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RECONSTRUCTING IMMUNE PHYLOGENY: NEW PERSPECTIVES

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Abstract

Numerous studies of the mammalian immune system have begun to uncover profound interrelationships, as well as fundamental differences, between the adaptive and innate systems of immune recognition. Coincident with these investigations, the increasing experimental accessibility of non-mammalian jawed vertebrates, jawless vertebrates, protochordates and invertebrates has provided intriguing new information regarding the likely patterns of emergence of immune-related molecules during metazoan phylogeny, as well as the evolution of alternative mechanisms for receptor diversification. Such findings blur traditional distinctions between adaptive and innate immunity and emphasize that, throughout evolution, the immune system has used a remarkably extensive variety of solutions to meet fundamentally similar requirements for host protection.

The evolutionary development of the METAZOANS was associated with the diversification of a wide range of specialized cell-surface molecules that mediate key metabolic processes, as well as provide crucial contact interfaces and carry out a broad range of other essential functions. It is not unexpected that some of these molecules also came to function as barriers to pathogenic invasion and, in doing so, began to carry out dedicated innate immune protective functions. Whereas the simplest form of protection, barrier formation, is essentially mechanical in nature, relentless pressure from genetic variation in pathogens probably drove the evolution of such innate immune protective molecules towards diversification and, in parallel, towards integration of signalling pathways to regulate cellular responses to external stimulation. However, despite the sophistication that such innate immune mediators achieved over time, their biological complexity, by definition, would be limited by genome space, so with increasing complexity of body plan and/or increasing pathogen sophistication, they could be overwhelmed.

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Online links DATABASES The following terms in this article are linked online to: Entrez Gene: <http://www.ncbi.nlm.nih.gov/entrez/query.fcgi?db=gene> BCR | RAG | $\gamma\delta$ TCR | TCR α -chain | TCR β -chain

FlyBase: <http://flybase.bio.indiana.edu/Dscam>

FURTHER INFORMATION Zebrafish: http://www.ensembl.org/Danio_rerio

More than 500 million years ago, a TRANSPOSITION event, probably involving a recombination-activating gene (RAG)-bearing element, might have given rise to the predecessors of the rearranging antigen-binding receptors of the jawed vertebrates, which encompass the vertebrate radiations that extend from the cartilaginous fish through to humans. This is considered the defining point in the emergence of RAG-mediated (conventional) adaptive immunity^{1,2}, which has evolved to create a mechanism for deriving almost limitless variation from very few genes. Studies in traditional and non-traditional animal models, such as sharks, bony fish and birds, have brought this event and its ramifications for host defence into sharper focus. We can now predict much about how these rearranging antigen-binding receptors probably arose, what alternative pathways of immune-receptor gene evolution have occurred, what relationships exist between B- and T-cell-mediated immunity and natural killer (NK)-cell function, how complex immune recognition might be solved in species that are below the phylogenetic level of the jawed vertebrates, and what general principles link adaptive and innate immune recognition^{3,4}. It is now clear that, contrary to traditional views, jawless vertebrates, protochordates and invertebrates have also evolved sophisticated RAG-independent strategies to effect recognition and facilitate elimination of pathogens, to respond to stress, and to distinguish self from non-self^{3,5-10}. Some of the molecules and mechanisms that are used to accomplish these processes are related to those used by jawed vertebrates, whereas others seem to be unique solutions to host survival, opening up the possibility that they, or alternative forms, might have broader phylogenetic distributions than are envisioned at present.

The comparative study of immune systems of extant vertebrate and invertebrate species holds important clues about how immune recognition has evolved from generalized to highly (antigen) specific reactions and how integral links between innate and adaptive immunity have been established and maintained. These efforts are being facilitated by the availability of whole-genome information, by the improved methodology for influencing the expression of individual genes (such as transgenesis or RNA interference) and by the development of approaches to examine gene expression and protein interactions in a global context. Studies in key representative vertebrate and invertebrate species — including insects such as *Drosophila melanogaster* (fruitfly), molluscs such as snails, protochordates such as *Botryllus schlosseri* and *Branchiostoma floridae* (amphioxus), and jawless vertebrates such as lampreys and hagfish — are beginning to provide a window on different mechanisms of GERMLINE DIVERSIFICATION and SOMATIC DIVERSIFICATION (FIG. 1). Specialized innovations that influence the generation of immune diversity and the subsequent implementation of powerful selective mechanisms are being recognized^{5,8-10}.

This Review describes and compares several key immune-related systems that have extensive genomic and/or somatic diversification of innate and adaptive immune-type receptors. Certain other types of receptor that are important in host defence³ are mentioned, but detailed discussion of these is beyond the scope and intent of the main focus of this Review. In the broadest sense, observations that have been made so far show parallel evolution of various forms of immune defence, and this can be viewed as a continuum in which multiple independent shifts from limited to complex germline variation, and of specialized commitments of somatic cells, have occurred. Even in the context of the limited number of metazoan phyla that have been studied in detail, we can now describe many of the alternative mechanisms of immune recognition that have emerged at varying points in phylogeny and can identify key processes that have bridged and functionally integrated innate and adaptive immunity throughout the evolution of immune competence.

An overview of adaptive immunity in phylogeny

To understand the relationships between innate and adaptive immunity, which are intertwined functionally in modern vertebrates^{11–16}, it is first instructive to consider some key events in the evolution of adaptive (RAG-mediated) immunity. Two central observations that were made two decades ago showed that markedly disparate mechanisms — differing from each other and from the human and mouse pattern of a single rearranging receptor-gene locus that undergoes marked somatic diversification — give rise to B-cell receptor (BCR) diversity in jawed vertebrates. In sharks, the ancestors of which diverged early in jawed-vertebrate phylogeny, BCRs are encoded by hundreds of independent immunoglobulin-gene loci¹⁷, some of which undergo RAG-dependent recombination and others of which are pre-joined in the germline and therefore do not undergo combinatorial joining or generate JUNCTIONAL VARIATION¹⁸. By contrast, in birds, which diverged more recently as a group than sharks, the immunoglobulin loci consist of single variable (V) and joining (J) elements (which together encode immunoglobulin light chains), and single V and J elements plus multiple diversity (D) elements (which together encode immunoglobulin heavy chains). GENE CONVERSION, which until this discovery was an underappreciated mechanism of immune-receptor diversification, uses pseudogenes that are encoded upstream of the main gene loci to somatically change V regions in birds^{19–21} (as well as in some mammals^{22,23}); the ‘converted’ loci can also be somatically hypermutated²⁴. Given these differences in the organization and the mechanisms of somatic diversification between sharks and birds, which strongly indicate co-evolution of immunoglobulin heavy- and light-chain gene organization and diversification mechanisms²⁵, it comes as little surprise that BCRs in other species also show lineage-dependent divergence of immunoglobulin-gene organization and diversification (FIG. 2). Furthermore, certain characteristics of BCR genomic organization that were previously thought to be restricted to cartilaginous fish, such as cluster-type gene organization, are also present in bony fish²⁶ (such as catfish) and fleshy-finned fish²⁷ (such as lungfish). Despite the divergence of heavy-chain types that has now been recognized in cartilaginous²⁸ and bony^{29–31} fish, prototypical heavy-chain CLASS-SWITCH RECOMBINATION is not seen below the phylogenetic level of amphibians^{32,33}. Given the DERIVED character of the extant bony fish, the resolution of whole genomes for the pufferfish³⁴, the zebrafish (see the Online links box) and other bony-fish species could offer considerable insight into the mechanisms that have driven the evolution of BCR diversification.

In marked contrast to the BCRs, T-cell receptors (TCRs) are relatively conserved throughout the jawed vertebrates in terms of the number, organization and multiplicity of rearranging elements²⁸. The remarkable conservation of MHC class I (REF. 35) and MHC class II (REFS 36,37) structure and polymorphism throughout jawed-vertebrate phylogeny mirrors the relatively stable history of the TCRs, despite organizational changes that have taken place in the loci encoding these genes³⁸ (BOX 1). This parallel stability is probably driven by the functional interdependence and possible co-evolution of the genes encoding the TCR α -chain and TCR β -chain and the MHC molecules. The endogenous ligands for the MHC-class-I- and MHC-class-II-independent $\gamma\delta$ TCR are less clear³⁹, and explanations for the organizational stability of the genes encoding the $\gamma\delta$ TCR are less apparent. The predicted stability of TCRs, and the nearly continuous variation in complexity, gene number and structure of BCRs, is one of the most consistent themes in the phylogeny of adaptive (RAG-mediated) immunity.

