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Expressed IgH μ and τ transcripts share diversity segment in ranched Thunnus orientalis

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ABSTRACT

It is now appreciated that in addition to the immunoglobulin (Ig)M and D isotypes fish also make the mucosal IgT. In this study we sequenced the full length of Ig τ as well as μ in the commercially important Thunnus orientalis (Pacific bluefin tuna), the first molecular analysis of these two Ig isotypes in a member of the order Perciformes. Tuna IgM and IgT are each composed of four constant (CH) domains. We cloned and sequenced 48 different variable (VH) domain gene rearrangements of tuna immunoglobulins and grouped the VH gene sequences to four VH gene segment families based on 70% nucleotide identity. Three VH gene families were used by both IgM and IgT but one group was only found to be used by IgM. Most interestingly, both μ and τ clones appear to use the same diversity (DH) segment, unlike what has been described in other species, although they have dedicated IgT and IgM joining (JH) gene segments. We complemented this repertoire study with phylogenetic and tissue expression analysis. In addition to supporting the development of humoral vaccines in this important aquaculture species, these data suggest that the DH-IH recombination rather than the VH-DH recombination may be instructive for IgT versus IgM/D bearing lymphocyte lineages in some fish.

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1. Introduction

The immunoglobulin (Ig) superfamily-based adaptive immune 47 system evolved in cartilaginous fish (sharks and skates) and is 48 maintained in all jawed vertebrates (Flajnik and Rumfelt, 2000). 49 50 One of the major characteristics of this adaptive immune system is the production of a repertoire of antibodies through somatic 51 52 V(D)J recombination of the loci that encode them. While mammals possess five functionally distinct Ig isotypes (IgM, IgD, IgG, IgA and 53 IgE), teleost fish have only three: IgM, IgD and IgT (Danilova et al., 54 55 2005a,b; Fillatreau et al., 2013; Hansen et al., 2005a; Wilson et al., 56 1997).

IgT was concomitantly discovered in trout (Oncorhynchus my-57 kiss) and zebrafish (Danio rerio, where it was given the appellative 58 59 IgZ) and IgT or forms of Ig with IgT domains have since been

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described in fugu (Fugu rubripes) (Savan et al., 2005b), carp (Cyprinus carpio) (Savan et al., 2005a), and stickleback (Gasterosteus aculeatus) (Gambon-Deza et al., 2010). IgT perhaps exists in most teleost groups, although it has yet to be found in catfish (Bengten et al., 2006; Salinas et al., 2011) and medaka (Magadan-Mompo et al., 2011). So far IgT is an isotype restricted to bony fish, and sequence characteristics (Hansen et al., 2005a), gut localization and functional work (Zhang et al., 2010) have suggested that it is a dedicated mucosal isotype (Zhang et al., 2011), functionally analogous but not orthologous with IgX/A of tetrapods (Mashoof et al., 2013). IgT was found to be expressed in gill of Chinese perch (Siniperca chuatsi) (Tian et al., 2009), IgT positive cells were identified in the epithelium of trout gill lamellae (Olsen et al., 2011), and clonal IgT responses were induced to trout viral pathogens (Castro et al., 2013), all further supporting the idea of this isotype filling a mucosal role in teleost humoral adaptive immunity. The IgT encoding DH-IH-CH elements are located 5' of the μ and δ DH-IH-CH regions in the fish genomes in which it has been studied, with most or all VH genes 5' to the τ block (Danilova et al., 2005a,b; Gambon-Deza et al., 2010; Savan et al., 2005b). Although class switch recombination has been described in shark (Zhu et al., 2012) and fish activation-induced cytidine deaminase (AID) is competent to induce somatic hypermutation and class switch in mammalian

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cells (Wakae et al., 2006), it does not appear that teleosts employ this for Ig heavy (H) chain isotype switching, instead they use deletional VH(DH)JH rearrangement to remove τ in IgM and IgD expressing cells and differential RNA splicing to control expression of IgM and IgD (Hikima et al., 2011), the τ/μ rearrangement appearing to have influence on lineage commitment similarly to the mechanism operating at the T cell receptor $\alpha\delta$ locus.

We recently turned our attentions to the expressed IgH transcripts of the Pacific bluefin tuna (*Thunnus orientalis*). *Thunnus* species are the most valuable global aquaculture product (Ottolenghi, 2008), yet infections from several groups of parasites plague high intensity tuna mariculture ranches (Fromentin and Powers, 2005), impeding the industry from optimal relief of fishing pressures upon wild adult stocks. In addition to their economic importance, the extreme physiological specializations of these migratory apex predators made their Ig of interest to us. Tuna are among the fastest fish and have countercurrent heat exchangers that minimize convective heat loss to maintain a form of endothermy distinct from that of birds and mammals (Block et al., 2001; Jusup et al., 2011). Specifically, we were curious whether tuna Ig harbored any special adaptations evident in their primary amino acid sequence to this rare form of fish endothermy.

105 Here we report the first full-length μ and τ sequences from 106 tuna. We have analyzed representative clones of the expressed 107 variable domain repertoire of these isotypes, performed phyloge-108 netic analysis of the IgH genes of this modern teleost, and analyzed 109 their relative expression in tuna primary and secondary lymphoid tissues, including the mucosal gill. Our results demonstrate that 110 these fish employ the same Ig VH gene families as other teleosts, 111 112 can use the same VH genes in both IgM and IgT heavy chains, make 113 diverse IgH complementarity determining region (CDR)3 regions, 114 and surprisingly employ the same DH segment in both τ and μ 115 rearrangements in what appears to be a previously undescribed 116 mechanism of B cell isotype determination.

117 2. Methods

118 2.1. Animals and collection of tissues

119 Sample tissues of spleen, gill and kidney from ranched T. orientalis were collected during the regular slaughter process from two 120 121 different commercial tuna facilities located off the coast of Ensena-122 da, Baja California, Mexico. At the time of harvest, fish weight and 123 fork length were 16.2 ± 6.5 kg and 96.3 ± 14.3 cm, respectively. 124 Samples were placed in RNAlater (Qiagen, Valencia CA), frozen in 125 liquid nitrogen, shipped to Texas A&M on dry ice and stored at 126 -80 °C until further use.

127 2.2. Total RNA isolation and cDNA synthesis

Total RNA was purified from spleen, gill and head kidney (pro-128 nephros, or anterior kidney) (35 mg from each tissue) using the 129 130 RNeasy Mini Kit (Qiagen) according to the manufacturer's instruc-131 tion. The quantity and quality of the RNA samples were assessed by NanoDrop 2000c spectrophotometer (Thermo Scientific, Wilming-132 133 ton, DE) and Agilent 2100 Bioanalyzer (Agilent, Santa Clara, CA) respectively. Message representation of RNA was assessed by PCR 134 135 of common (β -actin) and less common transcripts (TNF- α , IL1- β), 136 using previously published primer sets (Mladineo and Block, 137 2009). The GeneRacer kit (Life Technologies, Grand Island, NY) 138 with GeneRacer oligo dT and gene specific primers was used to produce 5' rapid amplification of cDNA ends (RACE) PCR products. 139 140 Pools of 3' RACE products were synthesized by Superscript III First-141 Strand Synthesis SuperMix kit (Life Technologies) using the oligo 142 dT primer.

2.3. IgH RACE PCR, cloning, and sequencing

5' and 3' RACE products were amplified by standard PCR using 144 various combinations of 5' GeneRacer (as forward primer in 5' 145 RACE), Oligo dT (as reverse primer in 3' RACE), and specifically de-146 signed primers for the conserved regions encoding the C domains 147 of T. orientalis IgM and IgT (as forward or reverse for 3' RACE or 148 5' RACE, respectively). Primers are listed in Supplemental Table 149 1. The PCR conditions were as follows: one cycle of 95 °C for 150 2 min, 35 cycles of 95 °C for 30 s, 50-53 °C for 30 s, 72 °C for 151 2 min, followed by one cycle of 72 °C for 7 min. The amplicons 152 were purified from a 0.8% agarose gel after electrophoresis in 153 tris/acetic acid/EDTA (TAE) buffer, cloned into pCR II vector with 154 the TOPO TA cloning kit (Life Technologies), and transformed into 155 chemically competent TOP10 Escherichia coli cells (Invitrogen). 156 Colonies were picked based on blue/white screening produced by 157 X-Gal (Sigma-Aldrich, Saint Louis, MO). The plasmid DNA was 158 purified using Zyppy Plasmid Miniprep kit (Zymo Research Corpo-159 ration, Irvine, CA) and was digested with EcoRI (Promega, Madison, 160 WI) to identify clones with inserts. Products for sequencing were 161 amplified using either M13 forward or reverse primers, purified 162 using ABI BigDye \times terminator purification kit (Life Technologies), 163 and sequenced by the DNA Technologies Core lab of the Depart-164 ment of Veterinary Pathobiology at Texas A&M University. 165

2.4. Sequence analysis of μ and τ gene rearrangements in Pacific bluefin tuna

BLASTX (http://blast.ncbi.nlm.nih.gov/Blast.cgi) and visual 168 inspection were used to identify the Ig isotype as well as VH, JH 169 and CH sequences of tuna amplicons based upon homology to 170 those from representative fish and other vertebrates. The amino 171 acid sequences were blasted to discriminate the VH segments 172 and CH domains. SignalP 4.1 was used to determine the leader 173 peptides (Emanuelsson et al., 2007). Three prediction methods 174 concurred upon the cleavage site of the representative µVH (after 175 the 18th residue) and were in less agreement for τVH (after the 176 20th residue) as shown in Supplemental Figs. 1 and 2. Sequences 177 were translated with Expasy translate tool (http://web.exp-178 asy.org/translate/), and the Clustal W program in Bioedit was em-179 ployed to align amino acid sequences (http://www.mbio.ncsu.edu/ 180 bioedit/bioedit.html) for figures. Sequences were managed and 181 assembled in Bioedit and have been deposited in Genbank under 182 accession numbers KF713322-KF713372. CDR3 length was calcu-183 lated using the "CDR3 length = exclusive number of amino acids 184 from C (of VH segment $Y \times C$) to F (of JH segment $FG \times G$)" IMGT 185 formula (Lefranc et al., 2003). 186

