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Ancient Use of Ig Variable Domains Contributes Significantly to the TCRδ Repertoire

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Abstract

The loci encoding B and T cell Ag receptors are generally distinct in commonly studied mammals, with each receptor's gene segments limited to intralocus, *cis* chromosomal rearrangements. The nurse shark (Ginglymostoma cirratum) represents the oldest vertebrate class, the cartilaginous fish, with adaptive immunity provided via Ig and TCR lineages, and is one species among a growing number of taxa employing Ig-TCR δ rearrangements that blend these distinct lineages. Analysis of the nurse shark Ig-TCR8 repertoire found that these rearrangements possess CDR3 characteristics highly similar to canonical TCR δ rearrangements. Furthermore, the Ig-TCR δ rearrangements are expressed with TCR γ , canonically found in the TCR δ heterodimer. We also quantified BCR and TCR transcripts in the thymus for BCR (IgHV-IgHC), chimeric (IgHV-TCR&C), and canonical $(TCR\delta V-TCR\delta C)$ transcripts, finding equivalent expression levels in both thymus and spleen. We also characterized the nurse shark TCRab locus with a targeted bacterial artifical chromosome sequencing approach and found that the TCR δ locus houses a complex of V segments from multiple lineages. An IgH-like V segment, nestled within the nurse shark TCR8 translocus, grouped with IgHV-like rearrangements we found expressed with TCR δ (but not IgH) rearrangements in our phylogenetic analysis. This distinct lineage of TCRδ-associated IgH-like V segments was termed "TAILVs." Our data illustrate a dynamic TCRδ repertoire employing TCR8Vs, NARTCRVs, bona fide trans-rearrangements from shark IgH clusters, and a novel lineage in the TCR δ -associated Ig-like V segments.

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Disclosures

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The nurse shark, *Ginglymostoma cirratum*, is a representative of the oldest vertebrate class, the chondrichthyans, with an adaptive immune system based on Ig superfamily (IgSF) Ag receptor somatic gene rearrangement in lymphocytes. The Ag receptors of B and T cells have significant similarities through all jawed vertebrate lineages, including the cartilaginous fishes (1-3). Shark BCR, or Ig, and TCR genes employ RAG-mediated V(D)J recombination with segmental junctional diversified by TdT activity. In sharks, both Ig and TCR (4) genes are modified by activation-induced cytidine deaminase-catalyzed somatic hypermutation (SHM) in response to Ag and repertoire generation, respectively (5–11). Shark IgH gene loci exist in many clusters, such as 15 in nurse shark (12) but possibly >100 in other species (13). The segmental ordering of an IgH cluster is V_H-D₁-D₂-J_H followed by a single set of C region exons, which delineate each cluster as either IgM (C μ), IgW [C $_{\omega}$, the ancestor of IgD (14, 15)], or IgNAR (C_{NAR}). This deviates from the typical IgH translocon topography (V_{1-n}-D_{1-n}-J_{1-n}....C_{μ}-C_{δ}-C_{γ}-C_{ϵ}-C_{α}) found in Euteleostomi lineages (16). Although nurse shark IgH rearrangements are generally intracluster, occurring within a single VDDJ cluster, class switch recombination was discovered to alter the C region class of VDJ rearrangements in the nurse shark (17).

The nurse shark has been used as a "primitive" model of the adaptive immune system for decades (18). In this species, polygenic and polymorphic MHC (19, 20); four TCR chains, including a doubly rearranging NARTCR δ (21); and multiple IgH and IgL isotypes have been well characterized (3, 6, 22, 23). Previously, we described unusual cDNA rearrangements of IgHV to TCR δ DJC and extremely rare cases of IgHV to TCR δ D to TCR α JC rearrangements (24). These could be attributed to either trans-rearrangements, occurring across vast genomic distances between distinct TCR and Ig receptor loci, or rearrangements originating from an IgH variable segment or cluster nested in the TCR δ locus.

Rare (1 out of >200,000 PBLs) interchromosomal, interarm, or distal intrachromosomal *trans*-rearrangements occur between different Ag receptor loci in mammals, despite regulatory mechanisms in place to prevent them (25). An inversion of human chromosome 7 can bring TCR β and γ loci, located at distinct telomeres, into proximity, facilitating *trans*-rearrangements (26, 27). Likewise, an inversion on human chromosome 14 juxtaposes the IgH and TCR α δ loci, enabling similar chimeric rearrangements (28). Such *trans*-rearrangements are associated with ataxia telangiectasia and childhood acute B lymphoblastic leukemia (29). The rates of *trans*-rearrangement can increase 50- to 100-fold when double-strand breaks occur during chemotherapy or radiation treatment of lymphoma and leukemia patients (reviewed in Ref. 30).

There is some logic to IgH-TCR δ rearrangements, as both Ag receptor chains have been documented to interact with nonpeptide Ags and display similar CDR3 length distributions (31). The presence of VH δ segments (V segments more similar to IgHV than TCR δ V located in the TCR δ loci of a growing number of vertebrate lineages) furthers the narrative of TCR δ rearrangements tolerating an Ig V domain. In the frog *Xenopus*, "VH δ " genes abound in the TCR α/δ locus and are commonly found in TCR δ transcripts (32). Such VH δ segments have also been detected in the TCR $\alpha\delta$ locus of the platypus, passeriform birds, and the coelacanth (33–35). In galliform birds, VH δ segments are encoded within a distinct