Transitions from innate to adaptive immunity

Understanding the forerunners of BCRs and TCRs, and the mechanisms that promoted their divergence during evolution, might shed light on past and present relationships between innate and conventional adaptive immune recognition. Although the pathways and patterns

of evolution of rearranging antigen-binding receptors will never be known with absolute certainty, one plausible progression could start with the recruitment of an immunoglobulin-gene superfamily (IgSF) member, such as those that are found in species as phylogenetically distant from vertebrates as *Caenorhabditis elegans* and *D. melanogaster*⁴⁰. The most recent common ancestor of the rearranging antigen-binding receptors would have been such an IgSF member; it would have either pre-existed as, or subsequently given rise to, an immunoglobulin V-region-type innate immune cell-surface receptor or a cell-surface adhesion molecule, as is indicated by one interpretation of the crystallographic analysis of shark NAR (new antigen receptor)⁴¹, an immunoglobulin heavy-chain homodimer that binds specific antigens⁴². The J motif (FGXGTXLXV, where X denotes any amino acid), which promotes homo- and heterodimer formation, might have been encoded by a separate exon and become contiguous with the V-encoding region through intron loss, a recognized feature of genome evolution³⁴. A VJ structure is the likely target of the key transposition event. Although it is overly simplistic to assume that such a VJ molecule persists in contemporary life forms, several single-V molecules or V-related molecules (or, more precisely, the features of their sequences) are interesting to consider⁴³; however, any interpretation of such data also needs to take into account canonical V regions that do not function in immune recognition.

The origins of RAG1 and RAG2 themselves are unclear; however, a convincing homologue of *RAG1* has been found in the sea-urchin genome^{3,44} and is contiguous with a structure that is homologous to *RAG2* (REF. 3, and S. Fugmann and J. Rast, personal communication). A single RAG-mediated transposition event into the VJ region could have created a potential junctional interface. Variation at the junctions during RECOMBINATION-SIGNAL-SEQUENCE-mediated recombination would then be introduced by the activation of DNA-binding proteins and repair enzymes that are not specific to the immune system, as well as terminal deoxynucleotidyltransferase, which is an error-prone DNA polymerase of the type X (PolX) family⁴⁵ that mediates non-templated base insertion during NON-HOMOLOGOUS END JOINING, the main source of somatic variation in BCRs and TCRs⁴⁶. The capture of seemingly unrelated mechanisms of gene rearrangement and diversification seems to be a generalized phenomenon in the development of immune strategies (discussed later). Class-switch recombination (another form of somatic reorganization that occurs in BCR-gene loci in organisms as phylogenetically distant from mammals as amphibians³²), as well as somatic hypermutation and gene conversion, is mediated by many of the same DNA-binding proteins that are involved in V, D and J joining, as well as by activation-induced cytidine deaminase (AID)⁴⁷. AID is present in bony fish, and interestingly, zebrafish AID can catalyse class-switch recombination when introduced into mammalian B cells⁴⁸. So, the appearance of a class-switch reaction was dependent on the evolution of switch-region sequences. AID-like and uracil-DNA glycosylase (UNG)-like proteins, which are involved in these processes, had ancient origins, co-evolving with innate immune antiviral defences⁴⁹⁻⁵¹.

Over time, V-gene duplication and germline substitutions, which were focused in COMPLEMENTARITY-DETERMINING REGION 1 (CDR1) and CDR2, coupled with combinatorial joining, imparted a considerable selective advantage in terms of deriving receptor diversity. Somatic hypermutation of immunoglobulin genes, which is a central feature of BCR maturation, occurs in the earliest extant jawed vertebrates — the cartilaginous fish^{52,53} — and might be an accentuation of a genome-wide process^{54,55}. ALLELIC EXCLUSION of BCR- and TCR-gene loci, which is an integral component of single-cell immune commitment that is present throughout jawed-vertebrate phylogeny⁵⁶, is also a feature of the loci encoding olfactory receptors^{57,58} and protocadherins⁵⁹, emphasizing again the integration of pre-existing mechanisms in the evolution of immunity. The full potential of these features did not come

to fruition until the co-evolution of mechanisms to select and expand individual clones of immune cells.

Collectively, these observations indicate that the evolution of the V-type antigen-binding receptors from mediators of innate immunity to the fore runners of adaptive (RAG-mediated) immunity probably occurred incrementally through multiple recruitment events rather than as an immediate consequence of an immunological ‘big bang’⁶⁰, a term that more aptly relates to the apparent absence of RAG in jawless vertebrates. RAG-mediated adaptive immunity is one of the most elaborate examples of the integration of disparate cellular and developmental processes to achieve a highly specialized, lineage-restricted biological function: that is, the use of segmental gene re arrangement to produce a fertile receptor-gene pool for the selection and clonal expansion of individual somatic cell lineages. Many of the elements that we collectively recognize to be common features of jawed-vertebrate adaptive immune receptors, including multigene complexity, somatic hypermutation and gene conversion, also diversify immune-type receptors in jawless vertebrates, protochordates and invertebrates (discussed later).

NK and NK-type receptors in jawed vertebrates

Further insight into relationships between innate and adaptive immunity can be gained from a basic consideration of NK cells, which share a common progenitor with T and B cells but do not somatically rearrange their receptors. NK cells are primary innate immune sentinels for the detection of certain viral infections and malignant transformation. The main classes of NK-cell receptors in mammals are either related to C-type lectins or members of the IgSF, and they are expressed by NK cells and natural killer T (NKT) cells⁶¹. In humans, the main diversified NK-cell receptors are killer-cell immunoglobulin-like receptors (KIRs), which are activating or inhibitory TYPE I TRANSMEMBRANE PROTEINS with immunoglobulin C2 (constant) ectodomains. In mice, the corresponding receptors are members of the Ly49 gene family and are activating or inhibitory TYPE II TRANSMEMBRANE PROTEINS. The extracellular regions of Ly49 gene products are single C-type-lectin domains. Haplotypic variant and invariant genetic regions of KIRs and Ly49 proteins are interspersed similarly⁶²; extensive haplotypic variability in KIRs arises through meiotic recombination⁶¹. Certain KIRs and Ly49 proteins recognize MHC class I molecules. Other NK-cell receptors recognize a broad range of stress-induced molecules and other non-classical MHC molecules^{63,64}, as well as viral gene products^{65,66} and other NK-cell receptors⁶⁷. NK-cell activity has a broad phylogenetic distribution extending to invertebrates⁶⁸. A lectin-like, NK-cell-type receptor has been characterized in a protochordate⁶⁹; however, its sequence similarity to other NK-cell receptors is remote, and its function is unknown. Putative C-type-lectin NK-cell receptors have been reported in bony fish^{70–72}; however, they encode members of the type 2 family of C-type-lectin domains and not the type 5 family, in which mammalian NK-cell receptors are classified.

Determining the evolutionary origins of NK cells and their receptors could be important for understanding the phylogeny of adaptive immunity. However, given the diversity of NK-cell receptors that have already been identified in mammals, this could prove daunting. The NITRs (novel immune-type receptors) — a large multigene family of NK-cell-like activating and inhibitory cell-surface receptors in bony fish — show an intriguing link between conventional adaptive immune receptors and innate immune receptors⁷³ (TABLE 1). Most NITRs consist of an amino (N)-terminal membrane-distal immunoglobulin V-region-type extracellular domain and a membrane-proximal intermediate (I)-type extracellular domain; other NITRs have only a single V-region-type extracellular domain. Most NITRs have IMMUNORECEPTOR TYROSINE-BASED INHIBITORY MOTIFS (ITIMs) in the cytoplasmic domain and therefore are probably inhibitory. Other NITRs have positively charged residues in their transmembrane regions and probably interact with adaptor molecules that contain

IMMUNORECEPTOR TYROSINE-BASED ACTIVATION MOTIFS (ITAMs)⁷⁴. Some NITRs have amino-acid motifs that are related to the IMMUNORECEPTOR TYROSINE-BASED SWITCH MOTIF (ITSM) that has been described in members of the mammalian CD2, SIRP (signal-regulatory protein), SIGLEC (sialic-acid-binding immunoglobulin-like lectin), CEA (carcinoembryonic antigen) and PIR (paired immunoglobulin-like receptor) families, potentially allowing the modulation of signalling by differential interaction with SRC homology 2 (SH2)-domain-containing adaptor proteins (thereby allowing either activating or inhibitory function)⁷⁵. Other NITRs lack a transmembrane region and resemble the decoy molecules that have been described in other activating and inhibitory receptor systems⁷⁶.

