrences comprise a minority (~3%) of the total number of comparisons. Notably, these losses did not show any phylogenetic pattern and hence are more likely to have resulted from incomplete genome assembly (Supplementary Note). As MHC II pathway genes are otherwise ubiquitous among vertebrates, the observed pattern shows that these genes were lost collectively in the common ancestor of all Gadiformes, following its divergence from Stylephoriformes at approximately 105 million years ago. This result implies that MHC II gene loss is shared by all 616 extant species of gadiform fishes. These taxa inhabit a wide variety of habitats, showing that their alternative immune system is highly versatile and not restricted to specific ecological niches.

#### MHC I copy number variation

MHC I molecules exist as five distinct lineages—L, S, P, U and Z—with the latter two comprising predominantly 'classical' (peptide-binding) MHC I molecules involved in antigen presentation<sup>32–35</sup>. The Atlantic cod genome has been shown to harbor 80–100 copies of MHC I (ref. 4), which is 15 times higher than the copy number determined to be optimal in the three-spined stickleback<sup>14,17</sup>. To investigate the evolutionary origin of the gadiform MHC I expansion and potentially detect other expansions, we estimated the MHC I copy numbers of the two peptide-binding lineages (U and Z) in the 66 sequenced species. MHC I copy number estimates were calculated on the basis of Illumina raw sequencing reads relative to a set of 19 putatively

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**Figure 1** Time-calibrated phylogeny of 76 teleost species. Dashed branches correspond to species for which genome data were already available in Ensembl. All nodes are supported by Bayesian posterior probability of 1.0 except where noted (**Supplementary Table 5**). Clade names to the right and on branches are given according to Betancur-R *et al.*<sup>24</sup>. Branch colors represent taxonomy at the level of orders. Teleost illustrations are by G. Holm.

single-copy reference regions. For both U- and Z-lineage genes, we determined the number of matching raw reads by aligning them to conserved regions of the MHC I genes ( $\alpha$ 3 domain) and the set of reference genes. The uncertainties of all copy number estimates were assessed with a double-bootstrapping procedure (**Supplementary Tables 3** and **8–11**, and **Supplementary Note**). Our copy number

estimates were cross-validated with previous results for Atlantic cod based on qPCR<sup>4</sup> and transcribed MHC I U-lineage genes<sup>6</sup>. The copy number estimation procedure was further validated by estimating the relatively conserved number of Hox gene copies for all species. Estimated Hox gene copy numbers (mean = 50.3, s.d. = 11.4) were in agreement with expectations for teleost fishes<sup>36</sup> and were uncorrelated

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**Figure 2** Immune gene repertoire and estimated MHC I copy number in the draft genome assemblies of 66 teleost species. (a) Presence of key genes in MHC I, MHC II and cross-presentation pathways for recognition of pathogen-derived antigens. Genes not detected are indicated by white squares. Colors reflect order-level taxonomy, and the cladistic representation follows that in **Figure 1**. (b) Copy number estimates for non-classical MHC I Z-lineage (light gray) and U-lineage (dark gray) genes. Error bars, 95% confidence intervals based on 1,000 bootstrap replicates (**Supplementary Note**).

with MHC I gene copy numbers ( $R^2 = 0.002$ ) (Supplementary Fig. 3, Supplementary Table 3 and Supplementary Note).

Mean copy number estimates for all species are shown in **Figure 2b**. Extremely high copy numbers of MHC I U-lineage genes were detected in Gadiformes, where several species had around 100 copies, followed by species within Percomorphaceae with up to 80 copies. Within Gadiformes, high copy numbers above 40 were observed in as many as 12 species. Although we also observed low copy numbers in a limited number of gadiform species (for example, five copies in *Trisopterus minutus*), nearly all representatives of this order appear to share an expanded MHC I gene repertoire with 15–30 copies of the U-lineage genes. Such gene family expansions may promote biological diversification by introducing new raw genetic material, potentially resulting in sub- or neofunctionalization and thus novel immunological pathways. In this regard, we identified the same cytoplasmic sorting motifs presumed to be part of a novel pathway