locus with a separate TCRδ C region. The membrane-distal domain of shark NARTCRδ employs a V segment most similar to those of the cartilaginous fish-exclusive IgH class, IgNAR (21, 22). Formulation of a functional NARTCRδ requires two RAG-mediated VDJ recombination events. The membrane-distal V domain is formed by the rearrangement of VDJ segments of the NARTCR lineage, which exist in VDJ clusters (21). This NARTCR domain exon is then spliced into a V domain exon formed by rearrangement between a leaderless, NARTCR-supporting TCR δ V, encoding an additional Cys residue, and canonical TCR8D and TCR8J segments. The rearranged NARTCR-supporting V domain exon is then spliced to the TCR8 C region, yielding a final receptor encoded by NARTCRV-NARTCRD-NARTCRJ-supporting TCR&V-TCR&D-TCR&J-TCR&C (21). Nonplacental mammals (marsupials and monotremes) have a TCR, TCR μ , with a quaternary structure similar to that proposed for NARTCR (36, 37): TCRµ also has two V domains, but both V domains of TCRµ are Ig-like with the membrane-proximal V encoded by a germline-fused VDJ gene. The TCR μ C region is clearly most similar to TCR δ but exists in a locus distinct from the TCRa δ locus. These Ig-TCR δ hybrid receptors corroborate earlier suggestions, based on CDR length, that $\gamma \delta TCR$ binding is structurally more akin to that of Ig receptors than MHCrestricted $\alpha\beta$ TCR (38).

In previous studies, we documented *trans*-rearrangements involving two IgMV and two IgWV rearrangements to TCR δ DJ yielding Ig-TCR δ chimeric receptor rearrangements in the nurse shark. In the current study, we aimed to further characterize the repertoire breadth and prevalence of Ig-TCR δ receptors, in addition to elucidating the genomic organization facilitating Ig-TCR δ rearrangements with a draft of the nurse shark TCR δ locus. To this end, we tested multiple hypotheses to conclude that Ig-TCR δ rearrangements were virtually indistinguishable from their canonical TCR δ counterparts in terms of CDR3 length and diversity and expression in both primary and secondary lymphoid tissue. Additionally, some, but not all, Ig-TCR δ rearrangements originate from within the TCR δ locus in nurse sharks. This study outlines Ig-TCR δ rearrangements that substantially contribute to the shark's TCR δ repertoire diversity, with prevalent expression levels suggesting their importance in an ancient adaptive immune system.

Materials and Methods

Animals

Nurse sharks (*G. cirratum*) were captured off the coast of the Florida Keys and maintained in artificial seawater at ~28°C in large indoor tanks at the Institute of Marine and Environmental Technology in Baltimore, MD, as per animal protocol (University of Maryland Institutional Animal Care and Use Committee no. 1012003, Texas A&M Institutional Animal Care and Use Committee no. 2013–0001). Animals were euthanized and bled out after an overdose of MS222. Tissues were harvested, and cells, DNA, RNA, and frozen histology blocks were prepared, as have been previously described (39–42).

PCR, cloning, and sequencing of IgHV-TCR&DJC transcripts

cDNA primed with reverse TCR δ C (primer FLAJ710, Supplemental Table I) was made with SuperScript III Reverse Transcriptase (Life Technologies, Carlsbad, CA) from 5 μ g of total

RNA from thymus, spleen, and spiral valve (intestine) of a mature (10-y-old) female nurse shark ("Joanie"). These cDNAs were used as a template for standard PCR amplification using GoTaq DNA polymerase (Promega, Madison, WI), with forward primers for IgM (FLAJ1701) or IgW (FLAJ1699 and MFC185) variable genes and reverse primers for the TCRδ C region gene (FLAJ767). After an initial 2-min 95°C denaturation, samples were cycled 35 times through a 30-s 95°C denaturation, 30-s annealing, and 1 min at 72°C, followed by a final 7-min extension at 72° C. Annealing temperatures used were 47° C for FLAJ1701, 51°C for FLAJ1699, and 58°C for MFC185. Bands were purified with the GeneCatcher (Gel Company, San Francisco, CA) isolation system and cloned into pCRII-TOPO (Life Technologies) or pBluescript II KS(+) (Stratagene, La Jolla), using One Shot TOP10 competent cells (Life Technologies). ZR Plasmid Minipreps (Zymo Research, Irvine, CA) of clones of appropriate size were sequenced using BigDye Terminator v1.1 Cycle (Life Technologies) through the Texas A&M University DNA Technologies Core Laboratory. Questionable base calls were corrected based upon other aligned sequences if they clearly occurred within V, J, or constant regions, but the sequence was excluded if ambiguous bases were in N/P regions or the D segment of CDR3. Sequence data were managed with Bioedit (www.mbio.ncsu.edu/BioEdit/bioedit.html) or Geneious Version 9.1 (Biomatters, Auckland, New Zealand) and submitted to GenBank (https://www.ncbi.nlm.nih.gov/popset? DbFrom=nuccore&Cmd=Link&LinkName=nuccore popset&IdsFromResult=374080000, accession numbers JF507709.1–JF507661.1). The CDR3 length was calculated using the "CDR3 length = exclusive number of amino acids from C (of V segment YxC motif) to F (of J segment FGxG motif)" ImMunoGeneTics equation (43). For example, the first clone (T0006M2J09) in Fig. 1A would have a CDR3 length of 21 aa.