2.5. Phylogenetic studies

Amino acid alignments were made with ClustalW employing 188 gap opening penalties of 10 and gap extension penalties of 0.1 189 for pairwise alignments, then 0.2 for multiple alignments using a 190 Dayhoff matrix based method (Schwarz and Dayhoff, 1979). Phylo-191 genetic trees were constructed using MEGA 5 software (Tamura 192 et al., 2011). Neighbor joining trees using the substitution method 193 of Jones et al. (1992) and pairwise deletion of empty positions were 194 constructed from alignments of VH and CH domain sequences. 195 Trees were bootstrapped 1000 times (Koichiro Tamura et al., 196 2011) and were viewed and adjusted using the Treeview Software 197 (Page, 2002). 198

2.6. Real time quantitative PCR

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Oligo-dT transcribed cDNA samples from spleen, gill and 200 anterior kidney were assayed for levels of μ and τ message using 201

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Fig. 1. Four VH families used by tuna IgH. Amino acid alignment of VH segment encoded sequences found within *T. orientalis* μ and τ cDNAs. Clone numbers are shown to the left and VH family designations are shown to the right. VH gene segment families were ascribed based upon 70% nucleotide identity in a pairwise matrix (Supplemental Fig. 2). Gaps introduced into the alignments are indicated by dashes ("-") and identity to the first sequence is indicated by a period (".") in the column. CDR1 and CDR2 are indicated below the alignment. If clone contained CH region encoding region, μ or τ is indicated at left of sequence after the clone name.

202 β-actin as a constitutively expressed control. Real-time PCR reac-203 tions were performed using 25 and 50 ng of cDNA with SYBR 204 Advantage qPCR Premix (Clontech, Mountain View, CA) per the 205 manufacturer's instructions. Primers were designed to span across 206 introns. Using a Roche LightCycler 480 a three-step thermal cycling 207 program was followed: 1 cycle at 95 °C for 5 min, then 45 cycles of 95 °C for 10 s, then 60 °C for 5 s, then 72 °C for 5 s. The Roche 208 LightCycler software was utilized for raw data acquisition and cal-209 culation of Ct (threshold cycle) values. Changes in gene expression 210 were estimated using the $2^{-\Delta\Delta Ct}$ method (Livak and Schmittgen, 211 2001), with $\beta\text{-actin}$ utilized as the stable reference gene for all 212 O3 experimental situations. The fold changes in gene expression were 213 214 calculated with respect to the expression level of the genes in the anterior kidney (the primary B lymphopoietic tissue of bony fish). 215

216 **3. Results**

217 3.1. Characterization of μ cDNA of T. orientalis

The initial full length tuna µ was cloned and sequenced using a
cDNA RACE library that was obtained from RNA pooled from several tuna anterior kidney, spleen and gill samples. The secretory
tuna µ sequence shown in Supplemental Fig. 1 is an 1827 bp open
reading frame which encodes a 609 amino acid protein containing
a leader peptide of 18 residues, one Ig VH and four CH domains.

The primary amino acid sequence showed two cysteine residues 224 (and intervening tryptophan) conserved for intra-domain disulfide 225 bond formation present in each of the Ig domains with the cyste-226 ines being spaced by approximately 70 residues in the VH domain 227 and 60 in the CH domains. The amino-terminal cysteine in the CH1 228 domain forms an interdomain disulfide bond between the IgH 229 chain to the IgL chain. The potential N-linked glycosylation site 230 near the carboxyl terminus of the IgM chain was found at this 231 232 position of the tuna IgM (Danilova et al., 2005a,b).

3.2. Characterization of tuna IgT

While sequencing 3' RACE PCR products employing VH primers designed from μ clones we found other clones with Ig CH region amino acid sequences distinct from IgM, although they often shared a VH domain highly homologous with μ clones. The CH1 domain of these clones shares 56% amino acid identity with the CH1 of *S. chuatsi*. More primers allowed the complete cloning of the IgT encoding cDNAs, with CH3 proving to be even more definitively of the isotype (60% identical amino acids to *S. chuatsi*) (Supplemental Fig. 2).

The secretory tuna τ cDNA is composed of 1614 base pairs translating to 539 amino acids forming a leader peptide, VH domain and four CH domains. As in tuna IgM, two conserved cysteine residues and one tryptophan were identified in the VH and each CH

