

# Article

# Origin of Immunoglobulin Isotype Switching

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# Summary

**Background:** From humans to frogs, immunoglobulin class switching introduces different effector functions to antibodies through an intrachromosomal DNA recombination process at the heavy-chain locus. Although there are two conventional antibody classes (IgM, IgW) in sharks, their heavy chains are encoded by 20 to >100 miniloci. These representatives of the earliest jawed vertebrates possess a primordial immunoglobulin gene organization where each gene cluster is autonomous and contains a few rearranging gene segments (*VH-D1-D2-JH*) with one constant region,  $\mu$  or  $\omega$ .

**Results:** V(D)J rearrangement always takes place within the  $\mu$  cluster, but here we show that the VDJ can be expressed with constant regions from different clusters, although *IgH* genes are spatially distant, at >120 kb. Moreover, reciprocal exchanges take place between *Ig* $\omega$  and *Ig* $\mu$  genes. Switching is augmented with deliberate immunization and is concomitant with somatic hypermutation activity. Because switching occurs independently of the partners' linkage position, some events involve transchromosomal recombination. The switch sites consist of direct joins between two genes in the 3' intron flanking *JH*.

**Conclusions:** Our data are consistent with a mechanism of cutting or joining of distal DNA lesions initiated by activation-induced cytidine deaminase (AID), in the absence of mammalian-type switch regions. We suggest that, in shark, with its many autonomous *IgH* targeted by programmed DNA breakage, factors predisposing broken DNA ends to translocate configured the earliest version of class switch recombination.

# Introduction

In mammals, a primary antibody response begins with a rising IgM titer that is followed, with a lag, by antibodies of a different class, such as IgG or IgA. The "switched" antibodies preserve the antigen-combining sites previously associated with IgM heavy (H) chain but replace the C-terminal portions that bear other functions such as the recruitment of effector immune cells via binding to Fc receptors, to bring about antigen clear-ance [1]. In disease states such as hyper-IgM syndrome where there is a deficiency in secreted Ig classes other than IgM, recurrent infections demonstrate the survival value of the

absent isotypes. Antibody isotype switching relies on a molecular process called class switch recombination (CSR), where deletional recombination juxtaposes the VDJ combining site to downstream C exon sets. In activated B lymphocytes, activation-induced cytidine deaminase (AID) initiates DNA lesions in the VDJ, promoting somatic hypermutation (SHM), and in the highly repetitive switch (S) regions 5' of the C exons, generating DNA double-stranded breaks (DSB) [2, 3]. S regions are critical to CSR, because their sequence and structure enhance targeting by AID, rendering the area recombinogenic [4]. How the DSB are achieved is not clear, but the ends are repaired and become recombined through nonhomologous end-joining (NHEJ) pathways [5, 6].

Ig classes exist in all vertebrates, but unambiguous parallels to the mammalian IgM-IgG switch extend only to amphibians [7, 8]. The representatives of the earliest jawed vertebrates, cartilaginous fishes like sharks and skates, are the oldest group to possess an adaptive immune system based on V(D)J recombination. They express two conventional Igs, IgM and IgW, and a third that is a single-domain binder, called IgNAR [9]. The IgM/IgW H chains are encoded by 20 to >100 miniloci or "clusters," a unique type of organization considered ancestral to the classical Ig locus in higher vertebrates (Figure 1) [10]. After defining the germline Igµ genes in the nurse shark, we were able to demonstrate that despite the multiple autonomous IgH, H-chain exclusion exists and shark antibody heterogeneity was mainly generated by junctional diversity and SHM [11-15]. Here we report the unanticipated finding that the multiple cluster organization also supports Ig isotype switching. Thus the basic features of humoral immunity-including V(D)J rearrangement, H-chain exclusion, SHM, and now CSR-emerged in the ancestor of all jawed vertebrates.

# Results

# Overview

The experimental results are presented as follows: (1) Screening of complementary DNA (cDNA) libraries revealed Ig transcripts composed of the VDJ belonging to one *IgH* gene cluster and the C region to another. (2) Parallel library screening and RT-PCR experiments show that the proportion of switched Ig is highest in immunized adults, less in nonimmunized individuals, not detectable in neonates. (3) Every *IgH* gene studied can switch. Switching to G5 C region and reciprocal switching of G5 VDJ to other C regions were observed. (4) The nature of mutations in productive VDJ of switched Ig suggests that the polypeptides were expressed and under selection. (5) Using cDNA primed in the J-C intron, sequences containing switch junctions were isolated. These are transcripts of *IgH* genes that appear to have undergone recombination.

# cDNA Sequences Not Correlating with Germline Organization

Characterization of nurse shark  $Ig\mu$  genes from bacteriophage and bacterial artificial chromosome (BAC) libraries respectively representing 4.5 and 11 genomes' coverage showed

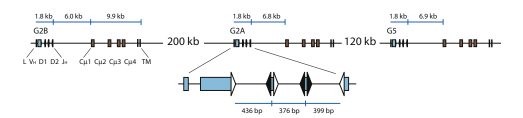


Figure 1. Organization of Three Linked IgM H Chain Genes in Nurse Shark

The *IgH* clusters were mapped in [13] and distances are indicated. Each cluster consists of a split leader (L) and the rearranging gene segments (VH, D1, D2, JH) depicted as blue boxes (enlarged) flanked by recombination signal sequences (RSS) shown as triangles. The RSS with 23 bp spacers are white, and those with 12 bp spacers are black. Brown boxes represent the four C region exons (Cµ1 Cµ4) and the transmembrane (TM) exons.

that each cluster consists of a single *VH*, two *D*, one *JH*, and one set of C $\mu$  exons (Figure 1) [12, 13]. G1, G2A, G2B, G3, and G5 are single-copy genes present in all sharks, and their *VH*, *JH*, and C $\mu$  exons are unique (Figure 2A). The 6–10 kb J-C intron was sequenced in each gene (accession numbers JQ272838–JQ272843), but the highly repetitive S region in tetrapods was not observed.