Bacterial artificial chromosome sequencing of the nurse shark TCRa8 locus

The G. cirratum bacterial artificial chromosome (BAC) library was probed using cloned segments of TCRaV, TCRaC, TCR8V, TCR8C, and NARTCR probes from splenic transcripts amplified with ³²P, as described under high-stringency conditions (42, 44). Selected BAC clones, positive for TCRa, TCR\delta, and/or NARTCR components, were isolated from 500-ml cultures using the Qiagen Large Construct Kit (Qiagen, Hilden, Germany). Purified BAC DNA was sent to the Duke Center for Genomic and Computational Biology for PacBio RSII large insert (15–20 kb) library sequencing. Assemblies were performed with the PacBio Corrected Reads pipeline, Celera Assembler version 8.3, using both raw PacBio reads, corrected in the PacBio Corrected Reads pipeline, and PacBio circular consensus sequences with minimum a read length of 500 bp were used as input files for the assembly (45, 46). Options to merge haplotypes to a single consensus were used to account for the BAC library containing both parental chromosomes. Reads were deposited into the Sequence Read Archive under Bioproject PRJNA530194. Annotation was performed using the Geneious software suite version 9.1. We employed a combination of BLAST, using a custom database of expressed nurse shark Ig and TCR sequences, and manual searches of assembled contigs for recombination signal sequences unveiled V, D, and J segments not yet in public databases.

Quantitative real-time PCR

Quantitative PCR (qPCR) was performed with 50 ng of random hexamer–primed cDNA generated with SuperScript III from thymic and spleen RNA samples of sharks ranging in age from 9 mo to 10 y. We used the SYBR Green PCR Master Mix (Bio-Rad, Hercules, CA), following the manufacturer's recommendations. Triplicate wells were assayed in a Bio-Rad CFX96 thermocycler for 40 cycles, annealing at 55°C. For the absolute quantification of each sample, a standard curve was created using serial dilutions of certain concentrations of a single-copy gene cloned into a vector. Additional interexonic, real-time PCR primers can also be found in Supplemental Table I. The resulting quantities, given in copy number per 50 ng of cDNA, were split into three groups (canonical BCR, canonical TCR, and chimeric Ig-TCR) for statistical analyses. Significance was determined via the median-based Kruskal-Wallis test, with post hoc Dunn test to determine specific differences between the groups.

Generation of anti-IgHV polyclonal antisera

Polyclonal nurse shark IgMV1, IgWV1, and IgWV2 group (47) Abs were generated in rabbits by Cocalico Biologicals (Reamstown, PA) by immunizing them with IgHV-maltose binding protein (MBP) fusion proteins. The IgHV sequences were amplified from shark (Joanie) spleen cDNA with 35 cycles of PCR, annealing at 59°C (IgM) or 63°C (IgW). Restriction endonuclease site-engineered primers used were MFC180 and MFC181 for IgMVI, MFC182 and MFC183 for IgWVI, and MFC184 and MFC183 for IgWVII (Supplemental Table I). These products were cloned into the pMAL-c2x (New England Biolabs, Beverly, MA) expression vector using SHuffle Express competent cells (New England Biolabs). Recombinant protein was produced in bacteria and cleared supernatant passed through amylose resin columns twice. Fusion proteins were eluted with maltose and precipitated with saturated ammonium sulfate, resuspended in PBS, and dialyzed in Slide-A-Lyzer cassettes (Pierce, Rockford, IL). Sizes were verified by 12% SDS-PAGE using Coomassie and silver staining. The immune serum of recombinant protein-immunized rabbits was passed through an MBP affinity column to remove MBP-specific Abs. Affinity purification of Abs to shark IgHV in the cleared sera was performed with the same immunizing Ag immobilized in an agarose bead column (AminoLink, Pierce) and verified by SDS-PAGE and Western blotting.

Flow cytometry

Thymocytes and splenocytes $(5 \times 10^5$ cells per treatment) were stained with either biotinylated mouse mAb LK14 (48) (against *G. cirratum* IgL) or unlabeled anti-IgHV rabbit polyclonal (against *G. cirratum* IgMV1, IgWV1, or IgWV2) at 1:100 in staining buffer (1% BSA in shark PBS) for 1 h at 4°C. Cells were then washed three times with staining buffer before staining with streptavidin-allophycocyanin (eBioscience, San Diego, CA) at 1:1500 and anti-rabbit Alexa Fluor 488 at 1:500 (Southern Biotech, Birmingham, AL), respectively, for 30 min at 4°C. All samples were washed and resuspended in 300 µl of staining buffer containing 0.1% sodium azide and examined by flow cytometry on a BD LSR II instrument (BD Biosciences, San Jose, CA). Fifty thousand events were collected, gated for live cells, and analyzed using the FlowJo software (Tree Star, Ashland, OR). Identical signal

thresholds could be applied to all samples except the 120-mo shark, which had lower fluorescent intensity across all experiments.

Phylogenetic analyses

V segment alignments for both cartilaginous fish and vertebrate lineage trees were performed in Geneious using ClustalW. Amino acid alignments containing the entire V segment, from FR1 and to the conserved Cys of FR3, were used in the cartilaginous fish alignments. The multispecies alignments used nucleotide sequences of only the framework regions as CDR length and composition vary greatly across multiple vertebrate lineages (49, 50). The resulting nucleotide alignment was manually adjusted to fit the b strand ImMunoGeneTics protein display for V domains, as previously described (51).

Maximum likelihood trees were constructed for all alignments in MEGAX using 1000 bootstrap replicates (52). For the cartilaginous fish phylogeny, we used a Poisson correction model with bootstrap values displayed for bifurcations with >50% consensus tree support. The multispecies tree used a general time reversal model, and the substitution rate was γ distributed with invariant sites with six discrete γ categories.

Stellaris RNA FISH

Frozen tissue sections were prepared from OCT embedded thymus tissue of a 10-y-old nurse shark, as previously described (24). Three tissue sections of the thymus, cut to a thickness of 10 μ m, were probed using the Stellaris (LGC Biosearch Technologies, Middlesex, U.K.) RNA FISH system. Custom probe sets were designed using the Stellaris custom probe designer for the IgWV1 segment, TCR δ C region, and TCR γ C region. The fluorophores used for each probe set were CAL Fluor Red 610 (IgWV1), Quasar 670 (TCR δ C), and CAL Fluor Orange 560 (TCR γ C). Nuclei were stained with DAPI (Sigma-Aldrich, St. Louis, MO). The probed slides were imaged on a Zeiss Stallion Digital Imaging Workstation, as previously described (4). Images were processed using ImageJ software version1.7 (53).