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		_	~						a 1		
		Tyr	Cy	TATA	CGGGGGGGGGGTACTO	3GG		1y	GLY		
1	VH1	TATTA	CTG	GCCAGATCCCCCC	GTAGTAG		CGCTGTTTTTGACTACTGG	GAAA	AGGCACG	J1	μ
2	VH1	TATTA	CTG	GCCAGAAGA	AGAGGGGGTACCCG-		ATTGACTACTGG	GAAA	AGGAACA	J8	μ
3	VH1	TATTA	CTG	GCCAGAGACCG	· GGC <mark>GTAC</mark> AA	ACGACGGCGCGC	TTGACTACTGG	GAAA	AGGAACA	J6	μ
4	VH1	TATTA	CTG	GCCAGAGGATACAGCTAC	CGGGAGTGGCGA		CTGGGCTTTTGACTACTGG	GAAA	AGGCACT	J4	μ
6	VH1	TAT TA	CTG	GCCTCC	CGATCAA <mark>GTA</mark> TTI	[A	CTATGCTTTTGACTACTGG <mark>G</mark>	<mark>GG</mark> AA	A <mark>GGC</mark> ACG	J3	μ
7	VH1	TAT TA	CTG	GCCAGAGACCGCTCC	CACAGT <mark>GGGT</mark> GGG•		TTTGACTACTGG <mark>G</mark>	<mark>GA</mark> AA	AGGGACC	J5	μ
9	VH1	TAT TA	CTG	GCCAGGCGCGAT	CGGGGGAG		AGATGCTTTTGACTACTGG <mark>G</mark>	<mark>GA</mark> AA	A <mark>GGC</mark> ACG	J1	μ
10	VH1	TAT TA	CTG	GCCAGAGAA	A <mark>GGGG</mark> TAA		CGATGCTTTTGACTACTGG <mark>G</mark>	<mark>GA</mark> AA	A <mark>GGC</mark> ACG	J1	μ
11	VH1	TAT TA	CTG	GCCAGAG	CICCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCCC	}	TTTGACTACTGG <mark>G</mark>	<mark>GA</mark> AA	A <mark>GGA</mark> ACA	J6	μ
12	VH1	TAT TA	C <mark>TG</mark>	GCCAGCAACCCCGTATAC	CGGGGGGTCCGGCT	BAC	TACTTTGACTACTGG <mark>G</mark>	<mark>GA</mark> AA	A <mark>GGG</mark> ACT	J5	μ
13	VH1	<mark>TAT</mark> TA	C <mark>TG</mark>	GCCAGAGACC	CCCGGGGGGGGTACT		CTGGGCTTTTGACTACTGG <mark>G</mark>	<mark>GA</mark> AA	A <mark>GGC</mark> ACT	J4	μ
14	VH1	<mark>tat</mark> ta	C <mark>TG</mark>	GCCAGAGAC <mark>ATAC</mark>	CGGGGGGGTCCG		CGATGCTTTTGACTACTGG <mark>G</mark>	<mark>GA</mark> AA	A <mark>GGC</mark> ACG	J1	μ
15	VH1	<mark>TAT</mark> TA	C <mark>TG</mark>	GCCAGAATCCCGGC	CAC <mark>GGGGGGGTAC</mark> C	CTACGGC	TTTGACTACTGG <mark>G</mark>	<mark>GA</mark> AA	A <mark>GGA</mark> ACA	J7	μ
16	VH1	<mark>TAT</mark> TA	C <mark>TG</mark>	GCCAGAAGGAGCGGGT	ATAC <mark>GGGGGT</mark> CAA ·		CTGGTCTTTTGACTACTGG <mark>G</mark>	<mark>GA</mark> AA	A <mark>GGC</mark> ACT	J4	μ
17	VH1	<mark>tat</mark> ta	.C <mark>TG</mark>	GCCAGGCGATC	CGGGGGG		CGATGCTTTTGACTACTGG <mark>G</mark>	<mark>GA</mark> AA	A <mark>GGC</mark> ACG	J1	μ
31	VH1	<mark>tat</mark> ta	.C <mark>TG</mark>	GCCAGAGCCC	GTGGCA <mark>GGT</mark> ·		ATGCTTTTGACTACTGG <mark>G</mark>	<mark>GA</mark> AA	A <mark>GGC</mark> ACG	J1	μ
33	VH1	<mark>tat</mark> ta	.C <mark>TG</mark>	GCCAAAG(GA <mark>GGGG</mark> ACGGCCG	TAC	TTTAACTACTGG <mark>G</mark>	<mark>GA</mark> AA	A <mark>GGG</mark> ACC	J5	μ
41	VH1	<mark>tat</mark> ta	.C <mark>TG</mark>	GCCAGAG	GGGG <mark>ACT</mark>	<mark>G</mark> AGCCGCGC	TACTTTGCCTACTGG <mark>G</mark>	<mark>GA</mark> AA	A <mark>GGG</mark> ACC	J5	μ
42	VH1	TAT TA	C <mark>TG</mark>	GCCAGGCAGAGCTC <mark>TAC</mark>	C <mark>GGGG</mark> CACTA ·		CTTTGACTACTGG <mark>G</mark>	<mark>GA</mark> AA	A <mark>GGA</mark> ACA	J6	μ
51	VH1	TAT TA	.C <mark>TG</mark>	GCCAGA	CGGGGGGAATCCCG	CAT	TTTGCCTACTGG <mark>G</mark>	<mark>GT</mark> CG	T <mark>GGG</mark> ACT	J9	τ
53	VH1	TAT TA	.C <mark>TG</mark>	GCCAGACTC	CGGACT <mark>GGGT</mark> GGC(CTTTTT	TTTGAGTACTGG <mark>G</mark>	<mark>GT</mark> CG	T <mark>GGG</mark> ACT	J9	τ
54	VH1	TAT TA	.C <mark>TG</mark>	GCCAGA	AT <mark>GTACT(</mark>	<mark>GGG</mark> GCGTC		<mark>GT</mark> CG	T <mark>GGG</mark> ACT	J9	τ
55	VH1	TAT TA	.C <mark>TG</mark>	GCCTAGAAC	CTTCT <mark>GGGGT</mark> GGC(GCCTTT	TATTTTGACTACTGG <mark>G</mark>	<mark>GT</mark> CG	T <mark>GGG</mark> ACT	J9	τ
56	VH1	TAT TA	CTG	GCCAGAGACCG		GA	TATTTTGACTACTGG <mark>G</mark>	GTCG	T <mark>GGG</mark> ACT	J9	τ
5	VH2	TAT TA	TTG.	GCGCGTGAT	GGCTATCG <mark>TC</mark>	GCTATTATGAA	TTTGATTATTGG <mark>G</mark>	<mark>GC</mark> AA	A <mark>GGC</mark> ACC	J6	μ
52	VH2	TATTA	.C <mark>TG</mark>	GCCAGA	CTT <mark>GTAC</mark> TC	GGAGAATAT	TATTTTGACTACTGG <mark>G</mark>	<mark>gt</mark> cg	T <mark>GGG</mark> ACT	J9	τ
18	VH3	TAT TA	TTG	GCTCGAC	<mark>GGGGGGTAC</mark> GA	AGC	CTATGCTTTTGACTACTGG <mark>C</mark>	GGAA	A <mark>GGC</mark> ACG	J2	μ
19	VH3	TATTA	TTG	GCTCGAGAC	CGGGGGCGGGGATGC	3	TTTTGACTACTGG <mark>G</mark>	GAAA	a <mark>ggc</mark> acg	J1	μ
20	VH3	TAT TA	TTG	GCTCGAGACTATATC	CGGGGAACAAT		GGTTTTGACTACTGG <mark>G</mark>	GAAA	a <mark>ggc</mark> acg	J1	μ
21	VH3	TATTA	TTG	GCTCGAGACTACTACG	T <mark>GGG</mark> CGCCGC		GGCTTTGACTACTGG <mark>G</mark>	GAAA	A <mark>GGA</mark> ACA	J6	μ
22	VH3	TATTA	TTG	GTTCGAGGCT	CCAGTGGGTGGGG	CTAC	TTTGCCTACTGG	GAAA	AGGGACC	J5	ů
27	VH3	TATTA	TTG	GCTCGAG	<mark>GGGGT</mark> CGCC	:G	CTTTGACTACTGG	GAAA	A <mark>GGA</mark> ACA	J6	ů
28	VH3	TATTA	TTG	GCCCGAGACACTTCCG	CGGGTGGCGAGT - ·		ACTTTGACTACTGG	GAAA	a <mark>gga</mark> aca	J6	ů
30	VH3	TATTA	TTG	GCTCGAGGATCA	TGGGT <mark>GGGT</mark> GGC	CCA	CTTTGACTACTGG	GAAA	a <mark>gga</mark> aca	J6	ů
32	VH3	TATTA	TTG	GCTCGAGA TAGGAGCTAC	GGGAGCAACAAT		GCTTTTGACTACTGG	GAAA	AGGCACG	J1	ū
43	VH3	TATTA	TTG	GCTCGAGA GGAGGGCTCTAC	GGGAAC		TTTGCCTACTGG	GAAA	AGGGACC	J10	ū
44	VH3	TATTA	TTG	GCTCGAGC	GGGGGACTGGAAT	GCGGAGGATTG	GCTTTTGACTACTGG	GGAA	AGGCACG	J1	ū
45	VH3	TATTA	TTG	GCTCG	GGGGCCAAATTA	TACGGTGCTC		GAAA	GGGTACA	J11	ū
46	VH3	TATTA	TTG	GCTCGAGAC	GTAGACT	C		GAAA	AGGGACC	JT10	
47	VH3	TATTA	TTG	GCTCGAGA TAGGAGCTA	CGGAGTAACAAT.		GCTTTTGACTACTGG	GAAA	AGGCACG	.T4	
48	VH3	TATTA	TTG	GCTCGAGGATACAATACAC	GGGGGGGGGGAGTCO	GTCGTCTACTA	CTTTGACTACTGG	GAAA	AGGAACA	.16	
49	VH3	TATTA	CTG	GCCAGAGAGCGGCCTACCG	GGGCTAC			GAAA	AGGACC	.15	
57	VH3	TATTA	CTG	CCAGAG	-TGGAGGGTCAGT	·CT	TTTGACTACTGG	anca	TagaaacT		<u>م</u>
59	VH3	TATTA	TTC	GCTCGAGAC	GTACCO	····	TTTGACTACTCC	GTCG	TGGGACT	.19	÷
60	VH3	TATTA	TTC	GCTCGAG	CTGGGGCAAT	,	ATTTTGACTACTCC	GTCG	TGGGACT	.19	÷
61	VH3		TTC	GCTCGAGACTACAC			TTTTGACTACTGC		TGGGACT		÷
62	VH3		TTC	CCTCCAGAC					TagaaacT		÷
29	WH4		CTC	CCCAGAGCT			TCCTTTTCACTACTCC		AGGCACC	.T2	
29	*114	TAT IA	10 17	N/D		N/D		<u></u>		52	μ
			v	N/P	<u>~</u>	14/1	J				

Fig. 2. Tuna VH(DH)JH junctional diversity. Nucleotide alignment arranged by VH family VH(DH)JH junctional region. Clone names and VH family are denoted on the left, JH gene and CH region is given to the right. Conserved tyrosine and cysteine codons of Y × C motif of VH segment as well as G × G glycines of JH gene are highlighted in yellow. Predicted DH segment is highlighted in blue. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

domain which are important for folding of the β-sandwich Ig do-247 248 mains. There is also one conserved cysteine in the CH1 domain 249 which forms a disulfide covalent linkage between the IgH chain 250 to the IgL chain. The secretory tail of tuna IgT is composed of 12 251 amino acids.

252 3.3. IgH μ and τ VH, DH and IH segments

The same cDNA pools were used to examine tuna IgH μ and τ 253 VH(DH)JH rearrangement diversity. In total 50 different sequences 254 encoding VH domains (Fig. 1) that possessed full or partial unique 255 256 VH regions were cloned, 36 spliced to μ CH regions and 11 with τ (three contained complete VH regions but were incompletely rear-257 ranged or did not splice to a CH). Based on percent identity the VH 258 segment sequences were divided into four separate families of IgH 259 V genes. Members of each family were more than 70% identical in 260 their nucleotide sequences (Brodeur and Riblet, 1984; Pascual and 261 Capra, 1991) (Supplemental Fig. 3). Analysis of the carboxy-termi-262 263 nal portion of the VH domains gave insight into the DH and JH gene segments used to rearrange mature VH exons. We predicted 11 dif-264 265 ferent JH segments used in these clones and one DH segment (TATACGGGGGGGGGGGTACTGGG) could be identified in the 48 unique 266 CDR3 encoding rearrangements analyzed (Fig. 2). The one DH seg-267 ment apparently was employed by both isotopes, as various 268 stretches of the sequence (including portions at each end) are 269 270 found in both μ and τ clones. The predicted DH germline nucleo-271 tide contribution to the final expressed CDR3 encoding sequence 272 ranges from 3 to 10 with a mean of 5.5 base pairs. All three reading

frame of the D segment were used (Fig. 3). The τ clones all used a 273 dedicated JH segment (J9). 274

IgH CDR3 is the crucial loop in the paratope of most antibody 275 antigen interactions. This sample of the tuna Ig heavy chains ex-276 pressed at the mRNA level allowed an initial analysis of the length 277 of IgH CDR3 of μ and τ . Table 1 shows that tuna μ clones display a 278 broader range of CDR3 lengths (from 9 to 18aa) as well as an aver-279 age of one amino acid longer CDR3 length than those found in tuna 280 281 τ.

3.4. IgM and IgT CH regions of tuna

The tuna IgM CH region amino acid sequence showed the most 283 identity to the mandarin fish (S. chuatsi, also known as the Chinese 284 perch and also a member of the Order Perciformes) with 53.6% and then to the rainbow trout with 39.8% identity and presented the least with chicken (23.8% identity) amongst the sequences we included in our analysis (Fig. 4). The tuna IgT CH region has the 288 highest identity also to that of the mandarin fish with 52.5% and 289 the least to grass carp with 20.4% (Fig. 5). Unlike the cyprinid grass 290 carp and zebrafish, the CH3 domain of tuna IgT conforms to the 291 canonical immunoglobulin domain fold with cysteines and trypto-292 phans in positions common for β -sandwich tertiary structure. 293

3.5. Phylogenetic analysis

To assess the phylogenetic relationship of the tuna Ig VH gene 295 segments with those of other teleosts we created dendrograms 296