In the course of screening the shark 33 epigonal organ (bone marrow equivalent) cDNA library, we isolated a few  $\mu$  chain sequences whose components did not correlate with the established germline organization [11] in the same individual. One cDNA consisted of VDJ from the lg $\mu$  gene G1 but with the C region from G2A (Figure 2B). Three other cDNAs contained VDJ from G2B but again C region from G2A (data not shown). The VDJ were intralocus rearrangements; chimerism began only 3' of JH.

To ascertain the frequency of chimeric cDNA, we looked for switching in all five subfamilies. Three more combinations were found: G3 VDJ to the G4C C region, G4A VDJ, and G4CG VDJ to G4E C region (Table 1, shark 33). The C regions were replaced in their entirety. Of six new chimeric sequences obtained from shark 33, one in particular was informative. Clone E176 carried the VDJ of G2B and the C region of G2A, and this VDJ was shared with one of the 11 conventional G2B sequences, E3-1 (Figure 2C). The common CDR3 and mutations shared by these two cDNAs established that both rearrangement and somatic hypermutation (SHM) had occurred in a parental clone previous to the event that produced E176. These cDNA sequences have all the hallmarks of products of Ig H-chain class switch recombination in tetrapods.

# Switch Frequency Is Higher in Immunized Sharks

Shark 33 was not deliberately immunized; therefore, its mutant Ig sequences were produced during natural exposure to environmental antigens. Two cDNA libraries were constructed using mRNA from immunized sharks—JS and GR (Table 1). Whereas in shark 33 1/12  $\mu$  cDNA bearing G2B VDJ have switched, in shark-JS it was 5/11 and in shark GR 20/23. For sequences bearing the C region of the G2 subfamily, 1/43 (2.3%) were switched in shark 33, compared to 5/41 (12%) in shark JS and 25/108 (23%) in shark GR. In view of these differences, we focused on G2 switches.

G2 sequences were amplified by RT-PCR from six animals, including immunized (sharks JS, GR, PI) and nonimmunized individuals (sharks AQ, 33, 626). The reverse primer targeted the G2A and G2B C region, but the universal forward primer allowed amplification of any VH subfamily (G1–G5). Most of the VDJ switching to G2A C region would be G2B, as observed in the cDNA libraries (Table 1). The two isotypes are

differentiated by two BstEII sites found in G2A but not in G2B C region (sites shown in Figure 2C). VH-hybridizing fragments with a single BstEII (marking the G2A C region) are  $\sim$  647 bp (Figure 3A); the absence of BstEII in JH means either a mutated G2A JH or a switched G2B VDJ. Individuals display varying amounts of the  $\sim$  647 bp fragment.

Neonatal shark AQ and three other pups show little mutated or switched G2 sequences (Figure 3A, lanes 1 and 2; see also Figure S1A available online). Sharks JS, GR, and 33 on the other hand serve as positive controls. The good correlation of the 777/647 bp ratio with G2B/switched-G2A frequencies observed in three cDNA libraries (Figure 3B) demonstrates how the RT-PCR products accurately reflect the switched Ig content. A marked difference in the 647 bp band can be observed between the immunized and nonimmunized samples.

In another experiment, the G2A and the G2B C regioncarrying populations were separately amplified (Figure S1B). Switched sequences bearing the G2A C region were observed in the same individuals and in the relative amounts as in Figure 3A, whereas in the parallel G2B samples, there was little reciprocal switching detectable. Because G2B is upstream of G2A (Figure 1), we investigated whether the biased switched combinations are a result of the relative physical location of the genes.

# Switching Is Not Restricted by Gene Position

Only switching events taking place between G5 and the other  $Ig\mu$  can be scored with confidence by PCR methods because in these cases subfamily-specific markers distinguishing JH from CH are separated by 2 bp (Figure 2A). The location of G5 with respect to two upstream  $Ig\mu$  has been determined (Figure 1), so we looked for all possible switches to the G5 C region as well as reciprocal events involving G5 VDJ.

G5 C region-containing Ig were amplified from sharks AQ, JS, GR, and PI and selected for switched sequences by differential digestion (Figures S1C–S1E). None was obtained from neonate shark AQ. In contrast, the switched G5 C region-containing sequences cloned from shark JS included G2A, G2B, G4A, and G4CG VDJ and those from shark GR G4A and G4CG VDJ (series I–V, Figure S2).

Direct amplification was also performed using subfamilyspecific primers, for example, in the G4 leader (G4L) and in the G5 C region (G5CH1R, Figure S2 diagram). Diverse sequences were successfully amplified, with several clones sharing the same VDJ as clones obtained in the first, bulk approach (paired, Figures S2I and S2II), showing that the same switched population was detected in both experiments. Virtually every *IgH* within 340 kb (G2A, G2B) and elsewhere (G1, G4A, G4CG) was found switched to the G5 C region.

# A JH-Cµ linkage

#### JH C region BstE II GCTATCTTGATCACTGGGGACAAGGGACCATGGTGACCGTGACTGCAG G1 G2A Α....Τ.....Τ.....Τ.....Τ.....ΤΤ... .....C.G.....C..G.CG......T..C......G.C..AT G2B .....C.G.....C..G.CG......T..A.....G.....G.....G....AT A....T...G.T.....TT.. .....C.G.....C..G.C.....T..A......CGG......G...T G3 G4A A....CT......A.........A........T......TT... .....C.G.....C..G.C......T..C......CG......G...T A....CT......A.......A.........T......TT... G4E $\ldots \ldots C.G.\ldots .C..G.C\ldots \ldots T\ldots C\ldots \ldots .CG\ldots \ldots .G...T$ G4C/G .....C.G......C..G.C......T..C......CGT......G...T CA...C.G.....G.C....T.A......GG....TT G5