High-confidence molecular modelling of NITR V regions shows them to resemble the V domains of immunoglobulin and TCRs⁷⁴; in some cases, they have motifs similar to the J regions of BCRs and TCRs. In pufferfish, 13 main families of NITR V regions are encoded at a single compact locus⁷³. In zebrafish, 12 families of NITR V domains are encoded at a single locus on chromosome 7 (REF. 74), and 2 are encoded on chromosome 14 (see zebrafish in the Online links box). Although the NITRs in pufferfish and zebra fish are unequivocal ORTHOLOGUES, the respective gene families do not show absolute SYNTENY. Approximately 150 alleles and 45 structural variants of NITRs have been identified in zebrafish⁷⁴; only one is an unequivocal activating form. Intergenic variation is regionalized within the large NITR1 gene family, and it is reminiscent of the CDR variation that is seen in BCRs and TCRs. NITR genes are transcribed in several different cell lineages in bony fish⁷⁷, including NK-type cells and cytotoxic T lymphocytes⁷⁸; at least one NITR protein has a high degree of specificity for an allogeneic target (J.P.C., D. Eason and N. Miller, unpublished observations). The evolution of NITRs is extraordinarily rapid, even within a species, a characteristic that is shared by immunoglobulins, TCRs and KIRs, and this has been attributed to a gene BIRTH-AND-DEATH process⁷⁹. The allelic and haplotypic variation in the NITR genes in a single strain of zebrafish is reminiscent of the variation in KIRs from various human subpopulations^{61,62}. KIRs and NITRs share other features, including differential expression by individual cells and expression by NK cells and T cells⁸⁰. Notably, a recent release of the zebrafish genome sequence has uncovered two groups of genes with sequence similarity to human KIRs, on chromosome 2 and chromosome 7 of the zebrafish (see zebrafish in the Online links box).

At least two explanations could account for the evolutionary origins of NITRs: they might be derived from a recombination event between a member of an activating and inhibitory gene family and elements of a locus encoding a rearranging receptor or V-type receptor; or they might have a common origin with the V-region-containing molecules NK-cell protein 30 (NKp30) and NKp44 (REFS 81,82) or, as has been suggested, with a family of activating and inhibitory membrane proteins containing V domains that has been found in a jawless vertebrate⁸³. In either case, the NITRs present a compelling case for parallel evolution of BCR- and TCR-type V-region diversity in the context of innate immunity.

Anticipatory immunity in jawless vertebrates

Lymphocytes can be traced to jawless vertebrates. In lampreys, mononuclear cells with morphological features of lymphocytes have been identified in various tissues and in the peripheral blood⁸⁴. Accelerated allograft rejection has been documented in lampreys⁸⁴, as have humoral responses to immunization with bacterial and cellular antigens; however, the circulating AGGLUTININS are not related structurally to antibodies²⁸. Extensive EXPRESSED SEQUENCE TAG (EST) surveys have failed to identify typical lymphocyte-specific cell-surface determinants in lampreys^{85,86}, with the exceptions of a predicted gene product that shows a distant relationship to CD4 (among other molecules)⁸⁷ and another monomorphic gene product that has cytosolic ITIMs and can be modelled to TCR V regions but shows only remote primary-

structure similarity to jawed-vertebrate TCRs⁸⁷. However, Ikaros and SPI, which are transcription factors that function in higher-vertebrate lymphocyte development and differentiation, have been identified in jawless vertebrates^{88–91}, as have ETS- (M. Anderson, personal communication), GATA-binding protein 2 (GATA2)- and GATA3-type molecules (J.P.C. and R. Haire, unpublished observations). Using a V-region-directed cloning technique in lamprey lymphocytes, a single gene was identified that resembles *VpreB*, which encodes a surrogate BCR light-chain component that is expressed early in lymphocyte ontogeny in higher vertebrates but does not participate directly in the function of the mature BCR⁹². The lamprey *VpreB*-like molecule might reflect the character of an early common ancestor that might or might not have mediated an immune function⁹³. In addition to the V-region-related sequences in hagfish⁸³, additional immunoglobulin V-region-related sequences that have so far been identified in lampreys include nectin and Lutheran blood-group-antigen-like genes, neither of which is related closely to BCRs or TCRs⁹³. Whole-genome analysis might yet uncover BCR- or TCR-related candidates.

Alternative mechanisms might mediate adaptive immunity in jawless vertebrates. Antigenic or mitogenic stimulation of short-term cultures of lamprey peripheral-blood leukocytes results in considerable proliferation of lymphocytes⁹. Subtracted cDNAs recovered from these cell cultures are highly enriched in sequences encoding a particularly complex set of LEUCINE-RICH REPEAT (LRR)-containing extracellular molecules. Notably, LRRs are a core feature of markedly diverse forms of pattern-recognition molecule that mediate receptor–ligand interactions throughout phylo geny^{94–98}. The LRR-encoding cDNAs in lampreys are encoded at a single locus. On the basis of the relatively high degree of hypervariation in the LRR cassette, the lack of variation in the N and carboxyl (C) termini, and expression in lymphocyte-like cells, the molecule has been designated a variable lymphocyte receptor (VLR). VLRs consist of the following regions: a 30–38-residue N-terminal region, an 18-residue N-terminal LRR, followed by up to nine 24-residue LRR cassettes, a 13-residue connecting peptide, a 48–58-residue C-terminal LRR, a threonine- and proline-containing stalk, a glycosylphosphatidylinositol (GPI) anchor and a C-terminal hydrophobic tail (TABLE 2). The N- and C-terminal domains are relatively invariant, but the number and sequence of the internal LRR motifs are variable. The *VLR* cDNAs seem to have arisen by recombination of LRR cassettes, which are found in different orientations in the germline locus and can be inserted into the coding frame of the active *VLR* gene, possibly through gene conversion. PCR amplification of genomic DNA and comparison of product lengths in erythrocytes and lymphocyte-like cells indicates that diversity in the VLR repertoire is derived by somatic recombination but is presumably RAG independent. The structural characteristics of VLRs, as well as cell-surface staining with VLR-specific antibodies, indicate that they are membrane bound. Disruption of the GPI linkage might function as the mechanism to release the membrane-bound VLR, in a manner that is generally reminiscent of lymphocyte-bound immunoglobulin. Single-cell PCR is consistent with haplotypic exclusion, a characteristic of lymphocyte-mediated immunity and a central feature of clonal selection in RAG-mediated adaptive immunity. VLRs that undergo similar genetic rearrangement have also been characterized in hagfish⁹⁹.

The postulated role of the VLRs as alternative mediators of adaptive immunity might have an even more ancient history within the broader functional roles of other metazoan LRR-containing proteins, which include the Toll-like receptor (TLR)⁹⁴, CATERPILLER (caspase-recruitment domain, transcription enhancer, R (purine)-binding, pyrin, lots of LRRs)¹⁰⁰ and Slit-homologue protein families^{101,102}. The TLRs have particularly broad distributions in phylogeny and are closely involved in mammalian immune surveillance^{103–105}. TLRs, which comprise one of the better-recognized functional relationships that links innate and adaptive immunity^{12,106}, recognize molecular patterns that mainly consist of conserved epitopes found on pathogens^{6,107}. Pattern recognition by

TLRs does not rely on somatic change; however, there is increasing evidence for functionally advantageous polymorphisms that, for example, improve resistance to bacterial infection¹⁰⁸, as well as increasing evidence for combinatorial pairing¹⁰⁹. Intracellular LRR-containing proteins that mediate recognition of intracellular infection, modulate signalling and regulate apoptosis have also been described^{100,110}. Our preliminary phylogenetic analysis of lamprey and hagfish VLRs indicates that a single LRR protein might have been recruited from a pre-existing family; this protein subsequently would have undergone rapid evolution and divergence (to function as an immune mediator). The VLRs might be a derived TLR family that could supplant and/or complement the function of these receptors in jawless vertebrates¹⁰³.