Results

Repertoire sequencing reveals additional IgHV usage

Transcripts in which an IgHV segment is rearranged to TCR δ gene segments have been previously documented in the nurse shark (24). To determine the breadth of this chimeric receptor repertoire, 5' RACE PCR was performed using a TCR δ C region primer. Nested forward primers were designed to selectively amplify the Ig-TCR δ hybrid receptors (Supplemental Table I). The sequenced amplicons totaled 137 unique IgV-TCR δ DJ rearrangements (chimeric, or *trans*-rearranged, clones are abbreviated Ig-TCR δ) from spiral valve (n = 38), spleen (n = 51), and thymus (n = 50) transcripts, displaying a diverse repertoire of Ig-TCR δ in both primary and secondary lymphoid tissues (Fig. 1A–C, Supplemental Fig. 1). Most of the recovered transcripts (95.6%) were in frame, indicative of selection mechanisms prohibiting nonfunctional rearrangements or mutations. We identified nine IgM/W V segments rearranged with TCR δ ; all but one segment, S0021TVJ09, could be found in BCR rearrangements bearing an IgM (or IgW) C region. This sequence encoded an IgH-like V segment and was most similar (68% nucleotide identity) to the nurse shark IgH

group 7 pseudogene cluster (EU312153). This genomic V segment was not found expressed in any rearrangement and was not determined to be the source of this peculiar V segment used in transcript S0021TVJ09. Accession numbers for IgH V segments found in both IgH and TCR& rearrangements were AY609260 (IgMV1), DQ857389 (IgMV2.1), AY609247.1 (IgMV2.2), AY609259.1 (IgMV4), AY609249.1 (IgMV5), KF192877.1 (IgWV1), KF192883.1 (IgWV2), and KF192884.1 (IgWV3).

Nurse shark Ig-TCR₈ and canonical TCR₈ repertoires possess equivalent CDR3 metrics

We compared the Ig-TCR8 CDR3 transcript metrics to that of the canonical TCR8 repertoire to ascertain whether chimeric Ig-TCR δ rearrangements bore similar CDR3 metrics to their canonical counterparts (Fig. 2, Table I). Two aberrant transcripts with extraordinarily long CDR3 lengths (S2223W1J08 and V0924W1J20 with CDR3 lengths of 41 and 49 aa) were removed from statistical analyses. The distribution of CDR3 length for functional TCR8 (n= 55) and Ig-TCR8 (n = 135) was nearly identical and clearly distinguishable from TCRB (n= 41) and IgH (n = 173) (Fig. 2A). Mean CDR3 length of Ig-TCR8 rearrangements was 17.44 ± 0.42 aa, statistically indistinguishable from that of canonical TCR8 (17.40 ± 0.67 aa), with each TCR δ subset encoding repertoire CDR3 content significantly longer than that of both TCR β (11.68 ± 1.01 aa) and IgH (12.55 ± 0.25) determined via ANOVA (Fig. 2B, Table I). The ranges of CDR3 lengths of receptors free from MHC restriction (TCR $\gamma\delta$ and Ig and especially IgHV and TCR δ) are typically much longer than that of TCR $\alpha\beta$ (38). Indeed, the range of the Ig-TCR δ subset was comparable to that of canonical TCR δ receptors (Fig. 2C, Table I), with both TCR δ subsets encompassing a larger range than either TCR β or IgH (Fig. 2C). Finally, we analyzed the isoelectric point (iP) of CDR3 residues and found that TCR δ (iP = 8.35) and Ig-TCR δ (iP = 8.38) both averaged slightly higher iP than either IgH (iP = 6.86) or TCR β (iP = 5.98) (Fig. 2D). These data show that chimeric Ig-TCR δ and canonical TCR δ receptors are comparable not only in CDR3 length but possess similar charge signatures integral to epitope-paratope interactions as well. These results were unsurprising as both Ig-TCR δ and canonical TCR δ receptors draw from the same TCR D and J segment pool, contributing heavily to the CDR3 similarities outlined above (Fig. 1, Supplemental Fig. 1). Additionally, we elucidated that the IgWV1 gene segment, found expressed in Ig-TCR δ rearrangements, was found in individual cells also expressing TCR γ using RNA FISH (Supplemental Fig.2). The highly similar CDR3 metrics between Ig-TCR8 and canonical TCR δ is indicative of an IgHV segment, with divergent CDR1 and CDR2 loops, which use provides substantial expansion of potential TCR8 paratopes.

Absolute quantification of TCR₈ by real-time PCR shows equivalent levels for canonical and chimeric transcripts

