ORIGINAL PAPER

Evolution of the B7 family: co-evolution of B7H6 and NKp30, identification of a new B7 family member, B7H7, and of B7's historical relationship with the MHC

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Abstract The B7 family of genes is essential in the regulation of the adaptive immune system. Most B7 family members contain both variable (V)- and constant (C)-type domains of the immunoglobulin superfamily (IgSF). Through in silico screening of the Xenopus genome and subsequent phylogenetic analysis, we found novel genes belonging to the B7 family, one of which is the recently discovered B7H6. Humans and rats have a single B7H6 gene; however, many B7H6 genes were detected in a single large cluster in the *Xenopus* genome. The *B7H6* expression patterns also varied in a species-specific manner. Human B7H6 binds to the activating natural killer receptor, NKp30. While the NKp30 gene is single-copy and maps to the MHC in most vertebrates, many Xenopus NKp30 genes were found in a cluster on a separate chromosome that does not harbor the MHC. Indeed, in all species so far analyzed from sharks to mammals, the number of NKp30 and B7H6 genes correlates well, suggestive of receptor-ligand co-evolution.

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Department of Veterinary Pathobiology, College of Veterinary Medicine and Biomedical Sciences, Texas A&M University, College Station, TX 77843, USA Furthermore, we identified a *Xenopus*-specific *B7* homolog (*B7HXen*) and revealed its close linkage to *B2M*, which we have demonstrated previously to have been originally encoded in the MHC. Thus, our study provides further proof that the B7 precursor was included in the proto MHC. Additionally, the comparative analysis revealed a new B7 family member, B7H7, which was previously designated in the literature as an unknown gene, *HHLA2*.

Keywords B7 family \cdot MHC \cdot Evolution \cdot Natural killer receptors \cdot Genetic linkage \cdot *Xenopus*

Introduction

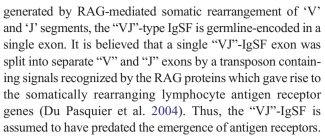
B7 ligands are expressed on the cell surface of many different cell types including APCs and their interaction with receptor molecules on T cells provide activating and/or inhibitory signals that regulate T cell activation and tolerance (Collins et al. 2005). Some inhibitory B7 ligands are also expressed on tumor cells, resulting in suppression of immune responses (Keir et al. 2008; Zou and Chen 2008). Therefore, stimulating or attenuating the interactions of B7 ligands and their receptors holds therapeutic potential for autoimmune diseases and cancer (Callahan et al. 2010; Dinesh et al. 2010; Loos et al. 2010). Most known B7 members contain so-called Ig superfamily (IgSF) domains (Williams and Barclay 1988), with an N-terminal variable (V)-type domain, followed by one constant (C)-type domain and a transmembrane (TM) region. V-type IgSF domains are usually responsible for interacting with their ligands. C-type IgSF domains generally fall into two major categories, C1 and C2, based on their beta strand and loop compositions (Williams and Barclay 1988). The C2-type is more primitive because it is found in certain immune molecules such as NK



receptors and FcRs in vertebrates, but it is most commonly found in adhesion molecules in both invertebrates and vertebrates (Du Pasquier et al. 2004). In contrast, the C1 type is found primarily in molecules involved in adaptive immunity, and the C domains found in B7 molecules are mostly of this unusual type. Many B7 family members have been identified from jawed vertebrate (gnathostome) classes (Hansen et al. 2009; Zhang et al. 2009) and phylogenetic analyses have shown their V domains to be related to tapasin (TAPBP), poliovirus receptor (PVR), and other virus receptor V domains (e.g., Coxsackievirus and Adenovirus receptors), and distantly related to butyrophilins (BTN) and myelin oligodendrocyte glycoproteins (MOG) (Fahrer et al. 2001; Henry et al. 1999; Linsley et al. 1994).

Xenopus species shared a common ancestor with humans ~350 million years ago, and these amphibians are "connecting" organisms linking fish to higher vertebrate taxa (Hellsten et al. 2010; Vogel 1999). Compared to the bony fish model, in which the genome has been disrupted presumably due to rapid expansion and contraction of species and a class-specific genome-wide duplication (Postlethwait et al. 2004), the Xenopus genome is much more stable, and some genes display unexpected syntenic relationships that are primordial (Ohta et al. 2006). Such genetic stability distinguishes Xenopus as a model taxon for comparative analysis. Thus, by examining syntenic relationships in Xenopus, in concert with overall comparative analyses in all chordates, we hope to shed light on the natural history of the vertebrate genome, especially regarding immunity.

While examining the Xenopus genome for MHC-related genes, we found two polygenic clusters of highly divergent genes that contain both V- and C1-type IgSF domains. Phylogenetic analysis showed that the V-IgSF domain clustered with the known B7 family members and subsequently we found orthologs from other vertebrate species, including human. Indeed, while we were preparing this paper, a human gene was reported as the newest B7 family member, B7H6 (Brandt et al. 2009). B7H6 binds to the activating natural killer receptor (NKR), NKp30, through its V domain (Brandt et al. 2009). B7H6 is upregulated under tumor transformation or stress conditions and NK cells eliminate such cells either directly via cytotoxicity or indirectly by cytokine secretion (Baratin and Vivier 2010). NKp30 is encoded in the MHC in mammals and possesses a single V-IgSF domain followed by a TM region (Pende et al. 1999). The cytoplasmic region of NKp30 interacts with the ITAM-containing adaptor molecule, CD3ζ, resulting in activation signals to NK cells (Pende et al. 1999). Interestingly, the NKp30 V domain is a unique type, the so-called "VJ"-domain, which resembles the precursor of antigen receptors that has been found in other genes involved in immunity and cellular interactions (Bartl et al. 1994; Du Pasquier 2002; Thompson 1995). Unlike the V domains of the antigen receptors in which diversity is



In this paper, we conducted a genomic analysis of *Xenopus B7H6* and *NKp30* loci. We found a positive correlation between the number of loci for these receptors and ligands throughout vertebrate evolution. We also uncovered another (and probably the last) B7 family member, which we designate B7H7. We further performed a phylogenetic analysis of all known B7 family members in all gnathostomes. Lastly, our analysis provides unequivocal evidence that B7 family members were originally encoded in the MHC.