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1	VH1	<mark>Y</mark> Y <mark>C</mark> ARSP <mark>R</mark> SSAVFDYW <mark>G</mark> K <mark>G</mark> TMVTVTSA	J1	<mark>2</mark> M
10	VH1	YY <mark>C</mark> ARE <mark>RG</mark> NDAFDYW <mark>G</mark> K <mark>G</mark> TMVTVTSA	J1	<mark>2</mark> M
9	VH1	YY <mark>C</mark> ARRD <mark>RG</mark> RDAFDYW <mark>G</mark> K <mark>G</mark> TMVTVTSA	J1	<mark>2</mark> M
14	VH1	YY <mark>C</mark> ARD <mark>IRG</mark> VRDAFDYW <mark>G</mark> K <mark>G</mark> TMVTVTSA	J1	<mark>2</mark> M
17	VH1	YY <mark>C</mark> ARRS <mark>GG</mark> DAFDYW <mark>G</mark> K <mark>G</mark> TMVTVTSA	J1	<mark>2</mark> M
32	VH3	YYCARDRS <mark>YG</mark> SNNAFDYW <mark>G</mark> KGTMVTVTSA	J1	<mark>З</mark> М
44	VH3	YY <mark>C</mark> ARAGDWNSGGLAFDYW <mark>G</mark> KGTMVTVTSA	J1	<mark>з</mark> м
19	VH3	YYCARDRGRDAFDYWGKGTMVTVTSA	J1	<mark>2</mark> M
20	VH3	YYCARDYI <mark>YTG</mark> NNGFDYW <mark>G</mark> K <mark>G</mark> TMVTVTSA	J1	1 M
31	VH1	YY <mark>C</mark> ARAGGRYAFDYW <mark>G</mark> KGTMVTVTSA	J1	<mark>1</mark> М
29	VH4	YY <mark>C</mark> ARALWQ <mark>L</mark> AFDYW <mark>G</mark> KGTMVTVTSA	J2	1 M
18	VH3	<mark>Y</mark> Y <mark>C</mark> ARR <mark>GV</mark> RAYAFDYW <mark>G</mark> K <mark>G</mark> TMVTVTSA	J2	1 M
6	VH1	YYCASRSSIYYAFDYW <mark>G</mark> KGTMVTVTSA	J3	<mark>2</mark> M
4	VH1	YY <mark>C</mark> ARGYS <mark>YG</mark> SGDWAFDYW <mark>G</mark> K <mark>G</mark> TMVTVTSA	J4	<mark>З</mark> М
16	VH1	YYCARRSGYT <mark>GV</mark> NWSFDYW <mark>G</mark> KGTMVTVTSA	J4	<mark>1</mark> м
13	VH1	YYCARDPRGVLWAFDYWGKGTMVTVTSA	J4	1 M
47	VH3	YYCARDRS <mark>YC</mark> SNNAFDYW <mark>G</mark> KGTMVTVTSA	J4	<mark>З</mark> М
12	VH1	YYCASNPVYGGPADYFDYWGKGTQVTVTSA	J5	<mark>1</mark> М
7	VH1	YYCARDRSTVCGFDYWGKGTQVTVTSA	J5	<mark>2</mark> M
33	VH1	YYCAKGGDGRYFNYWGKGTQVTVTSA	J5	<mark>2</mark> M
41	VH1	YYCARGGLEPRYFAYGWGKGTQVTVTSA	J5	1 м
22	VH3	YYCVRGSSCWAYFAYWGKCTQVTVTSA	J5	<mark>1</mark> М
49	VH3	YYCARERPTACYFAYWGKGTQVTVTSA	J5	1 M
3	VH1	YYCARDRAYNDGALDYWGKGTTVTVTAA	J6	<mark>З</mark> М
5	VH2	YYCARDGYRCYYEFDYWCKGTTVTVTAA	J6	<mark>з</mark> м
11	VH1	YYCARAGGWGFDYWGKGTTVTVTAA	J6	<mark>2</mark> M
42	VH1	YYCAROSS <mark>TG</mark> HYFDYWGKGTTVTVTAA	J6	1 м
48	VH3	YYCARGYNTLGGESVVYYFDYWGKGTTVTVTAA	J6	<mark>1</mark> М
30	VH3	YYCARGSVG <mark>G</mark> WPHFDYW <mark>G</mark> KGTTVTVTAA	J6	<mark>1</mark> М
27	VH3	YYCARGGRRFDYWGKGTTVTVTAA	J6	<mark>З</mark> М
28	VH3	YYCARDTSACGEYFDYWCKCTTVTVTAA	J6	1 м
21	VH3	YYCARDYYDCRRGFDYWCKCTTVTVTAA	J6	<mark>1</mark> М
15	VH1	YYCARIPARGVPYGFDYWGKGTTLTVTEA	J7	<mark>1</mark> М
2	VH1	YYCARREGYPIDYWGKGTTVTVTDS	J8	3 м
51	VH1	YYCARSAGESRHFAYWGRGTEVTVSSE	J9	3 т
52	VH2	YYCARLVLGEYYFDYWGRGTEVTVSSE	J9	1 т
53	VH1	YYCARLGLGGLFFEYWGRGTEVTVSSE	J9	<mark>1</mark> т
54	VH1	YYCARMYWGVFDYWGRGTGVTVSSE	J9	1 т
55	VH1	YYCA*TSGVAAFYFDYWGKGTEVTVSSE	J9	1 т
56	VH1	YYCARDRGGRRYFDYWGRGTEVTVSSE	J9	<mark>1</mark> т
57	VH3	YYCARVEGOSFDYWGRGTEVTVSSE	J9	2 T
59	Vh3	YYCARD	J9	1 т
60	VH3	YYCARVLGOYFDYWGRGTEVTVSSE	J9	1 T
61	VH3	YYCARDYTGGGFDYWGRGTEVTVSSE	J9	1 T
62	VH3	YYCARDRGLVGFDYWGRGTEVTVSSE	J9	2 T
43	VH3	YYCAREEGSTGNFAYWGKGTOVRVTSA	J10	<u>з</u> м
46	VH3	YYCARDVDYFDYWGKGTOVTVTSA	J10	2 M
45	VH3	YYCARGPNYYGALDYWGKGTTVTVSSA	J11	1 м
		V N/P D N/P J		
		.,, = = -,, = 0		

Fig. 3. Translated complementarity determining region (CDR) 3 repertoire sampling of tuna IgH. Amino acid alignment arranged by JH gene of the VH(DH)JH junctional region. Clone names and VH family are denoted on the left, JH gene, reading frame of DH used and CH region is given to the right. Conserved tyrosine and cysteine of $Y \times C$ motif of VH segment as well as $G \times G$ glycines of JH gene are highlighted in yellow. Predicted DH segment is highlighted in green, blue or magenta depending on the use of reading frame one, two or three, respectively, in panel B. Amino acids were assigned to VH, DH or JH based on at least two bases of codon matching consensus, grey highlighting indicates a residue partially encoded by D consensus that does not encode consensus amino acid. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

with their pairwise genetic distances (Fig. 6). The four tuna IgH V 297 gene families interleaved amongst those VH segment sequences 298 used by the other fish, indicating that they are using members of 299 300 the same ancient VH families that have been conserved by trans-301 species maintenance since at least the common ancestor of these divergent fish. However tuna families VH1 and VH2 appear to have 302 arisen from a more recent duplication in the order Perciformes. 303

We also explored the relationship of these new tuna IgH C 304 305 regions to those of other fish and other vertebrates (Fig. 7). As expected, the tuna IgM grouped with that isotype from other fish, 306 307 most closely the Perciforme trumpeter fish (Latris lineata). Within 308 the IgM, teleosts group together as a sister group to all of the other 309 vertebrates, including the shark which shares a more ancient com-310 mon ancestor and would be expected to branch outside of teleosts and tetrapods. However this incongruence with the organisms' 311 natural history is not unusual for phylogenetic analyses of teleost 312 antigen receptors, unless balancing numbers of operational taxo-313 314 nomic units fill the other vertebrate classes (Criscitiello and Flajnik, 315 2007). IgT of tuna clusters with that isotype from other represen-316 tative fish.

Table 1

CDR3 lengths in amino acids.

	IgM	IgT
Maximum	18	13
Minimum	9	8
Range	9	5
Median	12	11
Mean	12.49	11.00
Variance	3.31	2.00

3.6. IgH μ and τ relative tissue expression

Quantitative real-time PCR (Supplemental Fig. 4) was used to assess the expression of these isotypes at the mRNA level in secondary lymphoid tissues relative to the anterior kidney (the chief primary lymphoid tissue of fish (Fillatreau et al., 2013; Lam et al., 2004; Trede et al., 2004)). Relative levels of μ were higher than τ in both spleen and gill, but μ did not predominate τ to as great an extent in gill as it did in spleen. The averaged ratio of HCµ to HC τ in tuna spleen was 7.35 compared to 2.89 in the gill.

4. Discussion

4.1. Repertoire

The 20–30% sequence disparity between some VH family members in tuna suggests either ample somatic hypermutation for affinity maturation of these fish antibodies or an older divergence date of VH family members than has been seen in some other teleosts such as stickleback (Gambon-Deza et al., 2010). Families VH1 and VH2 share between 52% and 64% nucleotide identity (Supplemental Fig. 3) and appear recently diverged (Fig. 6), perhaps within a Perciformes branch including tuna.

Despite the initial report that found shorter CDR3 in trout IgM than IgT (Hansen et al., 2005a) we found a small skewing towards shorter IgT CDR3 (Table 1). We predict that this may be an effect of a different immunogenetic rearrangement mechanism involving a single shared DH gene segment that governs τ versus μ/δ in a clade including tuna and other fish (more below). IgH CDR3 often dominates antigen recognition properties of the six CDRs comprising the F_{ab} paratope (Davis, 2004; Xu and Davis, 2000). The three reading frames usually supplied by DH gene segments therefore contribute significantly to the eventual translated repertoire of antigenic specificities. Additionally, extended length of IgH CDR3 has been crucial in many clinically important antibodies against viral scourges (Kwong and Wilson, 2009; McLellan et al., 2011), and the loop has evolved into an entirely new domain in some antibodies of cattle (Wang et al., 2013). Thus, restricting the entire repertoire to rearrangements based on a single DH would be expected to place constraints on antigen recognition.