Although the number and diversity of chimeric clones suggested that nurse shark Ig-TCR δ rearrangements were of significance for the shark immune system, we performed absolute qPCR to determine their prevalence in comparison with other TCR δ rearranged transcripts. Specific primers for each V segment (TCR δ , IgM, and IgW) were used with either a TCR δ or IgH C region primer (Supplemental Table I). Interestingly, no significant differences were found between expression levels of canonical TCR δ and chimeric Ig-TCR δ transcripts in the thymus (Fig. 3A). Although expression of BCR was decidedly lower than either TCR group in the thymus, we found no significant differences between the three groups in the thymus of

young sharks aged 10 and 16 mo (Fig. 3A). Expression of transmembrane IgH message in the thymus of developing sharks has been documented, but consistent with our new quantitative data, such expression diminishes to background levels as the animals mature (6, 23). Expression of BCR in the spleen compared with thymus was markedly higher than both TCR δ groups, yet there remained no significant difference between canonical TCR δ and chimeric Ig-TCR δ expression in the periphery (Fig. 3B). Furthermore, flow cytometry performed on thymocytes, using anti-IgHV polyclonal and the LK14 mAb specific for κ L chains, provided preliminary evidence that chimeric receptors are present as a surface receptor (Supplemental Fig. 3). Although IgW has been shown to prefer an IgL other than κ (54), qPCR confirmed that expression of either IgWV segment was negligible in the thymus of these sharks (Fig. 3A). Importantly, these data show Ig-TCR δ and canonical TCR δ mRNA expression to be equivalent in both primary and secondary lymphoid tissues and suggest that they are used as a surface Ag receptor.

Draft assembly of the G. cirratum TCR⁶ locus reveals a novel V segment lineage

Because IgH V segments have been found to be associated with TCR δ loci in a growing number of vertebrate lineages, we also performed a draft assembly of the nurse shark TCR8 locus with targeted BAC sequencing. To this end, the G. cirratum BAC library was probed, and selected BAC clones, positive for TCR\delta, TCRa, and NARTCR components, were selected for long-read sequencing. The resulting assembly yielded a draft TCR δ translocus assembly totaling 169 kb with an average coverage depth of 61, with TCRa J segments lying downstream. Although the TCRa C region was not assembled within the contig housing the TCR δ translocus (Fig. 4, contig 10), the presence of TCRa J segments downstream of the TCR δ translocus indicate a linked TCR $\alpha\delta$ locus in this ancient model of IgSF receptor loci. This was corroborated by the BAC library screening, which identified multiple BACs probing positive for TCR δ , TCR α , and NARTCR gene segments (data not shown) and the linkage of TCRa and NARTCR in contigs that could not be linked to the grander TCR8 translocus (Fig. 4, contigs 36, 65, and 112). Although the targeted BAC sequencing approach allows confident assembly of IgSF loci, the technique is hindered in instances in which probed BACs contain no overlapping regions. Furthermore, without a nurse shark karyotype to probe chromosomes, selection of positive BACs with overlap is a random method that does not account for vast genomic regions without TCR gene segments.

The draft assembly of the nurse shark included four contigs with a total of 29 functional V segments, 4 D segments, 18 J segments, and 1 TCR δ C region. The largest contig (contig 10) revealed a translocon including 24 V segments (19 functional and five pseudo-V segments), a single TCR δ D segment, 11 TCR δ J segments, 4 TCR α J segments, and the TCR δ C region (Fig. 4A). We also assembled fragments with segments shown to be involved in formulating the TCR δ repertoire in three additional, nonoverlap-ping BACs. These BACs included three NARTCR VDJ blocks, with two functional NARTCRVs (contigs 36 and 112) and one pseudo-NARTCRV (contig 65). Each NARTCR block was found upstream of a supporting TCR δ V segment (functional on contigs 36 and 112, pseudogenized on contig 65). Two additional TCR δ V segments were identified upstream of the NARTCR block in contig 112. The first TCR δ V on contig 112 was truncated by the assembly, and the 164 bp assembled contained no frameshifts or stop codons; however, the second TCR δ V is

functional. Presumptively, these indicate an additional TCR&V translocon stretch unidentified thus far. Finally, we identified 6 TCRa V segments all oriented in reverse orientation in relation to the NARTCR blocks. Three functional TCRaVs were located downstream of the NARTCR block on contig 65. The remaining three were located on contig 36. One functional TCRaV was found upstream of the NARTCR block, whereas one functional TCRaV and a single pseudo-TCRaV were located downstream.

Additionally, we identified one functional IgHV-like segment and one pseudo-IgHV-like segment nested in the translocon stretch of TCR δ Vs (Fig. 4A). Although the IgHV-like segment is putatively functional, it could not be confidently ascribed to any Ig-TCR\delta rearrangement in our expression dataset; however, it was similar to the novel IgHV-like segment we found only in TCR δ rearrangements (Fig. 4B). In phylogenetic analyses of cartilaginous fish IgH and TCR8 V segments, these Ig-like V segments (not found in rearrangements bearing IgH C regions) interestingly grouped together in a clade nestled between the IgM and IgW segments (Fig. 5A). We dubbed the segments belonging to this clade TCRδ-associated Ig-like V segments (TAILVs) to distinguish them from the V segments found on both TCR δ and IgH receptors. We also performed a phylogenetic analysis that included TCRa, TCR\delta, VH\delta, IgH, and IgL genomic V segments from all vertebrate lineages. The nurse TAILV representative sequence was found in a branch that included nurse and whale shark IgMVs, recapitulating the findings in our cartilaginous fish tree (Fig. 5B, 5C). This multispecies phylogeny also unveiled a clade of V segments, previously labeled TCR\deltaV, sharing a common ancestry with Ig V segments rather than TCRV. The lack of a clade solely composed of Ig V segments found in TCR8 loci indicates that they have arisen multiple times through the course of vertebrate evolution.