Materials and methods

Data-mining the EST and genomic databases

Initially, the new Xenopus B7 family genes were found serendipitously with tBLASTn searches using IgSF domains of chicken MHC class II DM. Both the Xenopus tropicalis and Xenopus laevis EST databases and the X. tropicalis genomic assembly version 4.1 at the Department of Energy Joint Genome Institute (JGI: www.genome.psi/jgi.doe.gov/Xenopus) were used. All "hits" were further BLASTx searched against the eukaryotes protein databases to confirm their novelty. We then searched for all Xenopus B7 families by using B7 IgSF domains from other species and confirmed the orthology by BLAST and later phylogenetic trees. When there were Xenopus EST sequences available, genomic scaffolds were searched with BLASTn (using X. tropicalis) or tBLASTx (using X. laevis). Analysis of genomic synteny was obtained from either Gene Page at the NCBI web site or Map Viewer, with further manual annotation when necessary (as well as classical segregation analyses by Southern blotting, see below). Xenopus NKp30 genes were identified by using human NKp30 as bait and confirmed as detailed above. To search for NKp30 in shark databases, we used primarily *Xenopus NKp30* as bait and again confirmed as described above. Sequences were searched in the NCBI databases and "Elephant Shark Genome Project" web site (http://esharkgenome.imcb.a-star.edu.sg/). Transmembrane prediction was made using TMHMM server v2.0 (http:// www.cbs.dtu.dk/services/TMHMM/).

Phylogenetic tree analysis

The V domains of the deduced amino acid sequences of B7 family members were aligned using ClustalX and neighbor-



joining bootstrapping trees (500 trials) (Saitou and Nei 1987) were made and viewed in the TreeView 1.6.6. program (Page 1996). For phylogenetic tree analysis of NKp30, amino acid alignments of variable domains were initially made in Bioedit with ClustalW employing gap opening penalties of 10 and gap extension penalties of 0.1 for pairwise alignments then 0.2 for multiple alignments and the protein weighting matrix of Gonnett or Blossum (Hall 1999; Tamura et al. 2007). These alignments were then modified by hand and converted back to nucleotides for computation of distances and trees. MEGA was then used to infer the phylogenetic relationships. A neighbor-joining tree was built using distances computed with the maximum composite likelihood method. All three positions of codons were included in the analysis and 525 nucleotide columns were included. The maximum likelihood tree was computed with the method of Tamura et al. (2007) and had a log likelihood of -5,921.4319. The maximum likelihood tree employed 203 positions as all columns with gaps or missing data were excluded. The minimum evolution tree resulted in a sum branch length of 9.74464 and used all 525 positions. These NKp30 trees were bootstrapped 1,000 times.

Southern blotting

Genomic DNA (10 µg) isolated from the erythrocytes of an MHC-defined *X. laevis* family (Ohta et al. 1999) and *X. tropicalis* was digested with 80 units of restriction endonucleases overnight, followed by further digestion with 40 units of enzymes overnight. DNA was then separated on 0.8 % agarose gels and blotted onto nitrocellurose membranes by capillary transfer (Whatman). Membranes were then hybridized with a radiolabeled cDNA probe under high stringency conditions (50 % formamide) and washed under stringent conditions (Bartl et al. 1997). The same membranes were often re-used for different probes after the radioactivity decayed.

Fluorescence in situ hybridization on chromosomes

Pre-Sertoli cells as a source of metaphase spreads were obtained from testes of *X. tropicalis* juveniles. Hypotonic shock was done by 40 mM KCl followed by three washes with fixative (methanol:acetic acid, 3:1). The cell suspension was then dropped onto glass slides with subsequent immersion in 50 % acetic acid for 2 s. The ~1 kb cDNA probes were labeled by Dig-11-dUTP using the Decalabel Labelling Kit (Fermentas). Simultaneous hybridization of 100–150 ng of probe with chromosomes was performed using simple hybridization mixture of 50 % formamide in 2× SSC, in a humidifying chamber overnight at 37 °C. After several high stringency washes, slides were blocked and incubated with antidigoxigenin-POD antibody. Signal

visualization was done using tyramide coupled with tetramethylrhodamine (TSA-TMR kit-Perkin Elmer) (Krylov et al. 2007). Nuclei and chromosomes were counterstained with DAPI (shown in blue).

Expression

Gene expression was examined by northern blotting using 10–15 μg total RNA from various tissues. We extracted total RNA from various tissues of a Lewis-strain rat and *X. laevis*. Human total RNA from 18 different tissues was purchased from Invitrogen (Carlsbad, CA). The human and rat probes were made by subcloning the RT-PCR fragment amplified using primers encompassing the two IgSF domains (i.e., inter-exons) based on the GenBank sequence entries from liver, spleen, and testis RNA. The DNA fragments of expected size (~550 bp) were amplified from all the tissues in both human and rat.