**Figure 3** MHC class I copy number evolution and diversification rate analysis. (a) 3D representation of teleost phylogeny (**Fig. 1**) and MHC I copy number evolution, based on ancestral state reconstruction with the best fitting Ornstein–Uhlenbeck model. This model supports five shifts toward elevated copy number optima, marked by colored dots: 25.2 copies in Percomorphaceae excluding Ophidiiformes, 41.5 in Berycoidei, 22.1 in Polymixiiformes and Percopsiformes, 19.8 in Gadiformes excluding Bregmacerotidae, and 57.5 in Gadiformes excluding Bregmacerotidae and Gadidae. Loss of MHC II in the common ancestor of Gadiformes is represented by a black dot. The ordering of species is identical to that in **Figure 2**. Gray shading visualizes reconstructed MHC I copy numbers along branches. Steps of 20 copies are indicated by light gray lines on the 3D reconstruction. (**b**) Per-species MHC I copy numbers and mean net diversification rate estimates, connected according to phylogenetic relationships and colored as in **a**. (**c**) Diversification rate estimates for lineages with high and low MHC I copy numbers. Black lines represent mean estimates and gray shading represents standard deviation from 25 replicate analyses. (**d**) Improvement in log-transformed likelihood when speciation rates are unlinked for lineages with high and low copy numbers, in the analyses performed in **c**. The black line corresponds to the median and gray shading represents 0.05 and 0.95 quantiles from replicate analyses.

enhancing cross-presentation functionality in Atlantic cod<sup>6,37,38</sup> also in five additional species of the Gadinae. Because the origin of these signaling motifs only dates back to about 15 million years ago, it is likely that the enhanced cross-presentation functionality was not part of a preexisting machinery that could have favored the loss of MHC class II (**Supplementary Fig. 4**, **Supplementary Table 12** and **Supplementary Note**).

Two hypotheses have been put forward as possible explanations for the loss of MHC II: a metabolic gain by not maintaining a costly system or a functional shift in the immune gene repertoire, rendering it obsolete<sup>39</sup>. Whether the loss occurred before or following the expansion of MHC I genes is key in discriminating between these hypotheses. In contrast to all other Gadiformes, the most basal gadiform lineage, represented by *Bregmaceros cantori*, is characterized by a complete absence of MHC I U-lineage genes, in addition to MHC II loss. The only antigen-presenting molecules detected in Bregmacerotidae are encoded by the MHC I Z-lineage genes, of which 2–3 copies are still present. With minor exceptions, the Z-lineage genes were found in low copy numbers in all species, and, contrary to the U-lineage genes, they did not show a pattern of clade-specific expansions. Interestingly, the myctophiform *Benthosema glaciale* showed convergent loss of the MHC I U-lineage genes, and both *Bregmaceros* and *Benthosema* have experienced an additional loss of two genes involved in MHC I interactions with T cells (*cd8a* and *cd8b*), rendering the traditional pathway for endogenously derived pathogens non-functional as well (**Fig. 2**). The lack of MHC I U-lineage genes in *B. cantori* indicates that MHC I expansions in Gadiformes occurred after the divergence of Bregmacerotidae and thus subsequent to the loss of MHC II.

# Rate shifts in MHC I copy number evolution

It has previously been shown that MHC copy number and molecular diversity are linked to the habitat that a species or population occupies<sup>21,22</sup>. Our data show that the species-rich clades Gadiformes and Percomorphaceae, which are both distributed across a wide variety of habitats, contain the highest numbers of MHC I gene copies among teleosts. We therefore used phylogenetic comparative methods to test for putative adaptive evolution of MHC I copy numbers in all groups

included in our phylogeny. We used a Markov chain Monte Carlo (MCMC)-based reversible-jump Bayesian approach to fit multipleregime Ornstein-Uhlenbeck models in which trait evolution is directed toward different optimal values in different parts of the tree<sup>40</sup>. This approach identified the probability and direction of rate shifts in MHC I copy number optima on branches in the phylogeny (Supplementary Fig. 5 and Supplementary Note). On the basis of the inferred branch-specific rate shift probabilities and the presence of copy number outliers, we tested explicit hypotheses for rate shift combinations in a likelihood framework<sup>41,42</sup>. We found that the overall best fitting model included five shifts of the MHC copy number optimum (Fig. 3a, Supplementary Table 13 and Supplementary Note). This model is characterized by a phylogenetic half-life of 23 million years, a stationary variance of 0.38 squared log copy numbers and an optimum of 6.8 MHC I copies for basal branches of the phylogeny, which is in concordance with the hypothesized MHC I repertoire of early gnathostomes<sup>43</sup>. In contrast, substantially higher optimal MHC I copy numbers of up to 58 copies were inferred within gadiform and percomorph fishes. The phylogenetic half-life of 23 million years shows that the MHC optima are approached slowly and that there can be considerable lag in adaptation. The multiple-optima model vastly outperformed alternatives such as Brownian motion, white noise, single-peaked Ornstein-Uhlenbeck and early-burst models. This finding corroborates the hypothesis that MHC I copy number evolution is characterized by selection toward intermediate optima, resulting from a tradeoff between detection and elimination of pathogens.