Comparative analysis of cartilaginous fish TCR8 loci

A number of cartilaginous fish, including the whale shark, Rhincodon typus (55), bamboo shark, Chiloscyllium plagiosum, and cloudy cat shark, Scyliorhinus torazame (56), have joined the first cartilaginous fish genome of the elephant shark, Callorhinchus milii (57). Of the recently sequenced genomes, only the bamboo shark has a scaffold (BEZZ01002038.1) in which multiple TCR\deltaVs could be located; however, no TCRS J segments or C region was found. This left only the elephant shark assembly with a TCRS on the 694-kb scaffold NW_006890273 that is flanked by two IgMV segments found in typical cluster arrangement: one cluster possessing a full complement of IgM CH exons, the other with only a single CH exon. Also present in the elephant shark genome are NARTCRV and supporting TCR8V blocks, allowing syntenic comparisons to those found on smaller contigs in the nurse shark (Fig. 6A) (58). The absence of a TCR δ J segment and C region in either assembly made it impossible to confirm whether the TCR δ C proximal V translocon was inverted in other cartilaginous fish species, as well. Nonetheless, the elephant shark scaffold was still useful to formulate a predictive placement for drafted nurse shark NARTCR contigs, allowing us to posit the topography of a complete nurse shark TCR δ locus (Fig. 6A). Furthermore, the proximity of the IgM clusters to the TCR δ locus in the elephant shark allowed us to formulate a potential model for the locale of (at least some) IgH clusters used in Ig-TCR[®] rearrangements (Fig. 6B). Finally, we formulated a model of all nurse shark Ig-TCRδ rearrangements elucidated thus far: one intralocus TAILV and bona fide translocus

rearrangements between IgWV, from three clusters, and IgMV, from five clusters, which contribute to the TCR δ repertoire (Fig. 6B).

Discussion

From what is known of cartilaginous fish IgH isotypes and the Ag recognition modes of vertebrate $\gamma\delta$ T cells, there is a clear logic in IgMV and IgWV rearranging to TCR δ . Both share the same intrinsic structure of the IgSF V domain and the capacity to interact with free Ag. Drawing from the same pool of D and J segments as canonical TCR δ rearrangements, the Ig-TCR δ repertoire was found to be just as diverse as their canonical counterparts, characterized by large nontemplate TdT additions at the V-D and D-J junctions (Fig. 1, Supplemental Fig. 1). The large nontemplate additions found in Ig-TCRδ rearrangements were constrained to encode CDR3 lengths consistent with that of canonical TCR8 rearrangements in the nurse shark (Fig. 2, Table I). We were surprised to find that the predicted CDR3 iP in the two TCR8 rearrangement classes were also roughly equivalent, despite the bulk of CDR3 residues stemming from nontemplate nucleotides (Fig. 2). Similarities observed in CDR3s of Ig-TCR δ and canonical TCR δ suggest the two groups make use of the same dimeric partner chain. The heterodimeric TCR complex, complete with signal transduction machinery, is required of all T cells during thymic selection (59-61). We have determined that Ig-TCR δ rearrangements also pair with TCR γ ; this is likely due to conservation of the Cys residue in TCR8 C regions, facilitating the interchain disulfide bond linking $\gamma\delta$ TCR complexes (Supplemental Fig. 2)(62). Any Ig-TCR δ chain rearrangements bearing incompatibilities with a partner chain, which prohibited paratope formation, would be selectively eliminated from the repertoire. Differences in the IgV domain are seemingly well tolerated by the partner chain, as the mRNA expression levels of Ig-TCRδ and canonical TCRδ were indistinguishable (Fig. 3). Thymic expression of aberrant receptors could be attributed to remnants of nonfunctional rearrangements (Fig. 3A). Equivalent expression levels for Ig-TCR\delta and canonical TCR\delta in the spleen of all five sharks (Fig. 3B) indicate that the Ig-TCRδ rearrangements regularly pass selection. Furthermore, the presence of Ig-TCR8 receptor rearrangements for at least three IgHV segments was identified on sorted thymocytes even in sharks that lacked support for BCR expression in the thymus (Supplemental Fig. 3).

Altogether, Ig-TCR δ rearrangements in the nurse shark are neither aberrant nor rare. With CDR3 metrics and expression levels so closely aligned to canonical TCR δ , it appears that Ig-TCR δ rearrangements, via their Ig-derived CDR1 and CDR2, are well tolerated by the partner chain and would diversify the recognition potential of shark TCR δ receptors. Such diversity would likely be disallowed in MHC-restricted $\alpha\beta$ T cells but is tolerated in $\gamma\delta$ T cells unbound from these restrictions.

Recombination signal sequences are the regulatory elements governing rearrangements in sharks as well as in mammals (62–65). If an IgHV were in proximity to the TCR δ locus, one could expect Ig-TCR δ rearrangements to occur between IgH and TCR δ loci. Indeed, this was the case, as several V segments identified in our Ig-TCR δ dataset originate from canonical IgH clusters. We found that bona fide *trans*-rearrangements between IgH and TCR δ loci are regular occurrences in the nurse shark. However, these were not the only type