Results and discussion

Identification of Xenopus B7H6 genes

While searching for Xenopus DM genes (an MHC class IIrelated gene) by BLAST using the immunoglobulin superfamily (IgSF) domain of DM genes from other species (e.g., chicken), we found many sequences containing C1-type IgSF domains in the databases. Subsequently, we performed BLASTp and BLASTx searches for each sequence and discovered that one partial EST (BX732750; X. tropicalis) sequence encodes a novel gene. Using this partial sequence, we found two closely related full-length X. laevis sequences (AAH84632 and AAH82955) and an AAH84632 ortholog (AAI66200) from the *Xenopus* databases (Fig. 1a). The deduced amino acid (aa) sequences of all clones had two extracellular domains, an N-terminal variable (V) and C-terminal C1-type, based on characteristic aa residues in each type of domain (shaded in grey in Fig. 1a), followed by a predicted type I transmembrane (TM) region (underlined in Fig. 1a). Additionally, there were often long cytoplasmic tails containing 4 to 5 IgSF domains (e.g., AAH84632, AAI66200), (Online Resource 1a), whereas there were no obvious cytoplasmic IgSF domains in AAH82955. BLASTp searches using V-IgSF domains showed that the Xenopus clones are most similar to the V domains of some genes of unknown function from various species (see below for further details) followed by the TAP-binding protein-like (TAPBP-R) with an E-value $\sim e^{-12}$, and the extracellular C1-IgSF was most similar to TAPBP and MHC class II C1-IgSF domains with E-values ~e⁻¹⁰. The presence of C1-IgSF domains that are distantly related to the C1-IgSF domains in various other immune genes (e.g., TCR, Igs, MHC I,



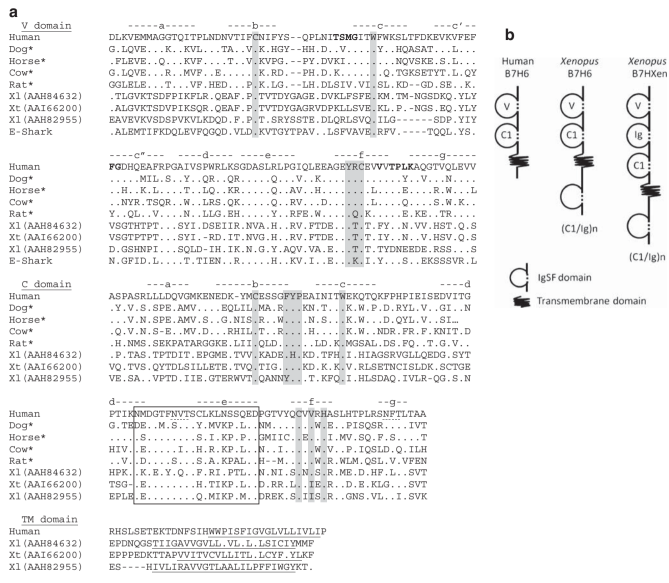


Fig. 1 a Alignment of B7H6 from various vertebrate species. Gaps and identical aa residues are indicated with *dashes* and *dots*, respectively. Canonical aa residues characteristic of IgSF domains are shaded in *gray*, potential glycosylation sites are marked with *dotted underlines*, and predicted TM regions are underlined. IgSF strand designation is shown above sequences and potential glycosylation sites are marked with *dotted underlines*. Only external IgSF and predicted TM (when available) domains are shown; the cytoplasmic regions including the internal Ig domains in *Xenopus* sequences are shown in Online Resource 1. *Bold letters* indicate the interaction sites with NKp30 based on the crystal structure (Li et al. 2011). *Asterisk* indicates

predicted genes. Conserved aa residues in the D-loop to E-strand region of the C1 domain are *boxed*. Accession numbers are listed in Fig. 2 legend. This figure includes sequences from representative vertebrates in different classes. **b** Domain structures of B7H6 and B7HXen. Human B7H6 represents all species except *Xenopus*. Number of cytoplasmic Ig domains varies depending on the sequences: *n*=5 for AAH84632 and AAI66200; *n*=0 for AAH82955 and AAI61104; *n*=4 for AAH85212; various numbers were predicted in gene models from the *X. tropicalis* genome (details in Online resources 1a, 1b, and 2)

MHC II, TAP-binding protein (TAPBP)) seems to be unique to these molecules. Interestingly, some of the cytoplasmic IgSF domains were C1-type and matched to C1-IgSF domains of MHC class I and IgH/L with low but significant similarities (~e⁻⁷ and ~e⁻⁴, respectively), while other non-C1-type domains seemed to be degenerate (e.g., domain-5 of AAH84632 "lost" the N-terminal-half (Online Resource 1a)) and did not significantly match other sequences. Although the

cytoplasmic domains may contribute to a unique signaling mechanism, and/or interaction with adaptor molecules that contribute to downstream signaling events; in this report, we focus only on the extracellular domains where the receptor—ligand interaction occurs in all B7 family members so far examined.

To examine whether such IgSF molecules were *Xenopus*-specific, we examined the genes with unknown function that



matched with the "best hits" (E-values < e⁻⁸) during the BLAST searches. Accession numbers of these genes were CAD97811 (cDNA from human uterus; also previously named AC406 (Du Pasquier 2000)), XP 001076811 (predicted gene; rat), XP 601599 (predicted gene; cow), XP 542518 (predicted gene; dog), XP 001501683 (predicted gene; horse), and CAH89395 (cDNA from orangutan kidney). Although some sequences were only predicted gene models and may not be full-length (marked as asterisk in Fig. 1a), all of them contained the V-C1 type domain architecture (Fig. 1b). The C1 domains from these genes show low but significant similarity to MHC class II IgSF domains. As mentioned in the "Introduction" section, while we were characterizing these molecules, the human ortholog was reported as the newest B7 molecule, B7H6, a ligand for the NKR NKp30 (Brandt et al. 2009).

Since the membrane distal V domains are involved in receptor-ligand interactions and contain conserved characteristic features of each gene family, we constructed phylogenetic trees using V-IgSF domains. Preliminary phylogenetic tree analysis with various molecules, including antigen receptors, NKRs, signal-regulatory protein (SIRPs) and poliovirus receptor (PVRs), revealed clustering of B7H6 to the other B7 family members (data now shown). Thus, we used only the B7 family members to identify the orthologous relationships (Fig. 2). Xenopus IgSF genes clustered tightly with B7H6 from other species, and based on the BLAST results and phylogenetic analysis, we concluded that the Xenopus IgSF gene is a B7H6 ortholog. Note that the cytoplasmic IgSF domains seen in *Xenopus* were not detected in other species (Fig. 1b). The deduced aa alignment shows conservation among mammalian species (Fig. 1a), and the variability is more apparent in the N-terminal halves of both the V- and C1-IgSF domains. There is a surprisingly high conservation of amino acid residues in the D-loop to E-strand region of the C1-IgSF domain (boxed in Fig. 1a), which may be used either for homodimerization or interaction with other molecules on the same cell. The TM regions are well predicted in all three *Xenopus* sequences (underlined in Fig. 1a). There is no GAG polyprotein homologous regions in the 3' untranslated region as was found in the original human sequence, thus this viral insertion seems to be primate-specific (Kaifu et al. 2011).