# **Diversification rate analyses**

The Gadiformes comprise 616 extant species<sup>1</sup> and are thus the most species-rich order within Paracanthopterygii, where Stylephoriformes are represented by a single species and Zeiformes, Percopsiformes and Polymixiformes collectively include 56 species. To investigate whether this pattern of diversity is the result of an elevated speciation rate in Gadiformes, we tested for differences in diversification among all clades included in our phylogenetic data set using a Bayesian framework (Supplementary Figs. 6-8 and Supplementary Note). For this analysis, we identified a set of 37 mutually exclusive clades that represent almost the entire extant diversity of the teleost supercohort Clupeocephala<sup>24</sup>. Assuming constant speciation and extinction rates within specific shift regimes, we identified two major shifts in diversification rates, one at the base of Gadiformes and a second within the taxonomically diverse Percomorphaceae<sup>24</sup>, that is, in the two clades featuring particularly high MHC I copy numbers (Fig. 3a). A comparison of per-species MHC I copy numbers and mean net diversification rates (Fig. 3b) indicates a relationship between the two measures. To quantify the association between MHC I copy numbers and rates of diversification, we carried out binary state speciation and extinction (BiSSE) analysis to estimate differences in diversification rate between lineages with high and low MHC I copy numbers (Supplementary Note). We found that diversification rates differed most when the threshold was placed between 20 and 25 copies (Fig. 3c). With a threshold in this range, the model with two separate speciation rates for lineages with high and low copy numbers was better supported than a model with a single speciation rate parameter ( $\Delta AIC > 37.1$ ; Fig. 3d and Supplementary Note). These results suggest that the influence of MHC I genes on speciation rates is stronger in species that have already evolved at least 20 copies.

# DISCUSSION

Teleost fishes are characterized by striking differences in species richness across different lineages, with some groups, such as cyprinids

and cichlids, containing thousands of species, while others include only a single or very few taxa. Although these differences can in part be explained by ecological opportunity<sup>44</sup> or key innovations<sup>45</sup>, a large proportion of lineage-specific variation in taxonomic diversity remains unexplained. It has previously been suggested that MHC genes can influence speciation rates through selection against hybrids with higher than optimal MHC copy numbers and consequent reinforcement<sup>22,23</sup>. For species with more MHC copies, the extra number of alleles in hybrids will be comparatively higher, and these species should therefore experience greater fitness reduction and stronger assortative mating to maintain co-adapted genes. The proposed role of MHC genes as speciation genes<sup>23</sup>, promoting diversification and the maintenance of recently diverged species, is therefore expected to be more pronounced in species exhibiting high copy numbers. The identification of an MHC I copy number threshold above which speciation rates are accelerated expands on this conceptual framework. Our analyses identify this threshold at 20-25 MHC I copies, suggesting that the effect of T cell depletion on hybrid fitness becomes more pronounced in this range and that this might affect mate choice in species with copy numbers above this threshold, promoting inbreeding and reinforcement.

By pinpointing the loss of MHC II pathway genes to the common ancestor of Gadiformes and by reconstructing MHC I copy number evolution, we show that loss of MHC II predated the MHC I expansions that occurred within this group. This implies that MHC II functionality was not outcompeted by a preexisting alternative immunological strategy such as an expanded MHC I repertoire but may instead have been lost as a result of the metabolic costs associated with its maintenance<sup>39</sup>. The temporal relationship between the loss and expansions in Gadiformes and the identification of putative cross-presentation signaling motifs in Atlantic cod<sup>6</sup>, as well as in several other gadiform species, indicate that a number of genes in the expanded MHC I repertoire evolved to compensate for loss of MHC II. Nevertheless, marked expansions of MHC I genes were also observed outside of Gadiformes in groups with intact MHC II pathways. These independent expansions clearly show that loss of MHC II is not the only potential trigger for MHC I expansion but that MHC copy number evolution has also been driven by other factors, highlighting the extensive plasticity of the teleost adaptive immune system.