of chimeric Ig-TCR8 rearrangements we documented, as several V segments were not found in any IgH cluster nor expressed in the nurse shark Ab repertoire. Our draft nurse shark TCR δ locus assembly identified a single IgHV-like segment integrated in a translocon stretch of TCR\deltaVs, with no downstream Ig C region, that likewise was not found expressed as an IgH (Fig. 4). Phylogenetic analysis of shark TCRS and IgH V segments determined that these peculiar segments both clustered in a distinct clade in the phylogenetic analysis (Fig. 5). As both are clearly more Ig-like than TCR, we decided to call these V segments TAILVs. Presence of TAILVs in the TCR δ locus in one of the earliest iterations of IgSFbased adaptive immunity adds to the narrative that Ig and TCR δ V segments are largely interchangeable. This is also consistent with the emerging paradigm of IgHV segments associated with TCR8 loci of a growing number of species. Indeed, we identified a clade of canonically labeled TCR8Vs (located in the TCR8 locus of tetrapod, avian, and mammalian lineages) with a shared Ig ancestry (Fig. 5B, 5C). This evidence highlights implementation of Ig-like V segments on TCR8 receptors in all jawed vertebrate lineages from cartilaginous fish to eutherian mammals. Phylogenetic evidence did not support TAILV as the progenitor of VH δ nor the aforementioned Ig-like TCR δ V clade (Fig. 5C) but illustrates how IgH and TCR δ have freely exchanged gene segments over the course of vertebrate evolution. In fact, the location of the holocephalan (the most ancient chondrichthyes lineage) elephant shark IgMVs, which flank a TCR8 locus with no evidence of TAILVs, in the cartilaginous fish phylogeny suggest these proximal IgH clusters could be the ancient precursors to TAILVs (Fig. 6). Confirmation of this hypothesis via identification of syntenic blocks will require a more-complete elephant shark assembly and completion of the nurse shark TCR8 locus. Although we cannot claim that these IgMVs gave rise to the nurse shark TAILVs, we outlined hypothetical origins of TAILV as well as potential sources of IgHV segments found in both IgH clusters and Ig-TCR δ rearrangements in Fig. 6. This hypothetical syntemy also provided evidence that the NARTCR blocks are likely localized to the TCR8 locus in all chondrichthyan species with confirmed NARTCR expression.

The TCR α and TCR γ loci of sharks have been shown to employ SHM (4, 66, 67), a process normally associated with IgH/L loci. Although IgHV sequences in our dataset contained minor differences within identical CDR3 regions, we found no conclusive evidence (via mutations in the IgHV of transcripts with identical CDR3 content) suggesting that SHM acts to expand diversity within the IgHV segments used in chimeric rearrangements. This was surprising, as one would assume the Ig V segments would be primed for diversifying mutations, suggesting that the anatomic and developmental windows of activation-induced cytidine deaminase expression are tightly regulated in the shark as well as endothermic vertebrates.

In summary, although the evolutionary history of Ig-TCR δ remains complex in vertebrates, we have confirmed that Ig-TCR δ rearrangements contribute to the peripheral T cell repertoire. The cellular contribution of Ig-TCR δ within the nurse shark immune system was highlighted in sequence diversity and expression in an assortment of tissues. Indeed, at all ages included in this study, Ig-TCR δ receptors are as prevalent as canonical TCR δ rearrangements. Additionally, the Ig-TCR δ repertoire makes use of Ig V segments both from IgH clusters and from a separate lineage of IgHV-like segments, termed TAILVs because of their exclusive association with TCR δ (Fig. 6B). Comparative TCR δ locus analysis of

cartilaginous fish, both holocephali and elasmobranchii that split ~420 million years ago (68), exemplifies the versatility of TCR δ : a TCR employing receptors bearing the V domain of canonical TCR δ , IgHV (through chimeric Ig-TCR δ rearrangements originating both within the TCR δ locus [TAILV] and from IgH clusters of unknown proximity), and the doubly rearranging NARTCR (Fig. 6B).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Abbreviations used in this article:

BAC	bacterial artificial chromosome
IgSF	Ig superfamily
iP	isoelectric point
MBP	maltose binding protein
qPCR	quantitative PCR
SHM	somatic hypermutation
TAILV	TCR8-associated Ig-like V segment

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FIGURE 1.

Diverse, in-frame, IgHV-TCRDJC rearrangements are expressed in thymus and periphery of nurse shark. Amino acid translation showing CDR3 regions amplified from IgHV forward and TCR δ reverse primers from (**A**) thymus, (**B**) spleen, and (**C**) spiral valve. Clone name is to the left of alignment, followed by the IgH V segment origin, with the TCR δ J used to the right of the alignment. Sequences are aligned to the conserved cysteine of the YxC motif of the V domain and the FGxG motif of the J. Nucleotides within CDR3 are aligned in the center, and conserved residues of the genomic D segment are highlighted by frame (first frame, cyan; second frame, green; third frame, yellow) and flanked by nontemplate and palindromic residues. Asterisks (*) denote stop codons, and slashes (/) denote frameshifts.



FIGURE 2.

Chimeric Ig-TCR δ and canonical TCR δ encode comparable CDR3 metrics. Analysis of CDR3 for rearranged transcripts of TCR δ V-TCR δ C (red), IgHV-TCR δ C (green), IgHV-IgHC (blue), and TCR β V-TCR β C (gray) showcases the near-identical distribution (**A**), average length (**B**), range (**C**), and iP (**D**) of chimeric Ig-TCR δ and canonical TCR δ . Significance of CDR3 length indicated by lettering (B) was determined via ANOVA with post hoc Tukey honestly significant difference test, with each TCR δ subset being significantly longer than IgH and TCR 2β .



FIGURE 3.

Absolute qPCR reveals chimeric Ig-TCR δ expression to be comparable to canonical TCR δ in both primary (thymus) and secondary (spleen) lymphoid tissue. Absolute transcript values (displayed on the y-axis, calculated using known concentrations of an amplicon cloned into plasmid) are compared within each shark (displayed on the *x*-axis indicated by age in months) for (**A**) thymus and (**B**) spleen samples. To compare, rearrangements were classified as canonical B cell (BCR, blue), chimeric (Ig-TCR, green), and canonical TCR δ transcripts (TCR δ , red). The boxplot extends from the first to the third quartiles (25th and 75th percentiles, respectively), with centerline indicating median value. The whisker of each boxplot extends to the value no. >1.5 times the interquartile range (difference between the first and third quartiles), values beyond these whiskers are potential outliers, indicating a significant skew or preference for a given rearrangement. Because of the relatively high proportion of outliers present, the nonparametric Kruskall-Wallace test with post hoc Dunn

test was used to determine significant differences in groups (BCR, chimeric, TCR) within a given shark. Significance is indicated by asterisks (*). **p < 0.01, *p < 0.05, n.s. (p > 0.05).