B7H6 expression

A recent report showed that the human *B7H6* was not detected in normal human tissues but was expressed by human tumor cell serving as a damage-associated molecular pattern to trigger innate immunity (Baratin and Vivier 2010; Brandt et al. 2009). We examine whether this expression pattern was evolutionary conserved. High *B7H6* expression in an unimmunized rat was detected in lymph

node, muscle, spleen, and testis, and lower expression was found in mucosal tissues (colon, ileum, jejunum, and lung) (Fig. 3a). Thus, there is a different expression pattern in human and rat.

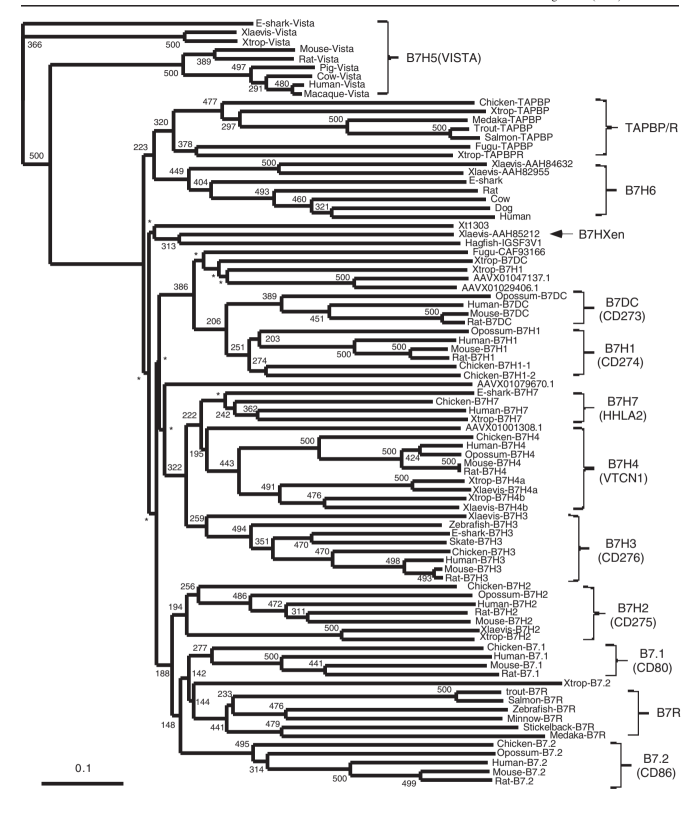
Xenopus B7H6 expression was also studied in unimmunized animals. There were at least five distinct transcripts showing tissue-specific expression patterns in blood, colon, intestine, and spleen, and lower amounts in skin and other tissues (Fig. 3b). Multiple transcripts were expected since we identified many different-sized EST sequences from various tissues and developmental stages (see below; and Online Resource 2). Thus, the diverse bands likely reflect expression of different genes and/or differentially spliced forms. Since we used a probe derived from the BX732750 sequence that only contains two cytoplasmic IgSF domains (Online Resource 1a), we repeated the northern blotting with the V-IgSF domain of AAH84632, which under low stringency conditions should recognize all frog B7H6 members (Online Resource 1b), and we obtained identical results (data not shown). Collectively, the differential expression pattern in various species implies that B7H6 can acquire new functions in different vertebrate taxa.

Genomic analysis of Xenopus B7H6 gene

To obtain B7H6 genomic information, we searched the X. tropicalis genomic scaffolds (http://genome.jgi-psf.org/ Xentr4/Xentr4.home.html) by BLASTn and tBLASTn and found 12-13 predicted gene models in two scaffolds (707 and 1,420) (Online Resource 2). Most of them contain a domain structure similar to AAH824632 and AAI66200 (Fig. 1a, b; Online Resource 1a) with external V-C1, TM, and cytoplasmic C1-IgSF and "Ig-like" domains. Fiftyseven unique IgSF domains were annotated, each of which being encoded by a single exon. We found that some C1 or "Ig-like" domains are more similar to each other and they were often found in the same order like cassettes in multiple gene models (data not shown); thus, these IgSF domains were likely generated by duplications, further suggesting that the expansion of the Xenopus B7H6 genes was driven by cis-duplications and subsequent exon shuffling.

B7H6 genes from all mammalian species are either single-copy or completely absent (Figs. 1, 2, and Table 1), and therefore the expansion of B7H6 genes seems to be species-specific. The orthologous genes were lost in mouse (Brandt et al. 2009), chicken and opossum, and are not present in fish (Table 1). The species-specific expression pattern, in combination with expansion of genes in Xenopus and loss of the gene in some species, suggest that B7H6 evolves rapidly and might be co-evolving with its receptor genes. The deduced as sequences for all V domains contain the conserved cysteine (C), tryptophan (W), but no GXG





motif in the G-strand (i.e., not a "VJ"-type IgSF domain). The alignment of the *Xenopus* V-IgSF domains showed broad variability (57–93 %) throughout the entire domain (Online Resource 1b).