The 66 new teleost draft genome assemblies provide unprecedented opportunities for comparative analyses of the largest clade of vertebrates. We have used this genomic information to unravel the evolutionary history of key immune genes and to show how MHC gene composition in teleost fishes has influenced diversification rates in this diverse vertebrate lineage.

**URLs.** GitHub repository for scripts used in this study, https://github. com/uio-cees/teleost\_genomes\_immune.

## METHODS

Methods and any associated references are available in the online version of the paper.

Accession codes. All reads generated for this project have been deposited in the European Nucleotide Archive (ENA) under study accession PRJEB12469 (sample identifiers ERS1199874–ERS1199939). All new assemblies (unitigs and scaffolds) reported on here have been deposited in the Dryad repository under doi:10.5061/dryad.326r8.

Note: Any Supplementary Information and Source Data files are available in the online version of the paper.

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#### AUTHOR CONTRIBUTIONS

M. Malmstrøm, K.S.J. and S.J. conceived the project. M. Malmstrøm, M. Matschiner, W.S., N.C.S., K.S.J. and S.J. planned and oversaw the project. M. Malmstrøm, R.H., W.S. and S.J. provided samples for genome sequencing. M. Malmstrøm prepared DNA and libraries for genome sequencing. M. Malmstrøm and S.J. were in charge of genome sequencing management. M. Malmstrøm, O.K.T., H.T.B. and A.J.N. performed genome assembly. M. Matschiner performed phylogenomic, diversification rate and comparative analyses. T.F.H. contributed to comparative analyses. M. Malmstrøm analyzed gene presence and absence. M. Malmstrøm and L.G.S. performed copy number estimation. M. Malmstrøm and M. Matschiner prepared the figures. M. Malmstrøm, M. Matschiner and B.S. wrote the manuscript with input from all other authors.

## COMPETING FINANCIAL INTERESTS

The authors declare no competing financial interests.

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# **ONLINE METHODS**

**Tissues, sequencing and assembly.** Genomic DNA was obtained from various tissues of the different species in this study. Most tissue samples were provided by museums and other collections, while some come from commercially caught fish in collaboration with local fishermen (see **Supplementary Table 1** for a full list of tissues and contributors). A single paired-end library, with an insert size of ~400 bp, was created for each species, using the Illumina TruSeq Sample Prep v2 Low-Throughput protocol. All species were sequenced (2 × 150 bp) to >9× coverage on the Illumina HiSeq 2000 platform, and sequences were assembled using the Celera Assembler<sup>26</sup> (**Supplementary Note**). Draft genome assembly quality, in terms of gene space completeness, was assessed using CEGMA<sup>27</sup> and BUSCO<sup>28</sup> (**Supplementary Table 3** and **Supplementary Note**).