Key questions regarding VLRs relate to the mechanisms and regulation of somatic-recombination events and to the clonal-selection characteristics of the lymphocyte-like cells that express VLRs. Notwithstanding the differences in terms of the mediators of adaptive immunity, a challenging and broader issue concerns the relationships between the VLR-expressing lymphocytes and those cells that express TCRs or BCRs and are distributed throughout the phylogeny of jawed vertebrates. A corollary issue is whether there are novel mechanisms for the generation of immune-receptor diversity in protochordates and other invertebrates.

VCBPs in protochordates

As indicated previously, the extant jawless vertebrates are highly derived, and despite the likelihood that they have an anticipatory adaptive immune system, their immune mediators might be more similar to other LRR-based systems than to the genes encoding immunoglobulin domains. In this regard, inferences drawn from more BASAL species can sometimes provide more insights than those drawn from species that are more related to the jawed vertebrates in terms of phylogenetic position¹¹¹: that is, although a bony fish shares a more recent common ancestor with humans than does a protochordate (FIG. 1), protochordates might still retain certain characteristics that are more closely related to those found in humans, as a consequence of evolutionary derivation. In this regard, the basal morphological and physiological characteristics of the CEPHALOCHORDATE amphioxus might shed light on links between innate immune recognition and immunoglobulin-type receptors that are related to those that mediate RAG-based adaptive immunity.

Early efforts to clone adaptive immune-like genes from amphioxus using methods such as cross hybridization and short-primer PCR were unsuccessful; however, a SECRETION-SIGNAL-PEPTIDE-SELECTION-based approach identified members of a multigene family that encode an immunoglobulin-type molecule (known as V-region-containing chitin-binding protein, VCBP) that consists of a pair of N-terminal V domains and a single C-terminal chitin-binding domain⁷ (TABLE 2). Five families of VCBPs are distinguished by sequence differences in their V regions⁷. The N-terminal V regions of VCBP2 and VCBP5 are closely related, although the remainder of each molecule is divergent. The structure of the N-terminal V domain of a VCBP3 molecule¹¹² has been solved to 1.15 Å (J. Hernandez Prada, R. Haire and D. Ostrov, personal communication).

VCBPs — unlike BCRs, TCRs and some NITRs — lack characteristic J segments and are not membrane anchored. However, the N-terminal V domains of VCBP2 and VCBP5 contain a region of particularly extensive diversity^{7,113}, which is consistent with immune-type function. Systematic sampling of the local amphioxus population indicates an extraordinary number of polymorphic alleles across this region of diversity that are inherited in a Mendelian manner (REF. 113, and N. Schnitker and L.J.D., unpublished observations). This hypervariable region probably has a function that is under strong selective pressure, such as

pathogen recognition¹¹³. VCBPs have also been identified in *Ciona* species¹¹³, and the possibility that somatic variation occurs in VCBPs is being evaluated.

RNA *in situ* hybridization shows that VCBPs are expressed abundantly and (apparently) specifically in the gut of amphioxus, which is an important site of potential pathogen infiltration in this exclusive filter feeder. Amphioxus accounts for up to 70% of the total biomass in certain coastal waters. In such large populations, gene selection can occur at the level of the population rather than on an individual level^{113–115}, which is in contrast to conventional adaptive immune mediators, for which selection mainly occurs at the level of the somatic cell lineage within an organism. Sea water sustains high levels of bacteria, fungi and viruses and undergoes marked seasonal variation. Although speculative, the function of VCBPs might be to recognize determinants on various pathogens. There is considerable precedent for IgSF-related viral receptors, including those for poliovirus (poliovirus receptor)¹¹⁶, reovirus (junctional adhesion molecule, JAM)¹¹⁷, HIV (CD4)¹¹⁸, mouse hepatitis virus (CEA-related cell-adhesion molecule 1, CEACAM1)¹¹⁹, measles virus (signalling lymphocytic activation molecule, SLAM)¹²⁰ and others. Other potential functions for VCBPs include binding to fungal or bacterial surfaces, direct opsonization, and self- and non-self-recognition^{3,7}. It is also possible that the V regions of VCBPs are pathogen specific and that the chitin-binding domains bind microbial *N*-acetylglucosamine, thereby inhibiting biofilm deposition, which is a key step in the colonization of a host by certain bacteria, through the formation of impenetrable barriers^{121–123}. Variation in VCBP V regions and BCR and TCR V regions is concentrated in different relative positions and is achieved through different mechanisms, presenting a likely case for evolutionary convergence of the V domain in innate and adaptive immune functions, respectively.

Variation in invertebrate immune receptors

Similar to vertebrates, invertebrate species are under nearly continuous assault from various bacterial, fungal, protozoan, parasitic and viral pathogens. Traditionally, it was thought that several distinct, but relatively invariant, innate immune mediators countered such challenges in invertebrates. However, invertebrates have relatively extensive polymorphism of their immune-gene loci¹¹⁵, and it is now recognized that they show memory in immune-type defence¹²⁴ and can probably mount various anticipatory immune responses¹²⁵. A series of investigations in molluscs has provided insight into a previously unrecognized aspect of innate immune-receptor diversification and, together with a recent report on *D. melanogaster*¹⁰ (discussed later), extends our consideration of ‘adaptive’ immunity beyond the vertebrates and protochordates.

In the snail *Biomphalaria glabrata*, a diverse family of IgSF-domain-containing fibrinogen-related proteins (FREPs) is secreted by circulatory haemocytes. The expression of FREPs is upregulated on infection with a trematode (a type of parasitic flatworm); FREPs then bind to trematode sporocysts and precipitate soluble parasite-derived products from solution¹²⁶. At the molecular level, FREPs consist of one or two tandem IgSF domains at the N terminus, as well as a fibrinogen-like domain at the C terminus, and they have lectin-like activity^{126,127} (TABLE 2). Fibrinogen is associated with blood coagulation in vertebrates; proteins that contain fibrinogen-like domains have previously been implicated in innate immune-defence functions in vertebrates and invertebrates¹²⁸. FREPs, similar to the VCBPs in protochordates, are examples of proteins that have recruited more than one type of framework domain during the independent evolution of immune-defence pathways (discussed later).

Sequence variation in FREPs occurs in the IgSF and the fibrinogen-like domains⁸. Each individual snail can express various unique alleles within the different subfamilies. The

FREP alleles seem to be under strong selective pressure, and no single allele has come to dominate the population; instead, a diverse repertoire is maintained, which is potentially an advantage for an immune mediator. Diversity in the sequence encoding the N-terminal immunoglobulin-like domain of FREP3 is three- to fourfold higher than in control, unrelated genes. Taken together, a mathematical estimate and Southern-blot data indicate that a specific number of germline sequences are maintained across each subfamily and could account for all of the known variation, including nucleotide changes and diversification through recombination^{8,129}. The observed diversity in *FREP3* cDNAs probably derives from somatic modification of *FREP3* genes⁸. It is unclear whether there is a component of cell-lineage specificity (and possibly selection) in FREPs that amplifies particular sequences from the pool of observed variants.