Identification of a Xenopus-specific B7H6 subfamily

During the searches, we found additional full-length sequences (AAH85212 (X. laevis) and AAI61104 (X. tropicalis))



Fig. 2 Phylogenetic tree analysis of all members in the B7 family. Deduced as sequences of V domains are used for this analysis. The bootstrapping tree was constructed using the Neighbor-Joining method with 500 runs (Saitou and Nei 1987). Bootstrapping values are shown at the branch, asterisk indicates less than 100 values. Our analysis consistently showed VISTA (gi24; B7H5) (Wang et al. 2011) to be the most divergent member of B7 family, thus we used it as outgroup. The scale correlates with the divergence time. Accession numbers used for this analysis are: **B7.1**—human (NP 005182), mouse (CAJ18316), Rat (AAC02262), chicken (CAJ18316); **B7.2**—human (CAG46642), mouse (P42082), rat (NP 064466), opossum (XP 001371519), chicken (XP 001371519), X. tropicalis (protein ID: 510857 at JGI genome site); B7R—trout (ACH58053), zebrafish (CN017839), minnow (DT293786), stickleback (DN656970), salmon (DW580717), medaka (AM306315); B7H1—human (NP 054862), mouse (EDL41681), rat (XP 001079572), opossum (XP 001371389), chicken (XP 424811, CF251191), fugu (CAF93166), X. tropicalis (protein ID: 16544 at JGI genome site); B7DC- human (AAP13471), mouse (Q9WUL5), rat (EDM13096), opossum (XP 001371408), fugu (CAF93166), X. tropicalis (Protein ID:162957 at JGI genome site), E-shark (AAVX14147467.1, AAVX01029406.1); B7H2—human (BAA31628), mouse (Q9JHJ8), rat (XP 574731), opossum (XP 001377700), chicken (CAA70058), X. laevis (translated from BX848980), X. tropicalis (Protein ID: 170158 at JGI genome site); B7H3—human (AAK15438), mouse (EDL25954), rat (Q7TPB4), opossum (XP_413702), chicken (Q68EV1), X. laevis (XP_700973), zebrafish (XP 001341518), skate (translated from DT378804), Eshark (AAVX01064573.1); B7H4—human (AAP37283), mouse (AAP37284), rat (AAH95842), opossum (XP 001365506), chicken (XP 416546), X. tropicalis (a: CAJ83577; b: protein ID:396923 at JGI genome site), X. laevis (a: AAH41253; b: AAH44000), E-shark (AAVX01001308.1); B7H5—human (AAH20568), Rhesus Monkey (NP 001177807), mouse (Q9D659), rat (AAH98723), pig (XP 001928572), cow (AAI23905), X. tropicalis (AAH88869), X. laevis (AAH73319), E-shark (AAVX01038049.1); B7H6—human (CAH18335), cow (NP 001193721), rat (XP 001076811), dog (XP 542518), X. tropicalis (AAI66200), X. laevis (AAH84632), Eshark (AAVX01166257.1); **B7HXen**—X. tropicalis (AAI61104), X. laevis (AAH85212), B7H7—human (Q9UM44), X. tropicalis (AAH96014), X. laevis (ACH85300), E-shark (AAVX01019552.1); TAPBP—chicken (CAA06327), X. tropicalis (XP 002938728), medaka (BAB83851), trout (AAZ66041), salmon (ABO13869), fugu (CAF97838); TAPBPR-X. tropicalis (XP 002941962); Hagfish IGSF3 (BAE93396); X tropicalis unknown B7 family in the scaffold-1303 (Protein ID: 461476). Opossum B7.1 and chicken B7H7 were retrieved from the genomic sequences, NW 001581960 and NW 001471429, respectively. BLAST searches with the V-IgSF of B7H5 (aliases; Gi24, VISTA, C10orf54) (Wang et al. 2011) matched to B7H4 with an E-value ~e⁻¹⁰, but consistently separated from other members of the B7 family with long branch lengths and therefore was used to root the tree. When orthology was not strong, E-shark sequences are shown with accession numbers starting with AAVX. All Eshark sequences are deduced from the genomic scaffolds

with low but significant identity to *Xenopus* B7H6. They contain multiple C1-IgSF domains as well as a distinctive V-IgSF domain (Online Resource 1c). The deduced as sequences revealed three extracellular IgSF domains (V-Ig-C1), followed by a TM region, and either a long cytoplasmic tail with four "Ig-like" domains (AAH85212) or no domains (AAI61104) (Fig. 1b). BLASTp searches revealed that the V-IgSF domains were most similar to the V

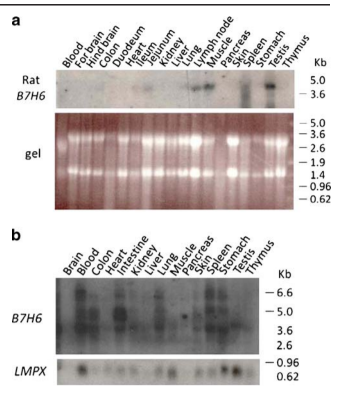


Fig. 3 Tissue expression pattern of rat (a) and *Xenopus* (b) *B7H6*. Northern blotting was done using total RNA from various tissues from a *Lewis* rat and adult *X. laevis*. Ubiquitously expressed *LMPX* was used as a control for *Xenopus*, and the gel image is shown for rat. Size markers are indicated at *right*. Extracellular IgSF domains were used as probes

domains of B7H1 and TAPBP-R (Du Pasquier 2000) with an E-value ~e⁻⁸. Like the results with B7H6, extracellular Ig-like and C1-IgSF domains of this related family were most similar to IgL chain and MHC class II (E-value ~e⁻⁵), and the four cytoplasmic Ig-like domains of AAH85212 matched weakly to IgH, IgL, and MHC (E-values ~e⁻⁷). Exhaustive BLAST searches did not uncover orthologs from other species, suggesting that these genes are *Xenopus*-specific B7 homologues, and thus we designate them as *B7HXen*.

Phylogenetic tree analysis did not show strong orthology to other B7 family members with all low bootstrap values (arrow in Fig. 2). Interestingly, B7HXen clustered with the membrane distal V-IgSF domain of hagfish IgSF3 (Haruta et al. 2006). The results were consistent between several sets of data and numerous analyses using different parameters (data not shown), and thus it is possible that B7HXen might be evolutionarily older than most B7 family members. Unfortunately, related genes were not found in the existing databases of lower deuterostomes such as amphioxus, sea squirt, or sea urchin so the hypothesis will be difficult to verify.