As Perciformes, tuna belong to the largest order of vertebrates that accounts for approximately 40% of all bony fishes. As T. orientalis is the first Perciformes to have either their IgM and T repertoire or IgH locus analyzed immunogenetically, there may be a great many fish that employ this system for Ig isotype control and B lineage commitment. As successful as the Perciformes have been in radiating to occupy most fresh and saltwater niches on Earth, the potential restriction in CDR3 length variability must not have too great a toll on the fitness of these fish.

4.2. Genomic organization

The generalized translocon configuration of the teleost IgH 363 locus with a set of VH genes and downstream μ and δ CH regions has been confirmed in many studies (Bengten et al., 2002;

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	CH1→
<pre>T. orientalis S. chautsi I. punctatus D. rerio L. lineata O. mykiss P. olivaceus S. salar G. cirratum X. laevis A. platyrhyncho B. taurus H. sapiens</pre>	PRAPTLEPUAGGSG IGDMVTLGCLAR SFTPSS-VTFSMT-KSGTALTDFTOYEPVOKGEFYEGVSOVOVRODWEAKOP-S-KCTVIHPLGT-SSVTPYSEP ISTGPTVFPLMCGGSG IGDMVTLGCLARGFTPSS-LTYAMS-KNGAALTDSIOYEPVOKGDVTRVSOIRVRICDWDARES-F-RCAVHEPAGN-GKADFMARKV -sAPKSLFFVMCGGSA SDGLVTLGCVTDLLSADGLSFIHKDASGSALTDVVOYEPVOATGCYTSVSBUKVKASDWNGNKK-F-TGGVKNCLGS-KDASIGAPFAA SAPCSVFGISGSSG GDGGITLGCLARGFFPASDLSFKKKDPJGGLSUDVYOYEPVOATGCYTSVSBUKVKASDWNGNKK-F-TGGVKNCLGS-KDASIGAPFAA PTTFFLMCGGGCSECUVLGCLARGFFPASDLSFKKNDSGGSSLTDVVOYEPVOATGCYTKUSBICVKKASDWNGNKK-F-TGGVKNCLGS-KDASIGAPFAA PTTFFLMCGGGCSECUVLGCLARGFFPAS-LIFYKNS-KNEVALNDFIOYEPVIKGNLYGCISQIKVSRDDWEAKPNTIKCAVHHAAGN-AQCDFRDHHV FFLMCGGGCGGVTLGCLARGFFPAS-LIFYKNDSGGSSLTDFVOYEPVIKGNLYGCISQIKVRBDWD-SKKFRCAVEBAGS-KVVVKOPE MPAIFTLFPUMCGGGCNGOMNTLGCLARGFFPAS-LIFFKNDGGGNSLTDFVOYEPVIGGSSVGVVCNGCDEVKRBDWD-SKKFRCAVEBAGS-KVVVKQPE SSTAPTLFPUAGGSGCNGOMNTLGCLARGFFPAS-LIFFKNDGGGNSLTDFVOYEPVIGGSSVGVVCNGVGVLKRBDWD-SKKFRCAVEBAGS-KVVVKQPE SSTAPTLFPUAGGSGCNGOMNTLGCLARGFFPAS-LIFFKNDGGSGNSLTDFVOYEPVIGGSSVGVVCNGVGVLKRBDWD-SKIFRCAVEBAGS-KVVVKQAE -PSSPTYGVSSCQOUNDGSVFGCLANGFFPAS-LIFFKNDGGSGNSLTDFVOYEPVIGGSSVGVVCNGVGVLKRBDWD-SKIFRCAVEBAGS-KVVVKQAE -PSSPTYFLVSCSSSSSSSNAVGCVAVGHVPAG-VYFSNTDVNAVATTIV-NFPARSGCJAVBSSDVVVKRADWD-SKIPSCOVMEBAGS-KVVVKQAE -PSSPTYFLSCGSSSSSSNAVGCVAVGHVPAG-VYFSNTDVTNANATTIV-NFPARSGGNNAASRLEDLDGGSGRGROFFQCMRBGSDKSGMP FGSPTLFPLISCGSSSSSSNAVGCVAVGHVPAG-VYFSNTDVTNANATTIV-NFPARSGCGNNAASRLEDLDLOGGKGROFFQCGARBGSDKSGMP FSCSPTLFFLISCGSSSSSSNAVGCVAVGHVPAG-VYFSNTDVTNANATTIV-NFPEARSGCGNNAASRLEDLDLOGGKGROFFQCGARBGSDKSGMP FSCSPTLFPLISCGSSSSSSNAVGCVAVGHVPAG-VYFSNTDVTNANATTIV-NFPEARSGCGNNAASRLEDLDLOGGKGROFFQCGARBGSDKSGGMP FSCSPTLFPLISCGSSSSSSNAVGCVAVGHVPAG-VYFSNTDVTNANATTIV-NFPEARSGCGNNAASRLEDLDLOGGKGROFFQCGARBGSDKSGGMP FSCSPTLFPLISCGSSSSSSNAVGCVAVGHVPAG-VYFSNTDVTNANATTIV-NFPEARSGCGNNAASRLEDLDLOGGKGROFFQCGARBFGSDXSGG
	CH2→
T. orientalis S. chautsi I. punctatus D. rerio L. lineata O. mykiss P. olivaceus S. salar G. cirratum X. laevis A. platyrhyncho B. taurus H. sapiens	LFELPT-LKULASTDDGSEASFSCFARVFSPENE-IKKULKDGDVDDVBVVGIKTFVIGKLVSVASFLVVFSDLRPNSMEFTCEFEKLNEGAFVNSTVTVGGSCPEPTCCE TVVPPTELKULASSGEQEASFSCFARDFSFKDE-IKKULKNEAEIPNKIYEIKKUFIKSCOTTDGKLVSVASFLVVPASEM-TVDRKFTCEFEKLNEGAFVNSTVTVGGSCPEPTCCE R-ELHASLLITTFTCTEIDNGTATFYCIATFSSFKSH-FKWTENKDEIPNKIYEIKNELGSVDKNGTLVSVASSFLVPASSM-TVDRKCDEFECKENGATFMNSSVTYKHTTP-GNCE R-ELHASLLITTFTCTEIDNGTATFYCIATFSSFKSH-FKWTENKDEIPNKIYEINKUKNIVSONGNFTALSVLESASSWTSSTBVCGEOCANHVFKASSTYNSTTYK-GTC PPDLLATVIGTAFTTKELEGGSAFFMCIARFSFKOYE-FKWYENDOEWNNYDDFFKDEKNGSVEYSATSILKINAEUWKQAESKWCVFEHNKN,DSRLOXKDTWDCI PCDLPTLKVJASSDESEASFSCFANDFSFRTH-IKWNENEVEIPNKIYEVTPACO-RDLNGSULYSASSLLVSESKWCDEFEKCKTSSTFKNSTVTK-DCKS-DVCD YLOOPSLVWTFSKEEMSENTTASFACFANDFSFRTH-IKWNENECTCEVVSDFKSSCESEKKSETLISTITSTE-HKSNGLVSASSEVTFCVFENNAGNVFKASTVTYK-DCKS-DVCD YLOOPSLVWTFSKEEMSENTTASFACFANDFSFRTH-IKWNENECTCEVVSDFKSSCESEKKSETLISTITSTE-HKSNGLVSASSEVTFCVFENNAGNVFKASTVTYK-DCKS-DVCD YLOOPSLVWTFSKEEMSENTTASFACFANDFSFRTHT-IKWNENECTEVSDFKSSCESEKKSETLISTITSTE-HKSNGLVSATSTLVNESEWKSEZVFTCVFENNAGNVFKTVTYK-DTCKGCM YLOOPSLVWTTSKEEMSENTTASFACFANDFSFRTHT-IKWNENECTEVSDFKSSCESEKKSKRLVSITSTSTSTK-PKNVTSDFVVTVCOVFENNAGNVFKTVTYK-DTCKGCM YLOOPSLVWTTSKEEMSENTTASFACFANDFSFRTH-IKWNENECTGSVHSDITSTE-HKSNGLVSATSTLVNESEWKSEZVFTCVFENNAGNVFRTVGTT-SSDAGPVH CGNGDFTVULTVSSEETESIKFATLVCSIDHSSTSS-VS-VSTD-VSTGAVEFVSSCESEKKSCNCDYTTGTV-VVTTOGVCVGFCSDAGPVH VICHPSKDALALNESLFIVCIANNFFFNTVVK-DTKSSCESEVSDFKSSCESEKKSCHVYSTSTSTSTSTSTSTSTVVVTOVFENNAGNVRTVGTT-SSDAGPVH VSSOFTAPVVSIHPSKDALALNESLFIVCIANNFFFNTVVK-DKKSCSCHDDIITSFVCVVSVGSSATSELVVTAEN-PDKAVTCOVVENASSLOEKNMSSLMCDFTT SSOFTAPVVSIHPSRDALALNESLFIVCIANNFFFNTVVK-NGKGAVEFVSDVKGSSCESEVSDFKSCASSUFCAVGFCANDESSLOEKNDSSCHVCPDY VIAELPVSVFVPFPNSLSCDGNSKSSLCCATDFSFK0TE-ISPKCTEVSSCVXEPVGSSVTJDVQAEAKESGFT-TVTSAEN-DAGTVTCQMADEMRNSKCSLOEKDKDS KAEVLSVVSVFVPPPNSSCDGGSNSSLCCATDFSFK0TE-ISPKCTATSSCVXEPVG-SSVTTDQVQAEAKESGFT-TVKVTSTLTTKESDN-LSONYTCOVENDEGSCUTATCTVDHKGAUSKC-
	CH3→
T. orientalis S.chautsi I. punctatus D. rerio L. lineata O. mykiss P. olivaceus S. salar G. cirratum X. laevis A. platyrhyncho B. taurus H. sapiens	VLDVEVETKGPTNIDMFVESKGTIVCOVKINKEOVITTEMEDEDGKSMIESVPA-DGFKGTVNVFLDITYDEWNAGIRXGVVOHINFL-EPIKRVYERKIV V-DUDIKITCPTTLADMFUNRSCTIVCOVKINEPIVGRIUMEDEKGNENGABSKTNDEGT-S-LHFEITYDEWSGCHRXGVVEHENLI-EPIKELYERSFG OP-OVKITCPSTEDLIKRACQUECREGDFOFKSIKKLICNREISISNISSKTNDEGT-S-LHFEITYDEWSGCHRXGVVEHENLI-EPIKEVFNERG DDNVHIDTIFSGPAVEDMFNRSCTITCUVNSS-EBIGKIMMEDGHGME
	CH4→ ^ ^
T. orientalis S. chautsi I. punctatus D. rerio L. lineata O. mykiss P. olivaceus S. salar G. cirratum X. laevis A. platyrhyncho B. taurus H. sapiens	GLEOHPSVFMLVDVEQANKETVTLTCFVKGFYPKEVIVSWLVDDVBADSNYDISTTNPVESNGFVSVGQLTLSLDDWKDSDKVYSCVVHGSVKNTTKATVKSMAHGST GQTQRPAVFMLPPVEHTREKTVTLTCFVKDFPQEVIVMALUDDEADSKYVFTTNPVESNGSVEAVGQLSLSLEOMKKNDVVSCVVHGSVKNTTNATVKST HTTFF HTTFF
	SEC→
 T. orientalis S. chautsi I. punctatus D. rerio L. lineata O. mykiss P. olivaceus S. salar G. cirratum X. laevis A. platyrhyncho B. taurus H. sapiens 	AQTNLVNLNMEVE ENTNLVNLNMNLPESGKAQ KTPTLVNLTTINPSGSGSGSTY JKSSITHLSNTPAQKAQ EXTNLVNLNNNLPETGKAQ ETTNLVNLNNLT-GKAQ STTNLVNLNNLT-GKAQ GKPSGVNLSLNVPSGKAQ GKPTGVNUSLVLSDTASGC GKPTGVNUSLVLSDTASGC GKPTTLVNLSLVLSDTASGCY- GKPTLYNVSLVLSDTASGCY-