Several other recent examples show extraordinary mechanisms for generating somatic variation that lie outside the traditional concepts of molecular genetics and evolution. In contrast to the mutation process that is used by FREPs, other invertebrates probably use different methods to diversify immune-related molecules. The penaeidin genes, which encode antimicrobial peptides in crustaceans, occur in sets of relatively few loci, but they express an extensive repertoire¹³⁰. Complex transcriptional processing of a lipopolysaccharide-inducible gene has been shown in sea urchins¹³¹. The possibilities for alternative splicing increase markedly when exons have undergone tandem duplication¹³², such as for the *D. melanogaster* Down's syndrome cell-adhesion molecule (*Dscam*) gene, which functions in complex neuronal processing by generating tens of thousands of protein variants from a single gene through differential RNA processing^{132,133}. Both the complexity and expression patterns of *Dscam* isoforms in haemocytes indicate that *Dscam* might have a role in immune function^{133,134}. Recent studies have defined differences in the expression of *Dscam* isoforms in haemocytes, fat cells and brain tissue¹⁰. Furthermore, secretion of specific *Dscam* isoforms has been confirmed at the protein level. *D. melanogaster* larvae carrying transallelic hypomorphic and amorphic mutations had reduced *Dscam* expression and less efficient phagocytosis of pathogenic bacteria, which was also achieved using *Dscam*-specific antibody and in RNA-interference studies. Directed binding of specific *Dscam* isoforms to bacteria was also shown, using soluble *Dscam*-immunoglobulin fusion proteins. *Dscam*-isoform variation is seen in multiple insect orders, which is consistent with it having an ancient origin. Collectively, these data underscore a role for complex RNA processing as another mechanism of immunological diversity. Other mechanisms can also introduce high levels of sequence diversity into receptor binding sites^{135,136}, for example, when accompanied by an adjacent retroelement.

Variation in immune receptors and pathogens

Phylogenetic studies of immune molecules clearly indicate that various mechanisms are used to diversify relatively limited amounts of genetic material to create a diverse set of receptor structures, the complexity of which is immense. The greater the amount of genetic variation that can be derived from a single gene or groups of genes, the more adaptable the host will be in responding to molecular variation shown by pathogens. The events that are likely to have occurred during the evolution of immunoglobulin and the TCR, as well as some of the other diverse immune receptors that are described here, show the successful integration of pathways that share important properties with those that are used to achieve genetic variation in viruses, bacteria, fungi and protozoan parasites. So, there is a co-evolutionary struggle in which selection acts on DNA in the host and in pathogens to mediate sequence variation and diversity for improving detection of infection in one case or for promoting evasion of immunity in the other (BOX 2).

What might be less appreciated is that the functional motifs (for example, LRRs) in the various receptors are related to those found in the pathogens themselves, including in protozoan parasites, bacteria (such as *Listeria monocytogenes* and group B *Streptococcus*) and fungi¹³⁷. Over evolutionary time, a large number of effector molecules, including those that mediate innate and adaptive immune responses, have been created using frameworks of immunoglobulin, lectin and/or LRR domains (FIG. 3). Molecules such as VCBPs, FREPs and Dscam have integrated immunoglobulin and lectin domains that might coordinately function in immune recognition. Molecules that are associated with these structures can recognize and interact with diverse arrays of protein and/or carbohydrate patterns to mediate complex signalling events and effector functions¹³⁸. Some pathogens have recruited these framework structures into some of their effector molecules, in effect creating a case in which selection acts on similar structures in the genetic repertoire of the host and the pathogen (FIG. 3).

Concluding remarks

Initially, adaptive and innate immunity were thought to be temporally separable processes involving different cell types. However, as more details have been learned about both forms of immunity, a growing awareness is emerging of a single, integrated process in which innate and adaptive effectors interact coordinately to counter a wide range of invasive pathological challenges⁶. Whereas innate immunity is generally considered to be the more phylogenetically ancient, this system has diversified and refined its functions over evolutionary time such that it persists in multiple forms and shows lineage-specific diversity. In this light, it is not unexpected that innate immune surveillance and recognition in modern organisms can achieve sophisticated discrimination between the ‘signatures’ of pathogens and those of non-pathogens or self-molecules. Indeed, a combination of specificity and degeneracy in recognition is common to innate and adaptive immune receptors¹⁴. The evolutionary success of the animal (as well as the plant) phyla outside the jawed vertebrates that do not undergo V(D)J recombination of immunoglobulin and TCR genes is a testament to the ability of non-RAG-mediated systems to defend the host against pathogenic threats.

Studies in various model systems allow an integrated view of adaptive and innate immune function. The dependence of mammalian immunity on T- and B-cell function and, in turn, the dependence of these cell types on RAG-mediated somatic recombination and subsequent diversification of antigen-receptor genes has understandably resulted in a focus on lymphocytes as the central components of mammalian immunity. However, increasing attention has recently been paid to the interactions of T and B cells with innate immune effectors, and this has produced important insights regarding functional bridges between innate and adaptive immune cells during infection^{11–16,139,140}. An increasingly rich picture of the diversity of innate immune surveillance and recognition in vertebrates is developing. So, although immunity in modern mammals heavily depends on T and B cells, the activation of these cells and their subsequent roles in elimination of infection depend on their interactions with cells that express innate immune receptors, specifically dendritic cells and other antigen-presenting cells, as well as on the influences of a myriad of cytokines and soluble factors, such as the interleukins, interferons and complement.

The recent discoveries of markedly diversified immune-type genes in species that lack RAG-mediated adaptive immune systems have blurred the distinction between purely innate and purely adaptive immune receptors. The germline-diversified V domains in NITRs are an example of an adaptive immune feature in an innate immune receptor^{73,74}, whereas germline joining of immunoglobulin-gene clusters in cartilaginous fish is an example of an innate immune feature in an adaptive immune receptor¹⁸. The Dscam variants in insects¹⁰, the FREPs in snails^{8,126}, the VLRs in the jawless vertebrates lampreys⁹ and hagfish⁹⁹, and

the VCBPs in the protochordate amphioxus^{7,113} all share certain fundamental characteristics with the adaptive antigen-binding immune receptors of jawed vertebrates.

Our historical considerations of the evolution of adaptive immunity have centred on the origins of the adaptive antigen-binding immune receptors themselves, as well as the MHC class I and class II molecules on which T-cell function depends so closely and the origins in phylogeny of which seem to coincide with the emergence of the antigen-binding receptors. Many efforts have been undertaken to label ‘primordial’ antigen-binding receptors in species outside the jawed vertebrates, under the presumption that such molecules exist and would appear in a recognizable form today. However, if one takes a more integrated view of the evolution of immunity as a whole and considers the wealth of diversity and discrimination that are available through using innate-type immune receptors, such intense focus on a single arm of a vast system is beginning to seem misplaced. A more realistic consideration of innate immune receptors might be necessary to understand the origins of adaptive immunity, particularly in cases in which modern genes have been assembled from ‘parts’ of other genes.

A wealth of examples of potential evolutionary vestiges occur in the modern mammalian immune system, including B1 cells and natural IgM, $\gamma\delta$ T cells, ‘invariant’ TCRs, NK cells and their sets of receptors, serum lectins and the various complement pathways. Immune responses in contemporary jawed vertebrates have integrated these diverse systems and cellular interactions to achieve a collective goal of host defence. As these systems are numerous, diverse and under constant evolutionary pressure from the adaptation of pathogens, it is probable that following the lineage of one particular immune effector (even immunoglobulin or the TCR itself) in phylogenetically ancient species will provide only a small part of a tremendously complex picture. It is possible that natural selection has produced (and even eliminated) innovative immune-type genes continuously over evolutionary time and that many such genes will be uncovered as additional non-traditional species are investigated. This view is supported by the diversity of immune mechanisms that are now being discovered in a lineage- and taxon-dependent manner. The successive steps in the evolution of integrated immune complexity might never be known; however, it is probable that unravelling the networks that guide host defence in disparate phyla will continue to provide valuable information that can direct us in approaching and manipulating the complex mechanisms that effect immune function in multicellular organisms.

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Glossary

METAZOANS	All animals that are above the phylogenetic level of protozoans and sponges.
TRANSPOSITION	Movement of a segment of DNA from one position in the genome to another position (or to a different genome).
GERMLINE DIVERSIFICATION	Changes in DNA sequence that occur in the germline of a species and are heritable.