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FIGURE 4.

Assembly and mapping of TCR δ loci reveals IgH-like V segment bearing similarity to those found rearranged with TCR δ but not IgH. (**A**) Segments located within the assembled nurse shark TCR δ locus are color coded according to lineage-differentiating TCR δ (red), TCR α (blue), and IgH/TAILV segments (purple). Pseudogene segments are indicated by lighter shading. The contigs in which the segments are housed are boxed and labeled beneath the map. The prospective location of the TCR α C region is also indicated by the light blue box downstream of the TCR α Js on contig 10. (**B** and **C**) Alignment of V segments identified in assemblies, including the TAILV and IgHVs found in both TCR δ and IgH rearrangements (**B**), as well as NARTCRV (C), and NARTCR supporting TCRV (**D**).

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Elephant TCR_δ

lumanTCRδV2 Armadillo TCRδV

- PigTCR_δV

Manatee TCRoV3-1p

Alpaca TCRδV2

Tenrec TCRδV Opossum TCRδV2

Opossum TCRδV3

Mouse TCR_δV4

Frog TCRδ1.1

Nurse Shark IgNARV

-Frog VHδ1.2 LFrog VHo1.4

-Frog VHδ1.5

Frog VHo1.6

Froa laHV4S1

Frog VHδ4

Manatee VHop

. Platypus VHδ

Coelacanth IgW2V5 Coelacanth IgWV1 Coelacanth IgW2V7 Coelacanth IgWV8

0.20

99- Frog IgHV5S1 Frog VHδ5.1

Coelacanth loW2V6 Nurse Shark IgWI-TCRV

-Coelacanth IgW2V1 IqHV (Mammals, Platypus, Chicken)

Frog laHV9S1

- Frog VH_{01.1} - Frog VHo1.3

Armadillo VHδp

Chicken VHo Zebra Finch VHδ

Opossum VHδp

Zebra Finch TCR₀V1

Zebra Finch TCR 8V3 IgHV (Mammals, Platypus, Frog)



FIGURE 5.

0.20

Phylogenetic analysis of Ag receptor V segments unveils a novel lineage of TCRSassociated IgH-like V segments (TAILV) in cartilaginous fish. (A) An amino acid alignment of chondrichthyes V segments, trimmed to the start of FR1 and ended at the conserved Cys of FR3, was used to construct a maximum likelihood tree. Species included in the alignment were Ginglymostoma cirratum (nurse), Scyliorhinus canicula (cat), Heterodontus francisci (horn), Carcharhinus plumbeus (sand), and Callorhinchus milii (elephant). The percentage of trees supporting branches after 1000 Bootstrap replications is displayed for bifurcations with >50%. The tree is drawn to scale, with branch lengths measured in the number of substitutions per site. The shaded regions indicate different V segment lineages found in Chondrichthyes, including elasmobranch TAILVs (green), elasmobranch IgMV (red), holocephali IgMV (pink), elasmobranch IgW (teal), and elasmobranch TCR δ V (blue). (**B**) Topology of the jawed vertebrate Ag receptor tree. V segment lineage indicated by color for IgL (gray), Ig-like TCR&V (green), IgNARV/NARTCRV (orange), IgH (blue, includes TAILV and VH δ), and TCR $\alpha\delta V$ (red). The tree includes representative sequences from

TCRαδ

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CD4

human, mouse, pig, tenrec, elephant, alpaca, manatee, opossum, platypus, zebra finch, chicken, frog, coelacanth, nurse shark, and elephant shark. (C) Subtree of the IgH and Iglike V segments. Green shading indicates lineages of Ig-like V segments that are exclusively found on TCR rearrangements, highlighting multiple exchanges of Ig V segments into TCR loci throughout vertebrate evolution.

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FIGURE 6.

Comparative synteny between holocephali and elasmobranchii TCR δ loci and model of nurse shark noncanonical TCR δ rearrangement origins. (**A**) Sequence comparison of the TCR δ locus structure using elephant shark scaffold NW_006890273.1 as a map for assembled nurse shark contigs. Light blue shading indicates syntenic nurse shark contigs from the draft assembly, with associated contigs labeled beneath the box. Green shading indicates prospective positions of nurse shark IgH V segments used in Ig-TCR rearrangements based on position within the elephant shark scaffold. (**B**) Model of the multiple sources of Ig-TCR δ rearrangements, including TAILV rearrangements from within the TCR δ locus, bona fide *trans*-rearrangements between the eight IgH clusters whose location in relation to the TCR δ locus is unknown, and the doubly rearranging NARTCR δ .

Table I.

CDR3 lengths of Ig-TCR8 chimeric receptor chains

Median 17 17 18 15 Mean 17.4 18.0 17.8 16.3 Mean 17.4 18.0 17.8 16.3 Variance 22.8 26.4 23.6 19.6 Maximum 34 34 28 29 Minimum 7 8 7 11	edian 17 17 ean 17.4 18.0	18 17.8		Calibilical I CNO
Mean 17.4 18.0 17.8 16.3 Variance 22.8 26.4 23.6 19.6 Maximum 34 34 28 29 Minimum 7 8 7 11	(ean 17.4 18.0 minnes 22.8 26.4	17.8	15	18
Variance 22.8 26.4 23.6 19.6 Maximum 34 34 28 29 Minimum 7 8 7 11	76 J		16.3	17.4
Maximum 34 28 29 Minimum 7 8 7 11	alallee 22.0 20.4	23.6	19.6	25.0
Minimum 7 8 7 11	aximum 34 34	28	29	30
	iinimum 7 8	7	11	7
Range 27 26 21 18	ange 27 26	21	18	23