Fig. 4. Amino acid sequence alignment of the heavy chain of IgM in different vertebrate species. The conserved (identical and similar) residues are highlighted in black. Arrows indicate CH1-CH4 and the secretory tail. An asterisk (*) is above the conserved cysteine that forms a disulfide bond with the light chain, a carrot (^) is above conserved cysteines that form intra-domain disulfide bonds. Gaps are indicated by dashes. Genbank accession numbers are: AAQ14846.1 Siniperca chuatsi (Chinese perch), A45804 Ictalurus punctatus (channel catfish), AF281480_1 Danio rerio (zebrafish), ADC45388.1 Latris lineata (striped trumpeter), AAW66973.1 Oncorhynchus mykiss (rainbow trout), AF226284_1 Paralichthys olivaceus (flounder), AAB24064.1 Salmo salar (salmon), AAU04507.1 Ginglymostoma cirratum (nurse shark), AAA49774.1 Xenopus laevis (African clawed frog), CAC43280.1 Anas platyrhyncho (duck), AAN60017.1 Bos taurus (cattle), and AAS01769.1 Homo sapiens (human).

Jørgensen, 2000; Samuel Aparicio et al., 2002), but many deviations on the theme are present as catfish and medaka appear to lack τ and many fish have duplications of blocks of the locus (Fillatreau et al., 2013). Although reported in shark (Zhu et al., 2012), class switch recombination (CSR) has not been described in a teleost. However, one study showed that teleost AID could induce CSR in mouse (Barreto et al., 2005).

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373 374 (DH-JH-CH) µ cluster in zebrafish, fugu (Takifugu rubripes) and 375 three-spined stickleback (G. aculeatus) (Danilova et al., 2005a,b, 376 2011; Gambon-Deza et al., 2010; Hansen et al., 2005a), or it is inserted within the VH gene segment array as in rainbow trout (Hansen et al., 2005b). Thus, in these fish the RAG mediated joining of a VH gene segment to either DH of τ or DH of μ/δ will determine whether the developing pro-B lymphocyte (using mammalian convention) becomes an IgT or IgM/D producer. Experiments in zebrafish (Schorpp et al., 2006) and trout (Zhang et al., 2010) have 383

The IgH τ gene together with its dedicated DH–JH gene segments are located between the VH gene segment block and the

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		CH1>
T. S. O. D. C. E.	orientalis chuatsi salar mykiss rerio idella coioides	PESSPTLYPLOCODSDTGNKVTVGCLARDFFEKSIDFOMNDARGTRÖDS-AQYISGONN-KYTGVSVVQVSRSDIKSSYNCSVDHLGRTMAVTVK 93 SVTVASPTLFPLVQCDSGPADKITVGCLARDFYEKSIDFOMNDARGTRÖDS-AQYISGONN-KYTGVSVVQVSRSDIKSSYNCSVDHLGRTMAVTVK 93 -AATAFSSLFPLMNCGPSNDIYSIGCVANGFSPSSITFKWTDASEGALTDFVQYFBVQSGANIGVSVVXKNDWEKSKSFRCSVBHQGKTAVIK 99 SAATAFSTLDINMCGPSNDIYSIGCVANGFSPSSITFKWTDASEGALTDFVQYFBVQSGANIGVSVVXKNDWEKSKSFRCSVBHQGKTAVIK 99 ETLTAFVVFKMSQCSS-SIDSUBICIAKGFSPSJTFKWTDASGALTDFVQYFBVQSGANITTKONAUSKSKSFRCSVBHQGKTAVIK 99 VSDSKSFPSILFINMCGFSNDIYSIGCVANGFSPSJTFKWTDASGALTDFVQYFBVQSGANITTKONAUSKSFRCSVBHQGKTAVIK 99 VSDSKSFPSILFINMCGSS-SIDSUBICIAKGFSDSJTFKWTDASGALTDFVQYFBVQSGANITTKONAUSKSFRCSVBHQGKTAVIK 99 VSDSKSFPSIVFKMSQCKS-SSDFJJIGCLASFDSLNIKKK-DNGKLINGIIQYFVKTGDKTFQVSLJANITKONAUSSFRCSVBHQGKT9VF 90 VSDSKSFPSIVFMSQCKS-SSDFJJIGCLASFDSLNIKKK-DNGKLINGIIQYFVKTGDKTFQVSLJANITKONAUSSFNCSFCSVBHQGKF 92 VSQTTAAPALFPIVQCKSGTAGTVTVGCLAQDFFFSSIFFOWTDASGTTQTF-KQYPTVMKDNKYTGVSVLDVSKSAWDSRSFCSVFHQGSFSVTLQ 99
		CH27 ^ ^
T. S. S. D. C. E.	orientalis chuatsi salar mykiss rerio idella coioides	IPSD - FRVTLLSVPNGDTOVLVCTTEEFLPETLS-VKWKKNRDYESDFTDWVPKQ-IGDVYSAVSVLKVKNADWESKAVYTCEVTHGKIYEKKASK-A 188 KPIPPRVTLVSVPSESSQLVCTIEDGRSGTLDSFKWKKNGAELNDVIGSFIGK-TGELESAVSVLKVKNTDMDSKAVYTCEVTYSGTOVRKKASK-A 195 KTVPKSPTVSLLSAPIGTTOVLMCMIEDFNSNTVT-VMWKKNDMEVEGOTPTLVKQ-PSGLVSGSLLKVINTNWNNKVKSCVVCHQEOTINKTISKTE 197 KPVFKSPTVSLLSAPIGTQVLMCMIEDFNSNTVK-VMWKKNDMEVEGOTPTLVKQ-PSGLVSGSLLKVINTNWNNKVKSCVVCHQGTISKTISKTE 198 STAPILSLVIVPTEKNTFAMCVIEDFYSTVK-VMWKKNDMEVEGOTPILGKR-PSGLVSGSLLKVINTDWNNKVKYSCVVCHQGTISKTISKTE 198 KVPFQBPILSLVPVTTQKSTFAMCVIEDFYTENIT-VRWKENNIVKOSOTNLEYKINNAGLHTALSLYKLNEIVIPN-TEYTCEVSHRGKTFEKTONFTA 185 KVPPBP-PENVILVAVPAGDTOTLVCTIEDFYTENIT-VRWKENNIVKOSVGFTDCPPQL-NGGVYTAVSILKVINSEWDSKAVYTCEVTNQCTIYPRKFF KPPPP-PENVILVAVPAGDTOTLVCTIEDFYSNVE-VKWKKDDNSVTGFTDCPPQL-NGGVYTAVSILKVINSEWDSKAVYTCEVTNQGTIYPKKVSK-V 195
		снз→
T. S. O. D. C. E.	orientalis chuatsi salar mykiss rerio idella coioides	PITVTLNPPSPKKFENNNOABLECIVEGODN IVSETEMTWQINGKNVAGNMGLLKTAGSOYSKINTLIRSLTEWLOVNIVRCSAKRKBVTVIKDLTFEKG 289 PITVTLNOPSPKEIFSNKOABLECIITGODTIVDEIKVTWQIDGODVSDNINETKSVDGORIKISTMTRSRTEWORVNKVRCSAIRDDDTLIQDLTVHKG 297 PLTVTLNPPRVREVFLDNQAVLECVITGIDODTVSGTIITWQVNGBKMDGIDLKNIESKGNLNSRVSTLTIGOTEWINVNKVQCSAMKSGEDTPVIQDLSFKKG 302 PLTVTLNPPRVREVFLDNQAVLECVITGIDODTVSGTIITWHINGDIQTAHIDLKNIESKGNLNSRVSTLTIGOTEWINVNKVQCSAMKSGEDTPVIQDLSFKKG 303 KFRULKFPWREMFINNRIVLECVITGIDOTVSGTIFWGVNGBKNDGIDLSVSQENESOHVKINNRVORSAMKRGEDTPVIQDLSFKG 285 TFALTLNPPIERELFVHNKIVLEAVVSGDLSIAVKEAVGASVSCKVKDANVASESITSEIVPSNDTSSFMKKHKVTIDINKWFGEVICTIRDTNNKDIGOKIHEDKG 299 PITVTLFOSSFKEIFSNNGAKFECVITGIDOTGP-DFCIIWQVDGONVTDNHEKISTMTRAHIDWOSINKVRCSAIR-DNMTPVIQELTICK 292
		Сн₄→
T. S. S. D. C. E.	orientalis chuatsi salar mykiss rerio idella coioides	DESKTVTVHILSEEEIRKNSDVTLVCLVSSSVQQTYYIAMSDDAGQNTGNYUDGITFPPQKTQHG-YSVTSVYTINKEKWNQQRSVFNGNVWLVGSNKSMIIRGV 394 DGREFKVTVHVLTEEDINKGAEVTLVCLVSSFVLODYYIAMSBDAGQNTGNYUDGITFPPQKTQHG-YSVTSVYTTNKEKWN-KFNMFYGNVWPAGSNDSMEPRGV S-VAPSVSVHLLPEEDINKEGEVTLVCLVVGPSLCDVYIMWQDJSGQVGEGVTSPPQKTQKANYEVTSVFTTKKKWN-KFNMFYGNVWPAGSNDSMEPRGV S-EAPSVSVHLLPEEDINKGGVTLVCLVVGPSLCDVYIMWKDJSGQVGEGVTSPPQKTVGKANYEVTSVFTTKKKWBRN-UFTCAVKHAGSDNNTAFKERRV 404 S-EAPSVSVHLLPEEDIXGISYVGVSSPSLCDVYIMWKBJSGGVGEGVTSPPQKTVKGNYEVTSVFTITKKKWBRN-VLFTCAVKHAGSDNSTSPKEMSV 96QEPSVEMYKPDDISTSGISYVGVSSPNLCDVYIMWKBJNTTFEGKSSFT02QGGS-TSVGSILTISKKFEWPFTINCAVVHANKDTAFSUKSS 384 DGRKPNVTIYRPDDIKADFVSLVCKVTSSDLCOVYIMWKIGNGPYTEGRTSAPIRONDS-TSILSILTITKKEVKNTTKFSCNVWPAGGNKSMKSRDV 398
		SEC→
T. S. O. D. E.	orientalis chuatsi salar mykiss rerio idella coioides	SKARGNSLECDK 406 SKSEGNSTECR 410 SKSRGNSCEDK- 416 SKSRGNSCEDK- 394 SKSRQAEQPVDY 411 SYAMSNSVECK 410