# B G1 VDJ with G2A C region

	Leader
G1 GL	AGAAC <mark>CGTTCTCCACAAGT</mark> TACCAGAAAGA <mark>TT ATGTC</mark> GACTATTTTT <mark>T</mark> TCAGTTTGTTGCTGACTTTCTTATC <mark>A</mark> TGTGAGTGTTTATTCTTTAAGTCTGTCTTTGTTTTCGTCTCAGTCTCACCTCAGCAATGTAAAAGACTCAGCCAT
CH22	AGAAC <mark>CGTTCTCCACAAGT</mark> TACCAGAA <mark>A</mark> GA <mark>TT ATGTC</mark> GACTATTTTT <mark>T</mark> TCAGTTTGTTGCTGACTTATTGTACAAT
G2A GL	AGAACATTCTCTCCCAGGATACCAGA ATGACGACGATGATAATTTTTCTCAGTTTGTTACTGAATTTCTTATCCTGTGAGTCTTTATTCTTTAGTTTTGTTATTTCCCCAGTGTTAGTCTCCCTCTGGAATGTAAAATATTCAGCAAT
G1 GL CH22	CATTGCTTCAGCGTAATGCTCATGTTTTATTTTTCATTTTCAAGGTGTCCA <mark>A</mark> TCG <mark>CAAA</mark> TTA <mark>T</mark> TTTGA <mark>C</mark> TCAA <u>AAAGT</u> GGCCAGAAACTGGCCGGGCCTGGAAGACCGGCCCGACGCCCGAAGACCGGCCCGGAGGTCCGAAAAAGCGGGCGTCCAAGACCAGCGGGCCCGGAGGCCCGGAGGCGCCCGAAGACCAGCGGGCCCGGAGGCGCCGGGCCGGGCCGGGCCGGGCCGGGCCGGGCCGGGCCGGGCCGGGCCGGGCCGGGCCGGGCCGGGCCGGGCCGGGCCGGGCCGGGCCGGGCCGGGCGGCCGGGCCGGGCGGCGGCGGCGGGCCGGGCGGGCCGGGCGGGCCGGGCGGGCCGGGCGGGCGGGCGGGCGGGCGGGG
G2A GL	
G1 GL	TAGALTAGE AND A CONTRACTOR AND A CONTRACTOR AND A CONTRACT AND A CONT
CH22	TGGATGCAG TGGATCCGACAAGTTCCAGGACAGGGCCTGGAGTGGCTGGC
G2A GL	<u>GCCATGAGT</u> TGGGTCCGACAGGTTCCCGGACAGGGGCTGGAGTGGATAGTT <u>TACTACTACTAGGGTCATATAGCAATGACCTATGCGCCAG<mark>CGA</mark>TTAAAGATCGCGTCCATAGACACTTGCAGAGTACTTCCCGGTCGAAATGAAGAG</u>
	D1 D2 JH
G1 GL	CCTGAAGATCGAAGACACCGCCATCTATTACTGTGCAAAAAGC AGTGGGT ATACACTGGATTGG GCTATCTTGAT <mark>C</mark> ACTGGGGACAAGGGACCATGGTGACCGTGACT <mark>GC</mark> AG
CH22	CCTGAAGATCGA <mark>g</mark> GACACCGCCAT <mark>E</mark> TATTACTGTGC GCG AGTG ATCC CACAT <mark>C</mark> ACTGGGGACAAGGGACCATGGTGACCGTGACT <mark>GC</mark> AG
G2A GL	CCTGAAGATCGAAGACACCGCCATCTATTACTGTGCAAGAGAC ATACTACAGTGGGT ACATACTGGGATAG ACTATTTTGATTACTGGGACAAGGGACCATGGTGACCGTGACTTTAG
	CH1
G1 C	
CH22	I GACARCETTE TE CECERTE ET TATORE ET GETE TE CETTE TE CARGARAGE ANALACIÓN TO TATORE ET TO TATORE TATORE AT TO TATORE TATORE AT TO TATORE AT
G2A C	TGACAL CETCTCC/CLALG/CTTTATCG/CTC/GCTCTCCTCCTCTCACACACACACAA GACAA GCGTTCTGTGATTTTGGTTG/TTGGCGATGACAACACCACLAG GTGACCTCGCAAAACGTGGACAACACCACCAGA
GEA C	
	VDJ CH1 CH2 CH3 CH4
	E3-1 E3-1
_	C28 VD1 with C24 C region F176
С	G2B VDJ with G2A C region
	Leader
G2B GL	AGAAC <mark>C</mark> TTCTCTCCCCAGGATACCAGA <b>ATG</b> ACGATGATGATGATGATGATGATGATGATGATGATGAGATTTCTTATCCTGTGAGTCTTTATTCTTTAGTTTTTGTTATTTCCCTAGTGTTAGTCTACCTCTGGAATGTAAAATATTCAGCAAT
E3-1	AGAAC <mark>C</mark> TTCTCTCCCAGGATACCAGA <b>ATG</b> ACG <mark>A</mark> TGATGATGATTTTTCTCAGTTTGTTACTGA <mark>C</mark> TTTCTTATCCT
E176	AGAAC <mark>C</mark> TTCTCTCCCAGGATACCAGA <b>ATG</b> AC <mark>T</mark> ACGATGATC <mark>A</mark> TTTTTCTCAGTTTGTTACTGA <mark>C</mark> TTTCTTATCCT
G2A GL	ΑGAACATTCTCTCCCAGGATACCAGA <b>ΑΤG</b> ACGACGATGATAATTTTTCTCAGTTTGTACTGAATTTCTTATCCTGTGAGTCTTTATTCTTTAGGTTTCTATTTGTTATTCCCCGAGTGTTAGTCTCCCCCTCTGGAATGTAAAATATTCAGCAAT
G2B GL	VH CDR1 CAGATAATCAGTAAAAATTGTCATGTTTTTGCTTTATTTCTGAAGGTGTCCAGTCGGAGGGAG
E3-1	
E176	
G2A GL	
02/1 02	CDR2
G2B GL	ACCATGAGTTGGGTCCGACAGGTTCCCGGACAGGGGGGGG
E3-1	ACCATGAGTTGGGTCCGACAGGTTCCCGG <mark>C</mark> CAGGG <mark>C</mark> CTGGAGTGGATAGTT <u>TACTtgcGTGCATGGCATGGCAGGCAGGCAGGCAGGGCAGGGCAGG</u>
E176	ACCATGAGTTGGGTCCGACAGGTTCCCGG <mark>c</mark> CAGGG <mark>c</mark> CTGGAGTGGATA <mark>cTcTACTtgcATGC</mark> CTCgagTAG <mark>t</mark> AATGACTATGCGCCAGCGATTAAAGATCGATTTACTGCGTCCAAAGACACTTCAAACAATATCTTCGCGTTGGAAATGAAGAG
G2A GL	GCCATGAGTTGGGTCCGACAGGTTCCCGGACAGGGGCTGGAGTGGATAGTT <u>TACTACTATGGTTCATATGGCAATGACTATGGCGCCAGCGATTAAAGAT</u> CGATTTACTGCGTCCATAGACACTTCAAACAATATCTTCGCGTTCGAAATGAAAGAG
	D1 D2 JH
G2B GL	CCTGAAGATCGAAGACACCGCCATCTATTACTGTGCAAGAGAC ATACTACAGTGGGT ACATACTGGGATAG ACTATTTTGGTTACTGGGGAACAAGGGACCATGGTGAC <mark>A</mark> GTGACTTTAG
E3-1	CCTGAAGATCGA <mark>g</mark> GACACCGCCATCTATTACTGTGCAAGA TTCGAGGGAGGCGG CAGTGGGT ACG ACTGGG GGGCCTACTGGGGACAAGGGACCATGGTGAC <mark>A</mark> GTGACTTTAG
E176	CCTGAAGATCGA <mark>G</mark> GACACCGCCATCTATTACTGTGCAAGA TTCGAGGGAGGCGG CAGTGGGT ACG ACTGGG GGGCCTACTGGGGACAAGGGACCATGGTGAC <mark>A</mark> GTGACTTTAG
G2A GL	CCTGAAGATCGAĀGACACCGCCATCTATTACTGTGCAAGAGAC ATACTACAGTGGGT ACATACTGGGATAG ACTATTTTGATTACTGGGGACAAGGGACCATGGTGACCTTTAG
	†BstEll (G2A)
	CH1 CH2
G2B C	TGACACCGTCTTCCCCGACGCTTTATGGTCTAGTCTCGCCTGTCAGCAACAGAACAATGAAGGGAACTTTCACGCTGAGCAGCCAGTTAGCC/ GTCGAATCGATGGCCCTCCAACTGTTCTCCTA 0Bst EII
E3-1 C	TGACACCGTCTTCCCCGACGCTTTATGGTCTAGTCTCCCCCCTGTCAGCAACAGAACAATGAAGGGAACTTTCACGCTGAGCAGCCAGTTAGCC/ GTCGAATGGACCCTCCAACTGTTCTCCTA 0Bst Ell
E176 C	TGACACCGTCTTCCCCGACGCTTTATGGTCTCGTCTCCTTCAGCAACAGACCAATGACGGGAACTTACACCCTGAGCAGCCAGTTAGCC/ GTGGAAACGGTGACCCTCCCGACTGTTCTCCTA 1 Bst Ell site in Cµ2
G2A C	TGACACCGTCTTCCCCGACGCTTTATGGTCTCGTCTCCTTCAGCAACAGACAG
	†BstEll (G2A)