We found 11 *X. tropicalis* B7HXen domain-containing scaffolds (scaffolds-1119, 867 etc.) containing a total of 31 predicted gene models (Online Resource 2), of which 13 V-IgSF domains were found at the N-termini of the gene



Table 1 Presence or absence of B7 family genes in different vertebrates

	Human	Mouse	Rat	Opossum	Chicken	Xenopus	Bony fish	Shark	Domain architecture
B7.1	V	V	√		√		B7R	?	VC-TM
B7.2	\checkmark	$\sqrt{}$	\checkmark	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$?	VC-TM
B7H1	\checkmark	$\sqrt{}$	\checkmark	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	VC-TM
B7DC	\checkmark	$\sqrt{}$	\checkmark	$\sqrt{}$		$\sqrt{}$?	VC-TM
B7H2	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$?	VC-TM
B7H3	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	\checkmark	VC-TM
									VCVC-TM
B7H4	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	\checkmark	\checkmark	VjC-GPI
B7H5	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	\checkmark	V-TM
B7H6	$\sqrt{}$		$\sqrt{}$?	?	$\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{$?	$\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{$	VC-TM
B7H7 (HHLA2)	$\sqrt{}$	Pseudo	Pseudo	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	$\sqrt{}$	VIgIg-TM
NKp30	\checkmark	pseudo	\checkmark	?	?	$\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{$?	$\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{\sqrt{$	Vj-TM

 $\sqrt{\cdot}$ single copy; $\sqrt{\cdot}$ 2 copies; $\sqrt{\cdot}$ $\sqrt{\cdot}$ multigene family; — absence of gene

Vj: V-IgSF domain with the presence of the "GXG" motif in the G strand. Ig: IgSF domain

models. Most gene models had a similar domain structure as AAH85212 (V-Ig-C1-TM) with zero to >10 cytoplasmic Ig domains (Fig. 1b, Online Resource 1c, and 2). Like B7H6, the deduced as sequences for all V domains contain the canonical conserved as but no G-strand GXG motif. Similar to B7H6, alignment of V-IgSF domains from the gene models showed high variability (Online Resource 1d) (some sequences shared only 36 % identity) distributed equally throughout the domain.

NKp30

As mentioned above, human B7H6 binds to the activating NKR, NKp30 (Brandt et al. 2009), and thus we examined the phylogenetic representation of NKp30 in the vertebrate lineage. It has been previously reported that NKp30 genes are not present outside of mammals (Kaifu et al. 2011), and furthermore that NKRs evolve rapidly, making it difficult to find orthologues among divergent taxa (Yoder and Litman 2011). Therefore, we were surprised that our BLAST searches using human NKp30 identified NKp30-like genes in the Xenopus databases (Online Resource 3). There were a total of 17 predicted gene models in six scaffolds in the X. tropicalis genome (88, 215, 3675, 3771, 15172, 18746), with 12 models clustered on scaffold-215. There are three gene models that contain two V-IgSF domains in scaffold-215; however, since NKp30 in all other vertebrates has only one V-IgSF domain and all Xenopus EST sequences only contain a single domain, most likely the models either identify pseudogenes or assembly artifacts owing to the difficulty in identifying the short leader and transmembrane exons. The number of gene models in *X tropicalis* (17–20) corresponds well with the number of bands detected by Southern blotting (~20) using a single-exon probe (Fig. 4a). The *Xenopus NKp30-like* genes are also quite polymorphic, showing different restriction fragment length polymorphism (RFLP) patterns in the 16 individuals tested (four are shown in Fig. 4a).

Like B7H6, many NKp30-like EST sequences were found in databases from various tissue sources (Online Resource 3). Based upon sequence similarity, X. tropicalis NKp30-like genes were categorized into four types: (1) BX-type (only one gene model 215-12 matches to EST sequence BX773925); (2) New (or 88-2)-type (gene model 88-2 has no matching EST sequence); (3) ES-type (four gene models (215-2, 88-1, 15172-1, 3771-1) matching EST sequence ES685492 and many other ESTs); and (4) DT-type (11 gene models (215-1, 3~11, 3675) matching the EST sequences DT447366 and EL837335) (see details in Online Resource 3). The DT-type is most abundant in the genome, but based on the relative levels of EST clones the ES-type may be expressed at the highest levels. The aa sequences of individual clones are >90 % identical within each type, but only 40-60 % identical between different types (Fig. 4b). Although, we do not know the function at this time, the phylogenetic analysis using V-IgSF domains from many different molecules supports the true orthology of Xenopus NKp30 to NKp30 of mammalian species (Fig. 4c, Online Resources 4). All four types of Xenopus NKp30 contain the "VJ" IgSF type "GXG" motif. Among the different types, there may be certain aa changes responsible for interaction with B7H6 based upon the human data (bold letters in Fig. 4b (Li et al. 2011)). Since Xenopus B7H6 and NKp30 are both polygenic and contain divergent V-IgSF domains, there may be a preference for binding between particular receptors and ligands, i.e., certain NKp30