SOMATIC DIVERSIFICATION	Changes in DNA sequence that occur in individual cells and their progeny. Traditionally, this process has been associated with lymphocytes and is brought about during gene rearrangements, as well as through gene conversion and somatic hypermutation.
JUNCTIONAL VARIATION	The introduction of genetic changes at the joining interfaces of rearranged variable (V), diversity (D) and joining (J) segmental elements in immunoglobulins and T-cell receptors.
GENE CONVERSION	A homologous-recombination event in which the donor gene(s) remains unmodified and an acceptor gene acquires the recombined segment. In chickens, variable (V)-region pseudogenes are donors that modify the functional, rearranged V gene in bursal follicles, and this process generates a diverse pre-immune repertoire.
CLASS-SWITCH RECOMBINATION	The process by which the heavy-chain class that is associated with a specific VDJ rearrangement is changed through recombination during B-cell ontogeny.
DERIVED	Reflecting characteristics that are acquired subsequent to divergence from a common ancestral form.
HISTOCOMPATIBILITY REACTION	The reaction to, and/or rejection of, individual cells or tissues that are introduced from one individual to a genetically different individual.
RECOMBINATION SIGNAL SEQUENCE	A conserved genetic element that constitutes a recognition site for the V(D)J recombinase proteins, which are encoded by the genes recombination-activating gene 1 (RAG1) and RAG2. These sites consist of a palindromic 7-base-pair sequence that is immediately adjacent to the coding gene segments — variable (V), diversity (D) or joining (J) — and is separated by a 12- or 23-base-pair spacer from a relatively conserved 9-base-pair sequence.
NON-HOMOLOGOUS END JOINING	A generic mechanism for repair of double-stranded DNA breaks. It joins hairpin coding ends that form during V(D)J recombination.
COMPLEMENTARITY-DETERMINING REGIONS	(CDRs). The most variable parts of immunoglobulin and the T-cell receptor. These form loops that make contact with specific ligands. There are three such regions (CDR1, CDR2 and CDR3) in each variable domain.
ALLELIC EXCLUSION	A mechanism by which antigen receptors of a single specificity are expressed at the cell surface of a lymphocyte. This is an integral step in the clonal commitment of a cell lineage.

**TYPE I
TRANSMEMBRANE
PROTEIN**

An integral membrane protein that is composed of an amino-terminal extracellular region, a transmembrane domain and a carboxy-terminal intracellular region (for example, T-cell receptors).

**TYPE II
TRANSMEMBRANE
PROTEIN**

An integral membrane protein that is composed of an amino-terminal intracellular region, a transmembrane domain and a carboxy-terminal extracellular region (for example, Ly49).

**IMMUNORECEPTOR
TYROSINE-BASED
INHIBITORY MOTIF**

(ITIM). A structural motif that contains tyrosine residues and is found in the cytoplasmic tails of several inhibitory receptors, such as the low-affinity Fc receptor for IgG Fc γ R1B and paired immunoglobulin-like receptor B (PIRB). The prototype six-amino-acid ITIM sequence is (I/V/L/S)XYXX(L/V), where X denotes any amino acid. Ligand-induced clustering of these inhibitory receptors results in tyrosine phosphorylation, often by SRC-family protein tyrosine kinases, and this provides a docking site for the recruitment of cytoplasmic phosphatases that have a SRC homology 2 (SH2) domain.

**IMMUNORECEPTOR
TYROSINE-BASED
ACTIVATION MOTIF**

(ITAM). B-cell, T-cell and natural-killer-cell receptors are non-covalently associated with transmembrane proteins that contain one or more ITAMs. The amino-acid sequence of an ITAM is (D/E)XXYXX(L/I)X₆₋₈YXX(L/I), where X denotes any amino acid. This is tyrosine phosphorylated after engagement of the ligand-binding subunits, which triggers a cascade of intracellular events that results in cellular activation.

**IMMUNORECEPTOR
TYROSINE-BASED
SWITCH MOTIF**

(ITSM). This motif has the amino-acid sequence TXYXX(V/I), where X denotes any amino acid. It recruits many of the same signalling molecules as do immunoreceptor tyrosine-based inhibitory motifs (ITIMs) and immunoreceptor tyrosine-based activation motifs (ITAMs), but it also recruits SAP (signalling lymphocytic activation molecule (SLAM)-associated protein).

ORTHOLOGUES

Similar sequences that are found in different species and can be attributed to descent from a common ancestral sequence.

SYNTENY

The same order of genes occurring on chromosomes that are present in different organisms.

BIRTH AND DEATH

In the context of multigene families encoding immune molecules, this involves the creation and extinction of alleles, and it is co-driven by pathogen load during evolution.

AGGLUTININ

A soluble protein that occurs naturally or in response to immunization and agglutinates a particulate antigen.

EXPRESSED SEQUENCE TAG	The term can be applied to an immunoglobulin molecule or to proteins that are unrelated to immunoglobulins.
LEUCINE-RICH REPEAT	(EST). A genetic sequence that corresponds to an mRNA or a region of an mRNA.
BASAL	(LRR). Domains that contain LRRs have a conserved solenoid structure, typically of 20–29 residues and containing an 11 amino-acid consensus sequence, LXXLXLXX(N/C)XL, where X denotes any amino acid. These domains lack considerable identity or similarity in the amino acids surrounding this structure, both between and among families. Sequence substitutions in LRR-containing proteins are associated with changes in specificity and relative affinity towards specific determinants.
CEPHALOCHORDATE	Reflecting the shared characteristic(s) of a common ancestral form.
SECRETION SIGNAL-PEPTIDE SELECTION	A member of the subphylum Cephalochordata. In this Review, cephalochordates are represented by amphioxus, which has a notochord, dorsal nerve cord and pharyngeal gills that are well developed in the adult stage.
	A cloning method that is based on the rescue of ampicillin resistance in a vector that encodes a defective β -lactamase, by the substitution of an exogenous leader, as well as additional coding sequence, from transmembrane or secreted proteins.

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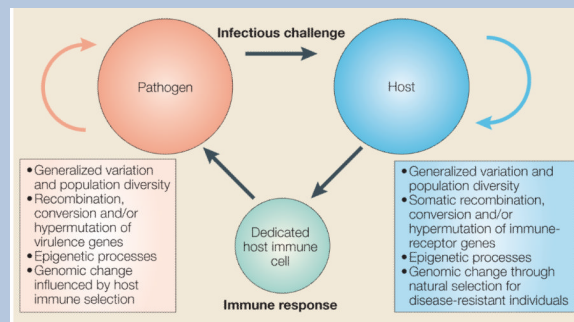
Box 1**Alternative mediators of histocompatibility**

Similar to the B-cell receptors (BCRs) and T-cell receptors (TCRs), MHC class I and class II molecules seem to be restricted to the jawed vertebrates. A rigorous genome-based search for homologues of the genes encoding these molecules in *Ciona intestinalis* (a solitary protochordate) was unsuccessful¹⁴¹. It is not yet possible to search whole genomes in the two orders of extant jawless vertebrates, the lampreys and hagfish; however, several molecular genetic searches have failed to detect the presence of MHC class I or class II molecules in these species. The apparent absence of MHC class I or class II molecules in species that occupy positions below the phylogenetic level of jawed vertebrates³⁸ confounds efforts to assign the ancestral form of the conventional adaptive immune receptors as being either BCR- or TCR-like and might present an intractable problem. Notably, despite the apparent absence of MHC molecules, *Botryllus schlosseri* (a colonial protochordate)¹⁴² and *Hydractinia* species (cnidarians; which are invertebrates)^{143,144} show HISTOCOMPATIBILITY REACTIONS (that is, allotransplantation reactions) through a process known as fusibility, which protects the germline by inhibiting chimerism and/or parasitism¹⁴⁵. Fusibility is determined by a single Mendelian trait and, in *Botryllus schlosseri*¹⁴⁶, is mediated by highly polymorphic cell-surface receptors that contain immunoglobulin-superfamily and epidermal-growth-factor domains that are encoded at a single locus¹⁴⁷. Notably, again, a chimeric structure is involved in an immune-type function.