Phylogenetic inference. Strict filtering criteria were applied for the identification of suitable orthologous phylogenetic markers. For the 33,737 annotated zebrafish genes in release 78 of the Ensembl database<sup>46</sup>, we selected the longest transcript if it had at least five stop-codon-free exons of 150 bp or greater in length. We removed genes that could not be assigned to an Ensembl gene tree and genes for which teleost fishes did not form a monophyletic group in the gene tree. We further excluded genes for which the Ensembl gene tree indicated gene duplications among teleosts or did not include all ten teleost species of Ensembl v.78 (Supplementary Note). The ten teleost reference genomes of Ensembl were used to calculate TBLASTN bitscores for each of the zebrafish exons, using the BLAST+ v.2.2.29 suite of tools<sup>47</sup>. Exon-specific bitscore thresholds for putative orthologs were defined on the basis of this bitscore information, and exons for which two or more of the known orthologs had bitscores lower than this threshold were excluded. Genes with fewer than five remaining exons were discarded, which resulted in a set of 2,251 exon sequences of 302 zebrafish genes that were then used as queries in TBLASTN searches against each of the 66 new teleost draft genome assemblies, the 10 Ensembl teleost genomes and the genome sequence of salmon<sup>48</sup>. For each species, the best hits were accepted as putative orthologs if their TBLASTN bitscores were above the exon-specific bitscore threshold (Supplementary Note). Alignments of TBLASTN hits for the 2,251 exons were further filtered on the basis of ratios of nonsynonymous to synonymous substitutions (dN/dS) determined with the software codeml of the PAML 4 package<sup>49</sup> to exclude exons putatively under positive selection. Unreliably aligned sites were identified with the software BMGE v.1.0 (ref. 50) and excluded from the alignments. We further removed all third codon positions from the alignments and excluded exons with high variation in GC content (Supplementary Note). For each gene, the concordance of individual exon trees was assessed with the software Concaterpillar v.1.7.2 (ref. 51), and we removed either individual exon alignments or all exon alignments for a given gene from the data set, depending on the number of congruent exon trees (Supplementary Note). To filter for genes with clock-like evolution, we estimated the coefficient of variation of rates of each gene with the software BEAST v.2.2.0 (ref. 52) and removed the genes with the highest estimated rate variation coefficients (Supplementary Note). After this step, our strictly filtered data set used for phylogenetic inference contained 567 exons from 111 genes, with a total alignment length of 71,418 bp and 7.3% missing data. To assess the consequences of strict filtering on phylogenetic inference, we compared maximum-likelihood phylogenies based on the strictly filtered data set (111 genes, 71,418 bp, 7.3% missing data) with phylogenies based on a data set that was substantially larger but less strictly filtered (302 genes, 252,442 bp, 18.2% missing data). Maximum-likelihood phylogenies for both data sets were inferred with the software RAxML v. 8.1.17 (ref. 53) (Supplementary Fig. 1 and Supplementary Note). The strictly filtered data set was further used to estimate the timeline of teleost diversification with the software BEAST v.2.2 (ref. 52). Calibration densities for time calibration were calculated with the BEAST add-on CladeAge<sup>54</sup>, on the basis of estimates of diversification rates and the fossil sampling rate. The earliest fossil occurrences of 17 clades in our phylogeny were identified and used to constrain the ages of these clades with CladeAge calibration densities, taking into account the uncertainties in the ages of these fossils (Supplementary Note). We further used coalescent-based species tree estimation to test for potentially misleading phylogenetic signal due to incomplete lineage sorting. This analysis was conducted both with individual gene trees and with trees based on alignments binned according to the binning approach of Mirarab et al.<sup>55</sup>.

Maximum-likelihood trees produced with RAxML and maximumclade-credibility trees resulting from BEAST analyses of binned and unbinned genes were used for species tree inference with the software ASTRAL v.4.7.8 (ref. 56) (**Supplementary Fig. 2**, **Supplementary Table 5** and **Supplementary Note**). To test for incomplete lineage sorting among the taxa included in our phylogeny, we compared branch lengths and the proportion of synapomorphic indels supporting each branch, following Jarvis *et al.*<sup>29</sup> (**Supplementary Note**).

**Gene mining of draft genome assemblies.** All draft genome assemblies were mined for genetic content on the unitig (UTG) assembly level, as assembly parameters are stricter for UTGs than for contigs or scaffolds. The presence or absence of each gene was determined through an automated pipeline, using full-length amino acid sequences for 27 immune-related genes and 3 control genes, from ten teleost reference genomes (Ensembl gene identifiers are listed in **Supplementary Table 6**). Potential orthologs were detected using TBLASTN with an acceptance level of *e* value =  $1 \times 10^{-10}$  and evaluated through identification of ORFs predicted by the software Genescan<sup>57</sup>. All ORFs were then aligned to the UniProt database (**Supplementary Note**), and reciprocal TBLASTN hits were recorded as potentially correct if their *e* value was below  $1 \times 10^{-10}$ . All recorded annotations for each gene were then manually inspected, and the best hit is reported (see the **Supplementary Note** for details and **Supplementary Table 7** for the location of each identified ortholog).