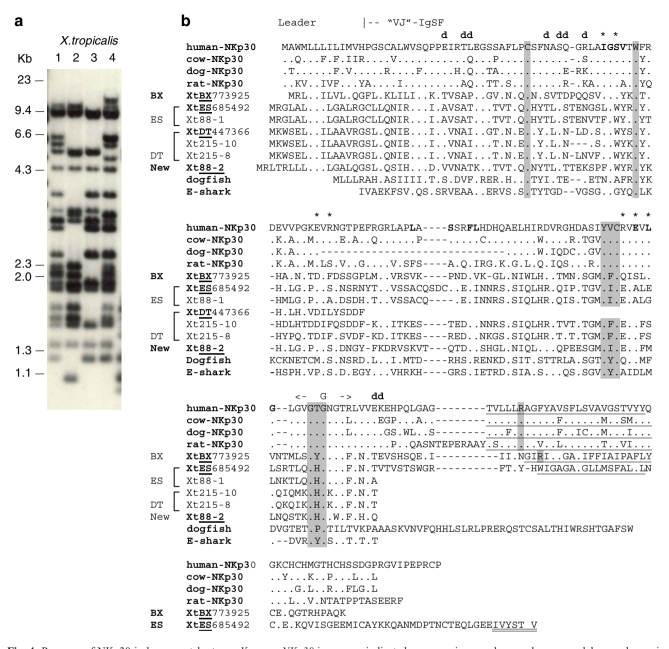


Fig. 4 Presence of NKp30 in lower vertebrates. a Xenopus NKp30 is polygenic and polymorphic. Southern blotting with a single V-IgSF exon probe shows many genes in the X. tropicalis genome. Four different unrelated individual X. tropicalis are shown, revealing RFLP-based polymorphism. Genomic DNA was digested with HindIII. Size markers are shown at left. b Deduced aa alignment of NKp30 from different vertebrate species. Canonical aa residues characteristics for IgSF domains and the "GXG" motif are shaded in gray and predicted TM regions are underlined. The potential dimerization interface, and residues that disrupt interaction with B7H6 when mutated, are noted as d and asterisk above the sequence, respectively (Joyce et al. 2011). Xenopus sequences retrieved from the genomic scaffolds are terminated after the V-IgSF exon, thus the TM and cytoplasmic regions are not known. Sequences belonging to the four types shown as BX, ES, DT, and New (88-2) on the left side of the alignment and representative sequences from gene models are shown under corresponding EST sequences and bundled with parentheses. Xenopus EST sequences

are indicated as accession numbers and gene models are shown in Online Resource 3. A potential ITIM motif in XtES685492 is indicated by double underline and positively charged as residues in TM regions are shaded in gray. Bold letters indicate the interaction sites with B7H6 based on the crystal structure (Li et al. 2011). Accession numbers used for this analysis are: NP_001138939 (human), XP_848928.1 (dog), NP 001035614.1 (cow), NP 861543.1 (rat), ES788778 (spiny dogshark; S. acanthias), and AAVX01127481 (elephant (E-) shark; Callorhinchus milii). c Neighbor joining phylogenetic tree analysis of NKp30. Nucleotide sequences of V domains are used for this analysis. Alternative methods yielding similar results are shown in Online Resource 4. Additional sequences encoding variable domains of lymphocyte antigen receptors were included as an outgroup: IgL κ from shark (L16765.1), frog (DT059572.1), and human (HE584554.1), TCR α from shark (FJ513688.1), and human (AY232282.1), TCRβ from shark (FJ513753.1), frog (NM 001091843.1), and human (L34734.1), and IgM from shark (M92851.1), frog (X14919.1), and human (M87789.1)



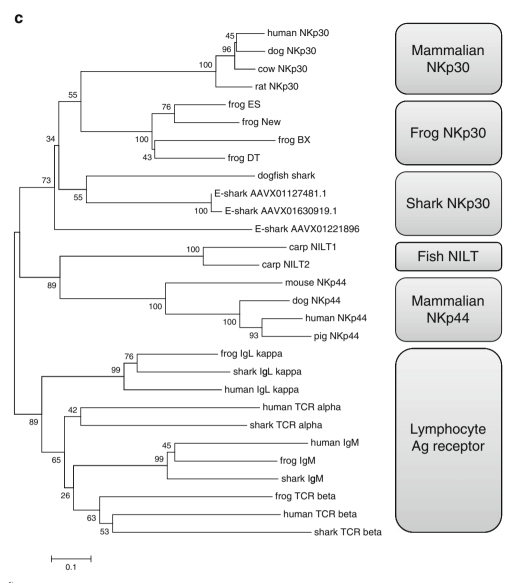


Fig. 4 (continued)

isoforms may be co-evolving with some *Xenopus* B7H6. Human NKp30 has six splice variants (Kaifu et al. 2011), some of which are differentially expressed in a tissue-specific manner (Delahaye et al. 2011). *Xenopus* EST ES685492 contains one ITIM motif (double underlined in Fig. 4b) in the cytoplasmic region and it lacks the arginine (R) in the TM that could interact with ITAM-containing adaptors; thus, as is typical for other vertebrate NKR—but not specifically for NKp30 in other species- *Xenopus* NKp30 differing in their cytoplasmic tails might be either stimulatory or inhibitory.

It was previously recognized that NKp30 is present in rat, cow, and dog (Fig. 4b), but that mouse *NKp30* is a pseudogene (Hollyoake et al. 2005; Kaifu et al. 2011). Neither could we identify *NKp30* in opossum, chicken, or bony fish. The presence or absence of both *NKp30*

and *B7H6* in different species correlate precisely (Table 1); this finding, in concert with the expansion of both genes in *Xenopus*, is strongly suggestive of co-evolution of this receptor/ligand pair over hundreds of millions of years.

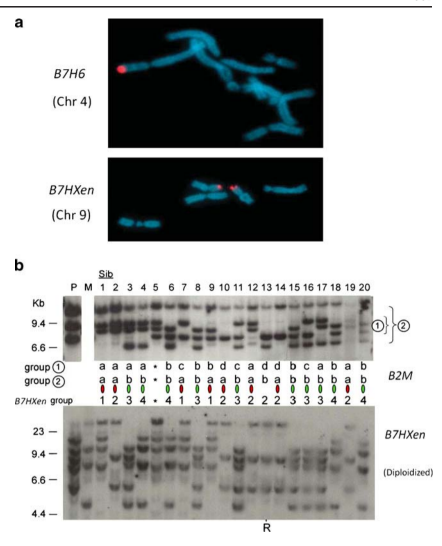
Mapping of B7H6, B7HXen, and NKp30 in the Xenopus genome

To further study the evolutionary origin of *B7H6*, we mapped both *B7H6* and *B7HXen* onto *X. tropicalis* chromosomes using the TSA-FISH system (Macha et al. 2003).