Box 2**Mechanistic similarities between pathogenicity and protection**

In a co-evolutionary struggle, the host and the pathogen use similar mechanisms to modify selection and outcome. In various pathogens, including bacteria and protozoan parasites, several fundamental mechanisms of DNA change are used during invasion of the host. DNA recombination^{148–151}, gene conversion^{152–154} and gene hypermutation can lead to the evasion of host detection and to the subsequent clonal selection of the pathogenic variant. For example, in trypanosomes, DNA recombination and gene conversion rapidly modify variant surface glycoproteins and allow evasion of host detection^{155,156}. In the parasite that causes malaria, these genetic events, which include gene activation and regulation, are modulated by various complex epigenetic mechanisms^{157,158}.

In the host, analogous processes, including DNA recombination and hypermutation of recognition receptors, counter these pathogenic challenges. For example, genome diversification occurs during meiotic recombination and can be further increased by the creation and maintenance (through sexual outbreeding) of allelic diversity (which is generated by selection and drift) and by the epigenetic control of gene expression. These events help to establish generalized variation and population diversity. The host immune cell, through its receptors, directly influences selection of the host or the pathogen. In vertebrates, somatic modification of the immune-cell receptor repertoire is common and can further improve recognition potential in clonal lineages. The direct interaction between immune cell and pathogen affects the selection of virulence genes and the diversity of pathogen populations. Since the origin of metazoans, a continuous exchange of selective pressures through a pathogen–host–immune-cell loop has driven the simultaneous diversification of cell-surface receptors, using parallel mechanisms of genetic modification, which has helped to drive the lineage-dependent divergences that are seen in the innate and adaptive immune repertoires.



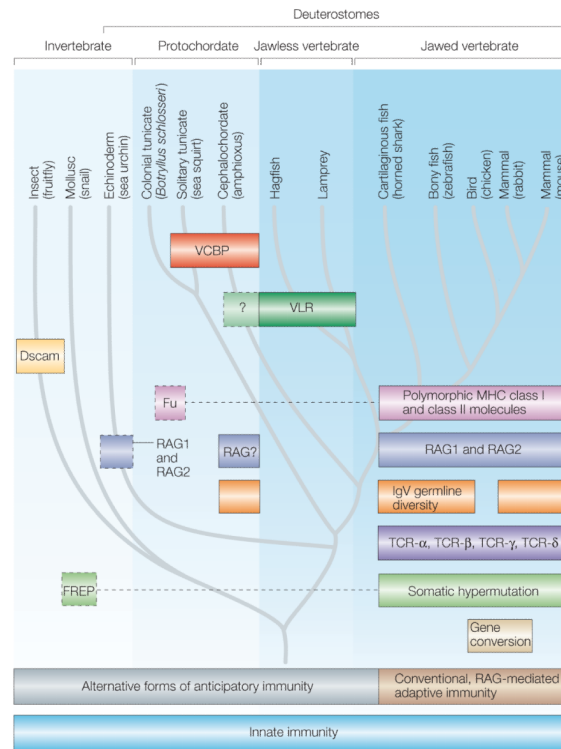


Figure 1. Phylogenetic occurrence of various immune functions, genetic organizations and mechanisms of receptor variation in selected representative species

The selection of individual invertebrate, protochordate, jawless-vertebrate and jawed-vertebrate species is based on emphasizing certain effects and is not intended to be comprehensive. Innate immunity uses various mediators that are not necessarily shared among the main groups that are represented; conventional, recombination-activating gene (RAG)-mediated adaptive immunity is thought to be confined to the jawed vertebrates, and alternative forms of anticipatory immunity are found in species that are below this level of phylogenetic development. Dashed lines are used to emphasize certain general or parallel relationships. Fusibility (*Fu*) genes are unrelated to the genes encoding MHC class I and class II molecules, but they are an alternative histocompatibility locus¹⁴⁶. A *Fu* locus has also been identified in *Hydractinia* species (cnidarians; which are invertebrates), and there might be homologous forms of *Fu* gene products in other species. *RAG1*-like gene sequences are found in amphioxus, which are cephalochordates; *RAG2* sequences have not yet been characterized. Differential RNA splicing creates diversity in insect Down's syndrome cell-adhesion molecule (Dscam), which is associated with bacterial binding and phagocytic function. Hypermutation occurs in the genes encoding fibrinogen-related proteins (FREPs) in a mollusc. Various leucine-rich repeat (LRR)-encoding genes, which are distantly related to the genes encoding the variable lymphocyte receptors (VLRs) of lampreys and hagfish and the Toll-like receptors of vertebrates, can be identified in the amphioxus genome (indicated by light-green shading); however, their relatedness to the genes encoding VLRs and to other LRR-encoding genes has yet to be determined, and the possibility that there might be VLRs in jawed vertebrates awaits further investigation. Throughout the metazoans, a huge number of different receptors and effector molecules are used to effect innate immunity, and these typically vary between the major phylogenetic groups. The background shows a simplified phylogeny of many of the species that feature in current considerations of the evolution of innate and adaptive immunity. IgV,

immunoglobulin variable region; TCR, T-cell receptor; VCBP, variable-region-containing chitin-binding protein.

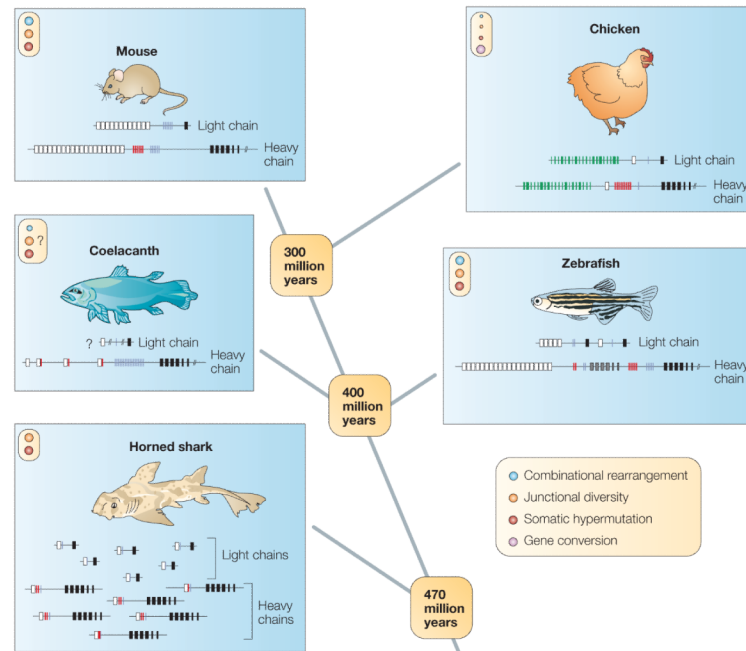


Figure 2. Main differences in the organization of rearranging segmental elements in immunoglobulin heavy and light chains in selected jawed-vertebrate models

Each jawed vertebrate is indicated by its common name: horned shark, zebrafish, coelacanth, chicken and mouse. The selection of the horned shark, chicken and mouse emphasizes the co-evolution of the organization of genes and the main mechanisms of diversification. In the case of mice, only an $Ig\kappa$ light-chain locus is shown; the $Ig\lambda$ light-chain locus has a cluster-type organization. Neither the number nor the relative position that is shown for variable (white), diversity (red), joining (blue) and pseudogene (green; used in gene conversion) elements in heavy and light chains is absolute. For the heavy chain, only the most proximal exons encoding the constant region (black) are indicated, except in the case of zebrafish, for which the exons encoding an additional heavy-chain class (grey) are shown. Additional pseudogenes are present in the continuous variable-element-encoding region in mice and zebrafish and probably also in coelacanths. The organization of genes in the type II light chain¹⁵⁹ in the zebrafish has been inferred by annotation of the zebrafish genome and is not complete as represented; note the presence of two constant regions in the cluster. Light-chain gene organization has not been reported in coelacanths. The relative contributions of various diversifying mechanisms have been approximated in circles: combinatorial rearrangement (blue), junctional diversity (orange); somatic hypermutation (red) and gene conversion (pink). The differences in circle area roughly approximate the relative differences in overall effect and are not quantitative. The relative time of divergence from the lineage that gave rise to mammals is indicated in million-year increments; a ~50-million-year error is assumed¹⁶⁰. This figure is produced from data taken from REFS 29,33.