#### Figure 2. cDNA Sequences Compared to Germline Components

(A) Germline JH gene segments and linked C $\mu$ 1 exons of the Ig $\mu$  genes G1 G5, as defined in genomic bacteriophage and BAC libraries [11 13]. BstE II site (shaded) in JH of G1, G2A, and G5. Dots indicate identity with G1 reference sequence.

(B) G1 VDJ with G2A C region. Comparison of germline VH gene segments and partial  $C\mu$ 1 exon from shark 33 G1 and G2A *IgH* with shark 33 cDNA CH22; drawn as the top, bottom, and center sequences, respectively. Nucleotide identities restricted between CH22 and one germline (GL) sequence are high lighted in yellow for VH and in pink for  $C\mu$ . Mutations are marked by lower case and highlighted in blue. The GL gene segments consist of leader, VH gene segment, two D genes, and JH gene segment, as labeled; GL intersegmental sequences are not shown. ATG is bolded, the CDR in the V gene segment are underlined and labeled. C is constant region, and dashes indicate gapping. Accession number is JQ272824.

(C) G2B VDJ with G2A C region. Diagram is color coded to depict the relationship of the two cDNA sequences. Comparison of shark 33 cDNA clones E3 1 (nonswitched G2B sequence) and E176 (switched) that share CDR3. See legend to (B). The number and location of BstE II sites in the GL G2A and E176 are indicated at lower right as well as by arrows. Accession numbers are JQ272833 and JQ272828.

We then looked for and found reciprocal combinations for G5 VDJ (Figure S2, series VIII–XI). For genes of established order (Figure 1), reciprocal switching of the G2B or G2A VDJ to the G5 C region and the G5 VDJ to G2B or the G2A C region demonstrates that switching combinations are independent of the genes' relative positions on the chromosome. In particular,

switching of VDJ to upstream C regions show that these events may involve the homologous chromosome.

# Related Clones Suggest that Switching Is Not Transient

The frequent finding of related clones (Figure 2C; Figure S3; Table 1 footnotes c and d) demonstrates that not only did

Table 1. IgM H Chain Switch among cDNA Clones from Three Shark cDNA	
Libraries	

 $^{a}\mbox{Accession}$  numbers for shark 33 and JS sequences are: JQ272822 JQ272837.