Fig. 5. Amino acid sequence alignment of the heavy chain of IgT CH domain. The conserved (identical and similar) residues are marked in black. Arrows indicate CH1–CH4 and the secretory tail. An asterisk (*) is above the conserved cysteine that forms a disulfide bond with the light chain, a carrot (^) is above conserved cysteine of intra-domain bond. Gaps are indicated by dashes. Genbank accession numbers are: ACZ54909.1 *Epinephelus coioides* (grouper), ABF19723.1 *Ctenopharyngodon idella* (grass carp), AAY42141.1 *S. chuatsi*, ACX50291. *S. salar*, AAW66981.1 *O. mykiss*, and CAI20890.1 *D. rerio.*

demonstrated heavy chain isotype exclusion at the cellular level infish.

The repertoire data presented here suggest that something 386 different may be occurring in tuna, however (Fig. 8). Like in other 387 388 fish, VH genes appear to be shared between both τ and μ/δ . Three of the four families we found expressed in these fish clearly were 389 390 used in both μ and τ , although a fourth was only found with μ . This 391 could easily be a case of low sampling depth as VH4 appeared as a 392 singular use in the described clones. Since this is a more parsimonious explanation than a dedicated μ VH rearranging to a shared 393 394 DH segment that rearranges to dedicated JH segments, Fig. 8 395 depicts an array of VH gene segments that can be used in either primary transcript type. 396

However, unlike in other fish, both τ and μ rearrangements of 397 tuna appear to employ the same DH gene segment. As all the tuna 398 399 JH genes appear with only μ or τ (none seem to be shared), this points to an arrangement where a single shared DH can rearrange 400 401 with JH segments upstream of either μ or τ to determine isotypic 402 fate of the cell, and this DH's rearrangement to several shared 403 VH's is not the event that stochastically determines isotype. So at 404 least two possibilities of IgT vs. IgM/D lineage fate are now 405 supported by data, one in which τ and μ/δ share VH genes from one block (as in zebrafish) or more than one array 5' and 3' to 406 the τ elements (as in trout) but DH–JH are dedicated to isotype, 407 408 and now the tuna paradigm where VH-DH are shared and JH is 409 dedicated to isotype. In one instance (tuna) the DH-IH join would 410 instruct lineage and in the other the VH-DH join would.

Importantly, we note that genomic sequencing of the locus has 411 not yet confirmed this organization in the tuna or the absence of 412 additional DH that we did not sample. Interestingly, this hypothe-413 sized organization could also explain why in trout a significant 414 difference was seen in CDR3 length and repertoire between τ and 415 μ clones (each using dedicated DH and JH gene segments, (Castro 416 et al., 2013)) while we do not see a great difference in tuna (sharing 417 418 VH and DH and only having dedicated JH, Table 1 and Fig. 3). 419 Future work must determine if this is truly stochastic in lymphocyte development or if there are more complex control mecha-420 nisms instructing this important juncture determining the B 421 cell's fate. 422

4.3. CH regions

IgM is the most conserved isotype in jawed vertebrates and was thought to be omnipresent until the discovery of its absence in coelacanth (Amemiya et al., 2013). The tuna IgM CH region seems very consistent with its orthologs in other fish.

As also noted in other IgT sequences (Hansen et al., 2005a), there are many prolines in the region of the tuna IgT CH1/CH2 juncture which may be indicative of hinge-like flexibility. Tuna IgT CH3 seems to conform to the classical Ig superfamily β -sandwich with canonical cysteines and tryptophan positions seen in the domain of the salmonids and grouper that are important in the folding of this domain (Fig. 5) (Lesk and Chothia, 1982). The tryptophan to cysteine replacement seen in zebrafish and grass

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Fig. 6. Tuna IgH V genes shared with other fish. Phylogenetic analysis of representatives of the four tuna IgH V families (clone 3 for VH1, 5 for VH2, 57 for VH3 and 29 for VH4) with VH genes from two of the better studied teleost models, rainbow trout and zebrafish. Trout and zebrafish accession numbers are labeled at each branch terminus.



Fig. 7. Tuna IgM and IgT CH regions group with those isotypes of other teleosts. Neighbor joining phylogeny using Dayhoff matrix and 1000 bootstrap replications. Alignment and accession numbers used in tree are shown in Figs. 5 and 6.

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Fig. 8. Hypothetical organization of elements in the tuna IgH locus suggests a novel method of lineage determination at the fish IgH locus. Simplified cartoon showing three paradigms in the locus organization and immunogenetic control of IgH τ vs μ/δ rearrangement.

carp appears to be a cyprinid characteristic, and has been suggested by modeling to still allow an immunoglobulin superfamily
domain fold (Danilova and Amemiya, 2009).

In this limited sampling, we found no evidence of the IgT hybrid
molecule with two CH domains identified in the common carp *Cyp*-*rinus carpio* (Savan et al., 2005a), the IgM/D hybrids (with or without VH domains) found in catfish (Edholm et al., 2010), the
IgMCH1/IgTCH4 variant IgT2 in carp (Ryo et al., 2010), nor the
run-on transcription secreted IgD form of trout (Ramirez-Gomez
et al., 2012).

446 In mammalian IgM a carboxyl terminal glycosylation site in the 447 secretory tail is important in J chain polymerization (Hohman 448 et al., 2003; Tacchi et al., 2013), but may have distinct physiology 449 in teleost such as catfish and zebrafish that have it (Wiersma et al., 1997). This conserved N-linked glycosylation site is part of 450 a larger sequence motif enabling polymerization of IgM and IgA 451 of mammals but is not present in the secretory tail of tuna IgM 452 or IgT, although there is a conserved cysteine in IgT shared with 453 other teleosts. Trout IgT was found as a monomer in serum but a 454 multimer in mucus (Zhang et al., 2010), however these IgT multi-455 456 mers did not appear to be covalently linked as they are known to 457 be for trout IgM (Kaattari et al., 1998). More biochemical studies are necessary to resolve the stoichiometry and functional avidity 458 459 of IgT.

460 4.4. Expression

461 Isotype expression studies in tuna echo what has been determined in other fish species: IgT and IgM both are present in pri-462 463 mary and secondary lymphoid tissues, yet more IgM than IgT, however the gap closes at mucosal sites (Hansen et al., 2005a; 464 465 Ryo et al., 2010; Savan et al., 2005b; Xiao et al., 2010). IgT1 in adult 466 zebrafish deviated from this pattern in being primarily in the headkidney and thymus (Hu et al., 2010). The molecular data presented 467 here could serve as a springboard for revisiting immunoglobulin 468 studies in tuna at the protein level that were initiated in the south-469 470 ern bluefin (Thunnus maccovii) (Watts et al., 2001). The work also 471 opens gates to explorations of B lineage development and commit-472 ment, where molecular markers might could be adapted from fish species such as zebrafish (Zimmerman et al., 2011) and trout (Barr et al., 2011; Macmurray et al., 2013) where more work has been performed. 473

5. Conclusions

Endothermic birds and mammals employ immunoglobulin isotypes IgM, IgY, IgE and IgG in systemic immunity but have specialized IgA for mucosal immunity. Poikilothermic vertebrates lack IgA, although amphibians do have an orthologous mucosal isotype in their IgX. IgM had long been the primary functional immunoglobulin isotype recognized in teleost until the recent discovery of the mucosal specialization of IgT. Mucosal epithelia is the barrier breached or exploited by most internal pathogens of vertebrates, and also ectoparasites of fish (Xu et al., 2013). This penetration of mucosal defense is also true of many pathogens of concern in the tuna ranching industry, including sea lice (Hayward et al., 2009), betanodaviruses (Gomez et al., 2010) and gill platyhelminths (Colquitt et al., 2001). It is hoped that this basic molecular characterization of humoral immunity in these economically important endothermic fish will enable more studies of host-pathogen interactions and the feasibility of vaccine development for offshore ranches. Increasing the productivity of these operations by reducing infectious disease mortality will reduce pressures on wild tuna stocks and the fish species used to feed ranched tuna.