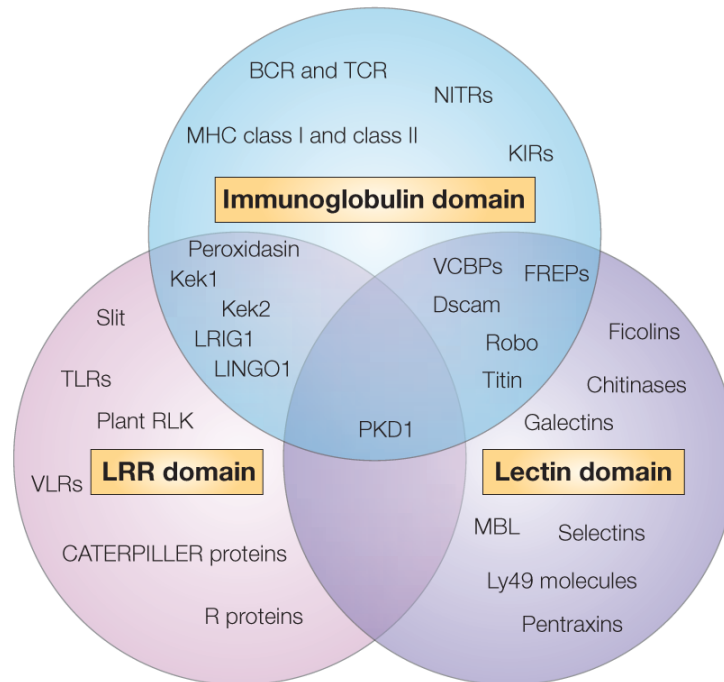
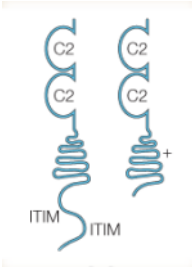
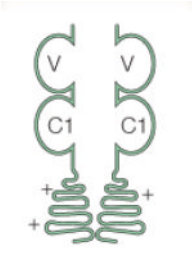
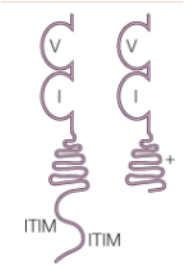


Figure 3. Three main framework domains are incorporated in various proteins, including those that mediate adaptive and innate immunity

The immunoglobulin-domain framework includes a superfamily of homologous types: variable (V), intermediate (I) and constant (C1 and C2). Single or multiple copies of one or more types of immunoglobulin domain are present in different receptors: for example, novel immune-type receptors (NITRs) can have V and I domains. Other proteins have more than one type of framework domain: for example, immunoglobulin V domains and a chitin-binding lectin domain in V-region-containing chitin-binding proteins (VCBPs). Fibrinogen-related proteins (FREPs) and Down's syndrome cell-adhesion molecule (Dscam) also contain immunoglobulin and lectin domains. Proteins that have both lectin and leucine-rich repeat (LRR) domains are rare; only polycystic kidney disease 1 (PKD1) is known to have all three domains. Proteins that share a specific domain type might not be related by common ancestry, so only the domains themselves have a homologous past (as products of recruitment). LRR-containing proteins are a special case. The LRR framework is a small motif (~11 amino-acid residues) that is generally encoded in contiguous copies. The functional site can be a single motif or a combination of motifs. LRR motifs are so divergent that even though the 11-residue-motif structure is highly conserved, together with the resulting solenoid structure of the LRR-containing protein, it is not possible to predict whether any two LRR motifs evolved from a common ancestor or whether they arose through evolutionary convergence. BCR, B-cell receptor; CATERPILLER, caspase-recruitment domain, transcription enhancer, R (purine)-binding, pyrin, lots of LRRs family; ficolin, collagen- and fibrinogen-domain-containing lectin; galectin, galactoside-binding lectin; Kek, kekkon; KIR, killer-cell immunoglobulin-like receptor; LINGO1, LRR- and immunoglobulin-domain-containing, NOGO (neurite-outgrowth inhibitor)-receptor-interacting protein; LRIG1, LRR and immunoglobulin-like domains 1; MBL, mannose-binding lectin; RLK, receptor-like kinase; Robo, roundabout; R proteins, plant disease-resistance proteins; TCR, T-cell receptor; TLR, Toll-like receptor; VLR, variable lymphocyte receptor.

Table 1

Natural-killer-cell, T-cell and related receptors

Feature*	KIR [‡]	TCR [‡]	NITR [‡]
			
Number of IgSF domains	2–3	2	1–2
Vdomains	No	Yes	Yes
Somatic change	No	Yes	No
Activating and inhibitory forms	Yes	No	Yes
Haplotypic variation	Yes	–	Yes
Maximum haplotype number [§]	14	~70 V α , 52 V β	36

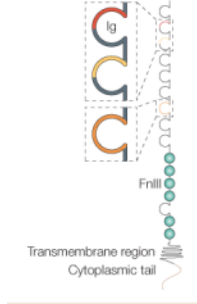




* The various characteristics that are indicated allow comparisons between the three receptor classes and are neither inclusive nor necessarily indicative of functional analogy, but they show that novel immune-type receptors (NITRs) share various characteristics with killer-cell immunoglobulin-like receptors (KIRs) and T-cell receptors (TCRs).

[‡]The KIRs and $\alpha\beta$ -TCR shown are of human origin; the NITRs that are described are from zebrafish. The structures have been drawn to emphasize potential similarities between these receptor types; the domain types — variable (V), intermediate (I) and constant (C1 and C2) — are indicated. Inhibitory motifs (immunoreceptor tyrosine-based inhibitory motifs, ITIMs) and positively charged residues in the transmembrane regions of KIRs and NITRs are shown; the pairings that are depicted are not intended to imply that KIRs and NITRs function as heterodimers. Positively charged residues in the TCR, which does function as a heterodimer, are also indicated. The structures are not drawn to scale.

[§]Haplotype comparisons reflect the highest allele content for various human populations and a laboratory line of zebrafish. IgSF, immunoglobulin superfamily; V α , variable region of α -chain; V β , variable region of β -chain.

Table 2

Organization of some alternative immune-type receptors

Feature*	Dscam [‡]	VLR [‡]	VCBP [‡]	FREP [‡]	IgL [‡]
					
IgSF domains	Yes	No	Yes	Yes	Yes
V domains	No	No	Yes	No	Yes
Leucine-rich repeats	No	Yes	No	No	No
Somatic reorganization	No	Yes	No	No	Yes
Junctional diversity	No	No	No	No	Yes
Complex RNA processing	Yes	No	No	No	No
Somatic hypermutation	No	No	No	Yes	Yes
Extensive population polymorphism	No	No	Yes	Yes	No
Regionalized hypervariation	Yes	No	Yes	No	Yes

*The various properties and characteristics that are indicated focus on those that best reflect the similarities and differences between the various receptors shown.

[‡]Schematic organization is shown for immune-type receptors that have various characteristics of innate and adaptive immune receptors and are found in traditional model systems: from jawless vertebrates, variable lymphocyte receptors (VLRs); from invertebrates, fibrinogen-related proteins (FREPs) and Down's syndrome cell-adhesion molecule (Dscam); from protochordates, variable (V)-region-containing chitin-binding proteins (VCBPs); and from jawed vertebrates, immunoglobulin light chains (IgLs). Coloured segments in the VLR indicate the regions in which extensive somatic variation occurs in leucine-rich repeats (LRRs). The relative sites of sequence variation in the various receptors are indicated by coloured symbols, irrespective of whether the variation occurs in a single gene (as is the case for IgL). Variation arises through several mechanisms: complex RNA processing, for Dscam; recombination of LRR cassettes, for VLRs; allelic complexity and population polymorphism, for VCBPs; and somatic mutations in multiple germline genes, for FREPs. Variation in Dscam is localized to only three immunoglobulin domains. Gene conversion (not shown) is a diversification mechanism for avian IgL and possibly for VLRs, although the latter has not been established. C1 domain, type 1 immunoglobulin constant-region domain; CBD, chitin-binding domain; FBG domain, fibrinogen-like domain; FcIII domain, fibronectin type III domain; GPI, glycosylphosphatidylinositol; IgSF, immunoglobulin superfamily.