<sup>b</sup>The VH gene segments of germline G4C and G4G are indistinguishable; it is the C regions that differ.

<sup>c</sup>Following clones indicated as "n = n" are identical, whereas as "n/n" share CDR3 but differ by mutation: GR 9 = 305, 12, 23, 47, 63, 99, 102, 157, 208, 51/154/214, 158/164/304/215/218, 36/235 (12 CDR3 among 19 unique sequences).

<sup>d</sup>Following clones indicated as "n = n" are identical, whereas "n/n" share CDR3 but differ by mutation: IgW VDJ to IgM G2A, w 2/3w = 4w/47w = 22w = 17w = 40w; 1w = 14w/7w/15w/52w/65w/6w/20w; 246w (3 CDR3 among 16 sequences, 11 unique). IgW VDJ to IgM G4C, 58w = 61w. Some clones shown in Figure S5 (accession numbers JQ272792 JQ272796). <sup>e</sup>Clone GR c57 (accession number JN802117), shown in Figure S6.

they originate from activated B cells but also that their switched configuration, even with a nonfunctional VDJ, is a stable feature in dividing cells undergoing SHM.

Because many cloned combinations in shark GR involved G4 VDJ and G5 C, we sought nonfunctional VDJ in the reciprocal combination (Figure S2, sets X, XI). In two such related clones, the mutations consist of in/del and alter important structural residues (Figure S3A). In the absence of shark B cell lines, we cannot ascertain whether they arose from a recombination process involving exchange for a productive VDJ. The deleterious changes contrast with the conservative nature of the FR changes in in-frame VDJ shown in Figure S3B and Figure 2, suggesting that in the latter, translated products were under selection.

# Switch Junctions

The reciprocal switches occurring between G2B, G2A, and G5 suggest that the process could involve transchromosomal

recombination and/or gene conversion. We attempted to isolate sequences containing a switchpoint located downstream of JH; the combination of the G4 VDJ switched to the G5 C was selected for the disparity in their J-C intronic sequences and because many G4 to G5 switched sequences were isolated from shark GR (Figure S2, I and II).

We had previously found extensive mutation and evidence of DNA breaks within the 500 bp 3' JH from genomic VDJ [16], an area prone to SHM. The fruitful cloning efforts were directed to this region in cDNA, with first-strand primers targeting points in the G5 J-C intron (Figure 4, top). Sequences isolated in these experiments are shown in Figure 4 and Figure S4. The switch junctions of all cloned sequences are shown in detail in Figure S4B-S4P, with arrows marking the 5'-most site where cleavage could have occurred on the donor sequence. Subsequent experiments involving cDNA primers located further downstream revealed switch junctions at primarily similar locations (Figure S4DII). Most of the switch joints are accompanied by insertions of unknown origin (Figure 4; Figures S4F-S4I and S4K-S4O) that suggest modification of DNA ends during repair processes such as NHEJ [5, 6]. This area did not contain non-B-DNA structure or greater levels of RGYW motifs.

What is more, we have discovered that some of these VDJ-switched intron sequences are clonally related to VDJ switched to C region from the G4G5 series (Figure S2I). This is schematically depicted in Figure 5. The isolation of four such related pairs demonstrated that some switched VDJ did derive the novel C region through recombination in the J-C intron (Figure 5; Figure S5). Thus, the isolation of related sequences, using successively (1) primers in VH FR1 and CH1 and (2) primers in the leader and J-C intron, demonstrates that the intronic junctions are in vivo gene products. Most importantly, these pairs show that the connection between VDJ and switched C region physically existed prior to VDJ to CH splicing. As such, we concluded that the switched VDJ and C were recombined at the DNA level.

At this time, we cannot conclude whether all or even the majority of switched *IgH* are formed by intra- or interchromosomal recombination in the J-C intron, but of 17 unique G4A VDJ switched to the G5 C region, four were related to the VDJ isolated with switch junctions in the intron (Figure S2I, asterisked). This proportion indicates that switching events apparently involving DNA DSB and repair by NHEJ are not infrequent. There is no definitive evidence for gene conversion for switch; any such event must extend from the J-C intron through the C exons (>7 kb). In the cDNA libraries, none of the sequences were chimeric within the C region or beginning within VH.

# Activation-Induced Cytidine Deaminase

Results presented in Figure 3A and Figure S1A show that little Ig switching occurs in four shark pups. This was confirmed in parallel experiments where we tried to clone intronic switch junctions among pup shark AQ Ig transcripts. In contrast to older sharks, AID is barely detectable in neonatal tissues like the epigonal organ and spleen (C. Zhang and E.H., unpublished data), correlating with their low levels of SHM [11, 13, 17].

The unique feature of shark SHM is that half of substitutions are adjacent nucleotide changes with a biased occurrence at RGYW/WGCY hotspots [11, 18, 19]. SHM extends into the JH 3' intronic flank for 1–1.5 kb, with a high frequency of duplications and deletions, sometimes accompanied by