Moreover, the apparent shift of isotype determination from VH– DH recombination to DH–JH recombination at the tuna IgH locus is interesting from a fundamental standpoint of lymphocyte antigen receptor immunogenetics, and begs many questions that must be verified and queried with new algorithms (Olivieri et al., 2013) at the levels of the tuna genome, the immunoglobulin proteins, tuna B cells, development in the pronephros, and the fish's response to pathogen. If the single tuna DH gene is verified at the genome, it will be interesting to know whether this IgH locus orientation is found only within this clade of endothermic fish or a broader set of Perciformes. These studies should provide insight into the natural history and fundamental physiology of antibodies while providing much needed tools for managing the health of ranched, and thereby wild, tuna stocks.

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515 Appendix A. Supplementary data

516 Supplementary data associated with this article can be found, in 517 the online version, at http://dx.doi.org/10.1016/j.dci.2013.10.015.

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Supplemental Table 1. Primers.

Primer ID	Primer name	Sequence (5'-3')	Designed to clone	Position	Amino acid
MFC 261	TolgMCF1	TGGATCMGACAGVMWCAGG	#6 KF713344	286-306	NGQNVQP
MFC 244	TolgMCR1	GCACARTAAAHACAGCDCTGTC	#6 KF713344	310-333	DTAVYYCA
MFC 260	TolgMCR	GCACARTAATAHACAGC	#6 KF713344	316-333	AVYYCA
MFC 340	MCF1	CCCTATTTCGCACAGACGCAACAA	#6 KF713344	1279-1302	GIKRVCVV
MFC 350	TolgMGR3F4	CTCATGGTACCCACCAGTGACTTGAGA	#6 KF713344	919- 945	LMVPTSDLR
MFC 351	TolgMGR3F5	ACTGTGACCTACGGAGGGTCATGT	#6 KF713344	1009-1032	TVTYGGSC
MFC 352	TolgMGR3F6	TTCTCTGTAGCTCTGCTGCTGCTGTTG	#6 KF713344	4- 30	FSVALLLLL
MFC 365	TolgZCR1	ACTTGGAGGGTTCAGTGTCACTGT	#59 KF713336	975- 996	TVTLNPPS
MFC 367	TolgZCR2	TGTGTTGACTTGCAGCCACTCAGT	#59 KF713336	1185-1206	TEWLQNQVNT
MFC 369	TolgZCR3	GAATGTGGACTGTCACTGTTGTCTTGCT	#59 KF713336	1278-1302	SKTTVTVHI
MFC 381	TolgMC1R4	TGTAAACTCGGTGGCGAGGCA	#6 KF713344	475- 495	CLATEFT
MFC 382	TolgMC1R5	TGGAGGATACTGGATGAAGTC	#6 KF713344	544-564	DFIQYPP
MFC 384	TolgZRC1R4	ACTGTTGGCTGCCTTGCACGTGAC	#59 KF713336	468- 489	CVGCLARD
MFC 386	TolgZRC1R6	TGGGAGTTTCACTGTCACAGCCATGGT	#59 KF713336	663- 687	TMAVTVKLP
MFC 387	TolgZRC1R7	ACCAAGATGATCGACGGA	#59 KF713336	642-657	SVDHLG

Primer name*	Sequence (5'-3')
βactin	ATCGTGGGGCGCCCCAGGCACC
βactin	GTCATCTTCTCYCTGTTGGC
TNF-α	CCAGGCRGCCATCCATTTAGAAG
TNF-α	CCGACCTCACCGCGCT
IL-1β	GGRSAGCGACATGGYRCGATTTCT
IL-1β	GGTGCTGATGTACCAGTTG

* Mladineo, I., Block, B.A., 2009. Expression of Hsp70, Na+/K+ ATP-ase, HIF-1 alpha, IL-1 beta and TNF-alpha in captive Pacific bluefin tuna (*Thunnus orientalis*) after chronic warm and cold exposure. J Exp Mar Biol Ecol 374, 51-57.

>signal peptide >variable	AC 100
ATGTTCTCTGTAGCTCTGCTGCTGCTGCTGCAGCTGGATCCTGTGTGAAGTGTGAACAGTTGACAGCCAGC	R 34
GTCTGACCATCACCTGCCAGGTCTCTTATTCTGTTGGCAGGTATTACACAGCTTGGATCAGACAGCCTGCAGGGAAAGGACTGGAATGGAATGGAATGG	AG 200
L T I T C Q V S Y S V G R Y Y T A I R Q P A G K G L E W I G M H	R 67
ATATACTGGAGGTTCATACTACAAAGATTCACTAAAGAACAAGTTCAGTATCGACTTAGACTCTTCCAGCAACAGAGTGACTCTAAACGGACAGAATGT	rg 300
Y T G G S Y Y K D S L K N K F S I D L D S S S N R V T L N G Q N V	100
>]olning CAGCCTGAAGACACTGCTGTGTATTACTGTGGCCTCCCGATCAAGTATTTACTATGCTTTTGACTACTGGGGGAAAGGCACGATGGTCACCGTCACCGTC Q P E D T A V Y Y A S R S S I Y Y A F D Y W G K G T M V T V T S >uCH1	AG 400 A 134
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GTCCAAGTGAGGAGACAGGACTGGGAGGGAAAGCAAAGC	AC 700 P 234
CACTTTTTGAGTTGCCAACTCTTAAAGTGCTGGCCTCGCCTCCACGAAGGAGGGCGAGGCTTCCTTC	IA 800 Z 267
TGAGATCAAATGGCTGAAAGATGACGGGGATGTCTTCGACAAAGTATATGAGATCAAAACACCCCATTAAGGAAAGCCAGACCACCGATGGAAAGACACT	'G 900
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TACAGTGTAGCAAGTTTTCTCATGGTACCCACCAGTGACTTGAGACCTAAATTCCACTAAGTTTACATGTGAGATTTAAGTTGAAAAATGAAAACGCATTT Y S V A S F L M V P T S D L R P <u>N S T</u> K F T \square E F K L K N E N A F > μ CH3	v 334
TGAATTCAACTGTGACCTACGGAGGGTCATGTCCTGGAGCCAACTGGATGTGGAAGTACTAGATGTGGAAGTAGAAGACCAAAGGCCCCACAATGACGGACA	AT 1100
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GTTTGTAGAGAGTAAAGGAACTTTAGTATGTCAAGTCAA	FT 1200 400
GAAGAATCAGTCCCCGCTGATGGATTTAAAGGCACAGTCAACGTTCCACTTGACATCACGTATGACGAATGGACTGCTGGGATAAAGCGTGTCTGCGT E E S V P A D G F K G T V N V P L D I T Y D E W T A G I K R V 💆 V >µCH4	rg 1300 v 434
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Q H T N F L E P I K R V Y E R K I V G L E Q H P S V F M L V P V H	S 467
ACAGGCTAATAAAGAAACGGTGACCCTGACTTGCTTTGTGAAAGGCTTCTACCCCAAGGAGGTGTTGGTGTCTTGGCTTGTTGATGATGATGACGCCAGCAGA	AC 1500
Q A N K E T V T L T C F V K G F Y P K E V L V S L V D D V P A D	500
TCAAATTACGATATCAGTACCACAAACCCTGTAGAGAGGAGCAATGGATTCTATTCTGTCTATGGCCAGTTAACACTCAGCCTTGACGATTGGAAGGACAG	rG 1600
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K V Y S G V V Y H E S L K <u>N T T</u> K A I V R S M A H G S T A Q T N I	567
GGTCAACCTCAACATGGAGGTCCAT 1725 VNLNMEVH 575	

Supplemental Figure 1. Nucleic acid and deduced amino acid sequence of *T. orientalis* IgHµ full length clone 6. The start of the predicted signal peptide, VH, JH, CH domains and secretory tail are marked above the sequence. Potential N-linked glycosylation sites are underlined. Cysteines and tryptophans necessary for the Ig superfamily fold are highlighted in black, the cysteine that forms the disulfide bond to the Ig light chain is highlighted in red.

>si ATG	>signal peptide >variable ATGATGGACTATAGGACAGTTGTGCTGCTTTTAACTATCTGCTGGGCAGGCGTTGATGGTCAGACTCTAACAGAATCCGAACCAGCAGTTAAAAAGCCTG														100																						
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Supplemental Figure 2. Nucleic acid and deduced amino acid sequence of *T. orientalis* IgHr full length clone 59. The start of the predicted signal peptide, VH, JH, CH domains and secretory tail are marked above the sequence. Features are annotated as in Supplemental Figure 1.



Supplemental Figure 3. Nucleotide identity matrix of VH region coding sequences to identity VH families. Clone numbers are in bold the left and right of rows and top and bottom of columns. Pairwise differences of less than 30% using nucleotide version of alignment in Figure 3 are highlighted in green and indicate inclusion in same VH family (by 70% nucleotide identity).



Supplemental Figure 4. IgH μ and τ expression in systemic and mucosal lymphoid tissues. Quantitative real-time PCR of secondary lymphoid tissue μ and τ C region mRNA expression relative to that in anterior kidney, standardized to β -actin. Two template concentrations were analyzed. Experiment performed in triplicate, error bars indicate standard deviation.