Copy number estimation of MHC I genes. High sequence similarity and conserved regions make the different MHC I genes difficult to assemble correctly. To estimate the number of copies of these genes in each of the sequenced genomes, we applied a new method for copy number estimation, based on a comparison of raw read counts for target and reference sequences. For MHC I U- and Z-lineage genes, we used 270 bp of the conserved  $\alpha 3$  domain as the target and equivalently sized fragments from 14 single- or low-copy genes as references (see Supplementary Table 9 for a full overview of all reference regions). MHC I target sequences were prepared through consensus by majority for all hits detected in the individual draft genome assemblies with TBLASTN (e-value cutoff set to  $1\times 10^{-5})$  using U- and Z-lineage MHC I  $\alpha 3\text{-}$ domain sequences from ten teleost reference genomes as queries. The number of copies of each of the target genes was determined on the basis of the number of unique sequencing reads mapping to this region, relative to the number of reads matching each of the reference gene regions. The copy numbers of each of the reference gene regions were estimated first, using an iterative method and four different BLAST stringencies. Not all reference regions fulfilled our criteria, and some references were discarded for some species (see the Supplementary Note for details and Supplementary Table 11 for a full list of the references used for each species). Copy numbers for both MHC I lineages were then estimated by comparing the number of raw reads matching both the target and reference sequences and taking estimated genome size, coverage variation and total number of reads into account. The uncertainties of all copy number estimates were assessed with a double-bootstrapping procedure (Supplementary Note).

Rate shifts in MHC I copy number evolution. Phylogenetic signal in MHC I copy number evolution was assessed with Blomberg's K statistic<sup>58</sup>, calculated using the phylosignal function of the picante R package v.1.6-2 (ref. 59), and with Pagel's lambda<sup>60</sup>, calculated with function phylosig of the phytools R package v.0.4-45 (ref. 61) (Supplementary Note). The fits of four general models of trait evolution were compared on the basis of their sample-size-corrected Akaike information criterion (AICc), using the function fitContinuous of the geiger R package v.2.0.3 (ref. 62): a white noise model, a Brownian motion model, an early-burst model<sup>63</sup> and a single-peak Ornstein–Uhlenbeck model<sup>40,64</sup> (Supplementary Note). The reversible-jump Bayesian approach of the bayou R package v.1.0.1 (ref. 65) was used to perform MCMC sampling of locations, magnitudes and numbers of shifts in multiple-optima Ornstein-Uhlenbeck models (Supplementary Fig. 5 and Supplementary Note). On the basis of the results of the bayou analysis, explicit hypotheses for shift combinations were tested in a likelihood framework, using the SLOUCH R package<sup>41,42</sup>. For each shift combination, the likelihood of the best fitting combination of optimum, half-life and stationary variance was recorded and used for model comparison based on AICc scores (Supplementary Table 13

and **Supplementary Note**). The ancestral states of log-transformed MHC I copy numbers were reconstructed for internal nodes of the time-calibrated phylogeny, on the basis of the best fitting Ornstein–Uhlenbeck model (**Supplementary Note**).

Diversification rate analyses. Patterns of species diversification were analyzed with the Bayesian framework implemented in BAMM v.2.2.0 (ref. 66), on the basis of the time-calibrated phylogeny and counts of species richness in each of the 37 mutually exclusive clades of teleost fishes (Supplementary Table 14). The 'MEDUSA-like' model of diversification, assuming constant speciation and extinction rates within specific shift regimes<sup>67</sup>, was used for this analysis (Supplementary Fig. 8 and Supplementary Note). To test whether high MHC I copy numbers are associated with lineages that have high diversification rates, we carried out BiSSE analyses<sup>68</sup> with the diversitree R package<sup>69</sup>. In these analyses, species were grouped into two categories for high and low MHC I copy numbers, on the basis of a given threshold value. Analyses were repeated for 26 equally spaced copy number threshold values between 10 and 60. As diversitree allows terminal clades with extant diversities of no more than 200 species, we used birth-death models of diversification in combination with the diversified sampling scheme of Höhna et al.<sup>70</sup> to stochastically resolve subclades of all clades with more than 200 extant species, which was repeated 25 times. BiSSE analyses were conducted for each of the 25 resulting phylogenies and with each of the 26 copy number thresholds, assuming symmetric transition rates between high and low copy numbers and identical extinction rates in taxa with high and low copy numbers (Supplementary Note and Supplementary Data).

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