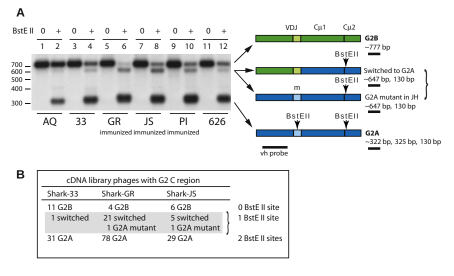


Figure 3. Frequency of Switch to G2A C Region Varies among Individuals

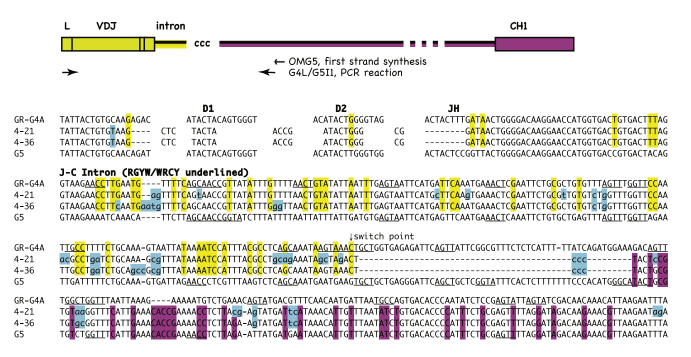
(A) Left shows RT PCR samples from nonimmu nized sharks AQ (lane 1, 2), 626 (11, 12), and 33 (3, 4) and from immunized sharks GR (lane 5, 6), JS (lane 7, 8), and PI (lane 9, 10). First strand cDNA was primed with a universal CH2 primer (CH2 3') and PCR was performed with G2 specific reverse primer (G2CH2R) but universal forward primers (V1 V5) for 30 cycles. The PCR products were incubated with (+) and without (0) BstEll and electrophoresed. The filter was hybridized with a vh probe (black bar in diagram). Right shows PCR products depicted with their BstEll sites and fragment sizes after BstEll incu bation. The underlined fragment is detected by vh probe. Nonswitched G2B (green) has no BstEll sites whereas nonswitched G2A (blue) contains one in JH segment and one in  $C\mu 2$ (arrows). Digestion with BstEll generates bands of  $\sim$  322 bp (G2A),  $\sim$  777 bp (G2B), and  $\sim$  647 bp (switched or mutated BstEII) as detected by vh probe in the samples.

(B) Tally of G2 sequences from cDNA library clones in Table 1. The relative ratios of G2B to switched Ig/G2A mutant (shark 33, 11:1; shark GR, 4:22; shark JS, 6:6), respectively, correlate with the 777 bp/647 bp signal intensity detected by phosphorimaging of (A) lanes (lane 4, 3719/396; lane 6, 497/2514; lane 8, 2567/3004).

nontemplated additions [16]. Although the current work was done on cDNA, all mutational changes other than the switching process are similar in character to previous observations made in shark B cell genomic DNA. Moreover the nature of the switch junctions resembles DNA DSB and repair processes found in the intron of nonswitched Ig.

# Switching to and from IgW

The epigonal organ in shark is analogous to bone marrow also in the sense that plasma cells tend to home there. After screening the shark GR library with  $\mu$  probe, we found not only more G2B VDJ switched to the G2A C region than nonswitched G2B (Table 1), but we also detected IgW VDJ



#### Figure 4. Two Clones Share CDR3 and Switch Junctions

Top diagram shows sequences generated by RT PCR using primers in the J C intron of G5 and leader of G4. Yellow indicates sequence identity with G4, pink with G5; "ccc" insertion at putative switch point. Not drawn to scale, as indicated by break symbol.

Bottom shows that G4 VDJ joined to G5 intron are compared to G4A and G5 reference sequences. Related clones 4 21 and 4 36 share CDR3 and mutations (blue) through VDJ and JH flank. The flanking intron is labeled. Hotspot motifs RGYW/WRCY are underlined in the reference sequences. Arrow (switch point) indicates where the G4A sequence identity ceases. This is followed nontemplated addition (ccc) and identity with G5 (in pink). Accession numbers are JQ272798 and JQ272799.

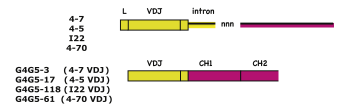


Figure 5. Clonally Related VDJ Are Flanked with Switched Intron or Spliced to Switched C Region

Top diagram depicts RT PCR product generated using primers in the G5 J C intron and the leader of G4, as in Figure 4.

Bottom diagram depicts RT PCR product generated using primers in the G5 C region and universal primers in VH, as in Figure S2. Yellow highlight indi cates G4 sequence, pink G5. Clones 4 7 and G4G5 3 share VDJ and muta tions; 4 7 sequence was switched to G5 in the J C intron, whereas G4G5 3 VDJ was spliced to G5 C region. Three other such pairs were found (G4G5 17 and 4 5, G4G5 118 and I22, G4G5 61 and 4 70). Sequence comparisons are shown in Figure S5. Full alignments with G5 intron are shown in Fig ure S4: (D) (4 5), (E) (4 7), (F) (4 70), (O) (I22). Accession numbers are JQ272797, JQ272804, GQ359826, and GQ359827.

switched to the G2A C region (Figure S6A). We then used IgW VH and CH probes to assess the extent of switching with Igµ genes, isolating not only additional IgW VDJ switched to Cµ sequences but also one cDNA that was IgM VDJ switched to the Ig $\omega$  C region (Figure S6B). As listed in Table 1 (footnotes c and d), the switched clones in the epigonal organ are highly amplified.

# Discussion

We have identified a somatic recombination process that occurs between IgH clusters in shark B lymphocytes, whose transcribed products are primarily productive VDJ switched to a novel C region. Our data are consistent with a SHM-mediated switching mechanism whose differences from tetrapod CSR may reflect an early process that existed in the ancestral vertebrate. The process is concomitant with AID/SHM activity: hardly detectable in neonates but vigorous after deliberate immunization in older animals. Furthermore, SHM occurs throughout the switching process because it is present both before and after the switching event. Not only is switching between IgM and IgW bidirectional, but every gene inspected can undergo switching, regardless of the relative chromosomal location of the partner. The discussion focuses on those differences from the well-studied mammalian systems in order to extract the elemental features of CSR as it evolved in the ancestral vertebrate Ig gene system.

# **Relationship of SHM and Isotype Switching**

The S regions in mammals contain long stretches of repetitive motifs that are 60% GC-rich and specialized features that promote switching through R-loop formation [20, 21]. Although the amphibian *Xenopus* S regions have normal

(40%) GC content and do not form R-loops, they contain palindromes and RGYW/WRCY motifs, and the region enables switch recombination when it replaces the mouse counterpart [22-24]. However, the shark J-C intron does not possess such structures. The location and nature of the shark switch junctions show that the DNA breaks leading to isotype switching arise from AID-mediated lesions that occur in proximity to the VH promoter. We have found no evidence for other promoters in the J-C intron. Both SHM and switching originate from AID lesions, and it may be that intrinsic properties of the J-C intron sequence render it a recombination hotspot. As such, in the earliest version of Ig switching, SHM and CSR are contiguous. We speculate that it was only after amphibian divergence, with the introduction of S regions, that SHM and CSR became independent, spatially and temporally, as they are in mammals [25-27].

CSR in mammals usually takes place intrachromosomally but can also occur between alleles. In rabbits, transrecombination occurs with a frequency of 3%-8% [28] and interallelic CSR in mice engineered to monitor this phenomenon takes place at similar and higher frequencies [29]. A rate-determining synapsis step for two IgH may render shark CSR less frequent, just as transrecombination constitutes a fraction of total isotype switching in rabbits. There is very little information on chromosomal distribution of IgH genes in cartilaginous fishes. In one study, the many  $Ig\omega$  and  $Ig\mu$  genes in clearnose skate appear to be in multiple chromosomal sites, as observed by fluorescence in situ hybridization [30]. Because it seems unlikely that all 9–12  $Ig\mu$  and >8  $Ig\omega$  genes would be linked in nurse shark, switching could involve breakage and joining between nonhomologous chromosomes, like a balanced translocation.

# Parallels with Aberrant Chromosomal Recombination

What factors might predispose recombination between the shark IgH? We suggest that a comparison can be made with AID-generated translocation in mammalian B cells. The broken DNA ends generated during SHM and CSR at Ig loci can participate in translocations where their non-lg partners tend to be AID "off"-targets [31, 32]. As found after genome-wide mapping of translocations in B cells [33, 34], those factors determining the joining partners include chromosomal proximity, proximity of the DSB, and transcriptional activity. Because AID is nonetheless preferentially recruited to Ig genes through unknown mechanisms that involve cis elements [35, 36], chromatin modifications [37-39], and AID-cofactors (reviewed in [40]), the available broken DNA ends in an activated shark B cell will primarily be from several IgH genes. We propose that the circumstances that congregate by chance to promote aberrant recombination in mammalian B cells are naturally converged in the case of shark B cells undergoing programmed DNA breaks at multiple locations.

Figure 6 shows our model for isotype switching at shark *IgH*. In single B cells, we have found 1–3 VDJ rearrangements among 9–10 autonomous Igµ genes and their alleles; primarily one VDJ is functional [15, 13]. We speculate that activation by antigen and T cell help upregulate *IgH* transcription. AID is recruited to transcribing Ig genes; mostly these are rearranged VDJ, but mutated cDNA sequence from partially rearranged and germline genes have also been isolated [15, 13]. In shark, plentiful evidence of DNA DSB and repair has been found in the J-C intron of nonswitched genes [16]. The shark *IgH* are gene duplications and may share nuclear space, as must rabbit *IgH* alleles where transchromosomal recombination occurs. There

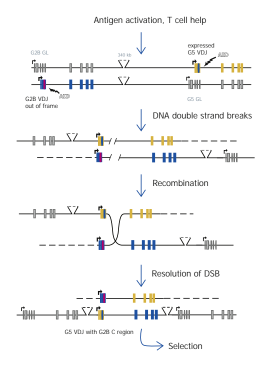


Figure 6. Proposed Switching Events at Shark IgH

(A) *IgH* genes in B lymphocyte. G2B and G5 genes are shown in relative order and orientation, as in Figure 1. Rearrangement has occurred two genes, a productive VDJ at G2B and nonproductive VDJ at G5. Both are transcribed (bent arrows).

(B) AID action in B cell. The B cell is activated by antigen and T cell help, and transcription is upregulated in G2B and G5 as well as in some nonrear ranged genes. AID is preferentially recruited to the highly transcribed genes. (C) DSB at *IgH* genes. AID mediated DNA lesions sometimes result in double strand breaks.

(D) Recombination at *IgH* genes. During DNA repair, recombination occurs between the two *IgH*. An example of the G5 VDJ switching to the upstream G2B C region is clone G5G2 21, Figure S2.

(E) Switched *IgH*. The B cells with novel antigen receptors undergo selec tion.

could be a clustering of similar *cis* elements and shared transcription factors, and the proximity of ensuing DSB presents the potential for intra- or interchromosomal recombination. What is more, AID from bony fish (zebrafish, catfish, Japanese puffer fish) has been shown to induce CSR in mouse cells [41, 42], demonstrating that early vertebrate AID already possessed the property of catalyzing recombination between two locations.

# Conclusions

Based on the established *IgH* germline gene complement in nurse shark, we have discovered that there exist cDNA sequences composed of VDJ from one gene cluster and C region from a different one: the classical feature of switched Ig H chains. The extent of switching correlated with increasing age, deliberate immunization, and AID expression, as well as being concomitant with SHM. Every Igµ gene inspected can undergo switching, and switching sites show evidence of DSB and recombination. There are no S regions; sharks possess the earliest version of Ig switching where SHM and CSR are contiguous activities that became independent only after amphibian divergence.

We hypothesize that programmed DNA lesions initiated by AID at multiple, functionally independent, and distant Ig genes prompt recombination in a shark B lymphocyte. Thus, determining factors that are predicted to lead to chromosomal aberrations and oncogenesis in other systems were co-opted by the evolving immune system in the ancestral vertebrate. Our findings demonstrate evolutionary pressure for flexible handling of pathogen during an antibody response.

#### **Experimental Procedures**

### Animals

Nurse sharks (*Ginglymostoma cirratum*) were captured off the coast of the Florida Keys. Some animals were sacrificed on arrival (shark 33, shark 626) and others had been immunized with DNP LPS (shark PI) or hen egg lysozyme (shark JS, shark J) or Ebola virus (shark GR) [11 13, 16]. These sharks range from 2 years (shark 626) to 7 years (shark GR) and are referred to as "adult" with respect to state of their immune systems. The pups were <1 week (AQ, EC, TH) to <2 months (LA) [13]. The cDNA libraries from shark 33 and shark GR epigonal organs and shark JS spleen were cloned into  $\lambda$ ZAP Express XR (Loftstrand Labs) [11, 12]. The animal protocols used in these studies were approved by the Institutional Animal Care and Use Committee of the University of Maryland and of SUNY Downstate.

## Library Screening

To distinguish the IgM subfamilies G1 G5 during cDNA library screening, we subjected all the IgM CH positive phage lysates to PCR using a primer combination (two forward primers 50% G2CH1: 5' GGACTACTC CCCTGACA 3' and 50% G4CH1: 5' GAAAGGTGGGAAAGCCCT 3' with re verse primer PANCH2 3: 5' GGAACTCAAAGTTAGGAG 3') that produced a 253 bp G2 CH fragment or a 217 bp G4 CH fragment. The remaining "nega tive" lysates would contain G1, G3, or G5 sequences and were identified by restriction endonuclease sites as described previously [11]. For direct sequencing, partial sequences were amplified from phage lysates using T3 and CH2 3' (5' ACCTGGCAKGTATARAC 3').

Screening for IgW containing sequences from the cDNA library involved double lifts hybridized to IgW VH and IgW CH. The selected phage lysates were separated into three categories: VH positive and CH positive, VH positive and CH negative, VH negative and CH positive. Nonswitched IgW sequences were verified by PCR with universal primers targeting IgW leader (JWF1: 5' GATTGCTCCWAATCTCKG 3') and IgW CH1 (two reverse primers 50% IgWCHa: 5' AGAGACTGTTTCAAATGT 3' and 50% IgWCHb: 5' GATTGCTCCGWAATCTCKG 3') and IgW CH1 (two reverse lysates containing only IgW CH but not VH were amplified with T3 and IgWCHa/b and sequenced; those containing IgW VH but not CH were ampli fied with JWF1 and IgM primer CH2 3'.

# PCR

Oligonucleotides were synthesized by Invitrogen and the areas targeted Ig H chain transcripts are illustrated in Figure S2, top. First strand synthesis of cDNA was performed using SuperScript III reverse transcriptase (Invitrogen) and oligo dT or specific primers targeting Cu: universal C region CH2 3', or G2 (G2CH2RR: 5' GATGTCAGAATGCACAC 3'), G2A specific (G2A C2a: 5' CTGAGAAGCTTCCATTCA 3'), G2B specific (G2B C2a: 5' ACACACGGG AGAAGTTAG 3'), G4 (G4CH1A: 5' TGTAAGTTCCTTTCTTGC 3'), G5 (G5C2: 5' ACTTTGAGTGGAAGTCAC 3'). This was followed by 30 cycles of PCR. Forward primers include universal primers in FR1 (V1 5 is a mix of 20% V1: 5' TGACTCAAAAAGTGGCAG 3', 80% V2 5: 5' TGAYTCAACCAGAG GCA 3'), V1 being G1 specific and V2 5 for G2 G5. Other forward primers targeted the leader: G2 specific G2V18 (5' ACCAGAATGACGACGATG 3'), G4 specific G4L (5' TTCTGACTTTCTTATCCC 3'), and G5 specific G5L2 (5' TGTTGCTGGCTTTATTAC 3'). Reverse primers targeted Cµ1 or Cµ2 and included: G2 (G2CH2R: 5' CCGTTCTTCAACCAATTG 3'), G2A specific (G2A C2b: 5' GATGTCAGAATGCACACT 3'), G2B specific (G2B C2b: 5' GA TGTCAGAATGCACACG 3'), G4 (G4CH1B: 5' ATAAATCCAGTCGTGAAG 3'), G5 (G5CH1R: 5' TCACAGGATATTTGGTCA 3').

First strand primers targeting the G5 J C intron include: OMG (at 651 bp, 5' GAATCCAATCAACTCAATTC 3'), G5I2 (at 557 bp, 5' ACTCAAATCAACA GATTGAGA 3'), OMG5 (at 551 bp, 5' AAACTCAACGACTCAAATCA 3'); PCR performed with G4L and G5I1 (5' AGAAAATTTGAAGACAGAAGT 3') produced products of about 900 bp (see Figure S4, top). The I3 series was obtained using first strand primer G5I4 (5' CAGCCTAATTGGTCAAAT 3') at

1,202 bp in the G5 J C intron; PCR with G4L and G5l3 (at 1,098 bp, 5' AGC AAATGAATGCACGA 3') generated fragments of  $\sim$ 1,498 bp.

#### **Probes and Blotting**

Probes to IgM H chain were derived from G2 cDNA (vh, cµ1, cµ2, cµ3 cµ4) had been described elsewhere [11]. Other probes were derived from an IgW cDNA sequence; the primers targeting V region (vhw, 285 bp; W5': 5' CAG GTCAAACCTTCAGTT 3'; W3': 5' AAACCAGGAAACCTGTAC 3') and CH1/ CH2 (chw, 560 bp, WCH1F1: 5' AAGATGAGATCAGCCTCC 3'; W543R: 5' GATCCTGGTACTGAAGCT 3') were generated by PCR, purified, and radiolabeled (Random Primed DNA Labeling Kit, Roche).

After hybridization, blots were subjected to autoradiography, and signal intensities of bands were quantified using a Storm 860 phosphorimaging system with ImageQuant software (GE Healthcare).

#### Accession Numbers

DNA sequences listed in Table 1, figure legends, and in supplemental fig ures were deposited at GenBank under the accession numbers JQ272792 JQ272843, JN802117, GQ359826 GQ359858, and GQ282627 GQ282630.

#### Supplemental Information

Supplemental Information includes six figures and can be found with this article online at doi:10.1016/j.cub.2012.03.060.

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