B7H6 and *B7HXen* were mapped onto the telomeres of *X. tropicalis* chromosomes 4 and 9, respectively (Fig. 5a). Since no other regions hybridized to the probes, all genes



Fig. 5 Genetic mapping of Xenopus B7H6 family members. a Chromosomal mapping of B7H6 and B7HXen in X. tropicalis using the TSA-FISH system. The Xenopus chromosome number was assigned following Wells et al. (2011). The coding region probes are shown in red; DAPI counterstaining is shown in blue. **b** Linkage of B7HXen to the B2M. Southern blotting was done using a previously typed X. laevis family with 20 siblings along with parents (Paternal and Maternal) (Ohta et al. 1999). Two *B2M* loci are indicated as groups ① and ②, while four segregating patterns with B7HXen are indicated as numbers 1-4. Matched groups are connected with colored ovals (red or green). Sibling 5 is a natural triploid. Size marker positions are marked at left



were tightly clustered only in these chromosomal regions, suggesting that expansion of the genes was likely the result of the *cis*-duplications. The expansion of genes in the telomeric regions seems to be a common phenomenon since it was observed for other *Xenopus* genes: e.g., non-classical *Xenopus* MHC class I genes are found in a large telomeric cluster on the same chromosome as MHC (Courtet et al. 2001; Flajnik et al. 1993).

B7H6 To gain more insight into the genomic synteny of B7H6 family members, we further analyzed genes neighboring B7H6 in the genomic scaffolds. Scaffold-707 contains two other genes (PLEKHA7 and RPS13) both of which map in the vicinity of B7H6 on human chromosome 11p15 (Online Resource 2). The scaffold-1420 contains only one other gene, SLC6A11 which maps to the telomeric region of human chromosome 3p25.3 (Online Resource 2). We also examined the X. tropicalis genetic map (http://tropmap.biology.uh.edu/) for synteny, but there were no microsatellites associated with the B7H6-containing scaffolds and thus no other associated genes or linkage groups could be established.

B7HXen Most of the B7HXen-containing scaffolds were small and did not contain other genes, and none of the gene models in larger scaffolds (e.g., 1119 and 867) revealed clear orthology to any human gene. However, upon examination of the X. tropicalis genetic map, we found that the B7HXen-containing scaffold-1146 was located near scaffold-673, which contains beta-2 microglobulin (B2M) (Stewart et al. 2005). To confirm their close linkage, we performed a segregation analysis using genomic Southern blotting of a X. laevis family with 20 siblings. Unlike the true diploid X. tropicalis, X. laevis is tetraploid (Kobel and Du Pasquier 1986); nevertheless, genetic synteny is usually conserved between these species. Consistent with the tetraploidy in X. laevis, two independently segregating B2M loci are present (groups 1) and 2 in Fig. 5b) (Stewart et al. 2005), whereas the B7HXen seemed to be diploidized (i.e., silenced on one chromosome), showing only four allelic segregation patterns derived from one locus (shown as groups 1–4). One set of B2M loci (group ②) matched perfectly to the B7HXen RFLP pattern except for sibling 13 (B2M RFLP "a" to B7HXen group 1 and 2; B2M RFLP



"b" to B7HXen group 3 and 4), confirming the close linkage of B7HXen and B2M (Fig. 5b). B2M is linked to the shark MHC, and in conjunction with other phylogenetic evidence the presence of B2M in the MHC is primordial (Ohta et al. 2011). Therefore, the linkage of B7HXen to B2M in Xenopus further implies that the precursor of B7 family genes was present in the proto MHC before genomic expansion and extensive reorganization during vertebrate evolution.

NKp30 NKp30 maps to the MHC in human and other species, but it is not in the *Xenopus* MHC; instead, we found a cluster of genes with a very similar domain structure as NKp30, the Xenopus MHC-linked Immunoglobulin superfamily Variable genes (XMIV), in the same location as human NKp30 (Ohta et al. 2006). Both XMIV and NKp30 contain "VJ"-type IgSF domain with the "GXG" motif and we predicted that these genes were derived from a common ancestor. However, phylogenetic tree analysis did not reveal a close relationship between XMIV and NKp30 genes (data not shown), suggesting that they may be evolving at different rates. Xenopus NKp30 maps to linkage group (LG) 7 (LG 4 in the new assignment (Wells et al. 2011)). Scaffold-88 contains three microsatellites, 036C01, 040A12, and 022B03, in the corresponding position 12.56-13.26 cM, and is sandwiched between scaffolds-128 and 1. Scaffolds-128 and 1 contain the *adamts18* and *e2f4* genes, respectively (Wells et al. 2011), and map to the long arm of chromosome 4 from centromere to telomere orientation (Khokha et al.

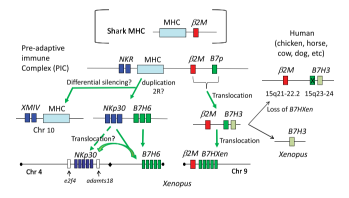


Fig. 6 Model for emergence of the two sets of NKR in *Xenopus* and B7 evolution from the MHC precursor. We hypothesize that the MHC precursor contained precursors of NKR, MHC class I and II, including *beta-2 microglobulin (B2M)*, and *TAPBP/B7* (denoted as B7p) genes. Via two rounds of duplication (2R), four MHC paralogous regions were formed followed by further genetic modification resulting in more paralogous fragments (also see Fig. 7). In *Xenopus*, two sets of MHC/NK regions can be followed by differential silencing of the loci, resulting with XMIV and MHC on chr 10 and *NKp30* and *B7H6* on chr 4. Chromosome 4 may have been further modified by a translocation of *NKp30*, followed by the expansion of *NKp30* and *B7H6* genes. The B7 precursor might have also been translocated along with *B2M*, generating *B7HXen* next to *B2M* on chromosome 9. The synteny of NKR was observed in many vertebrate genomes ((Belov et al. 2007; Kaufman et al. 1999)

2009; Wells et al. 2011) (Fig. 6). Thus, *Xenopus NKp30* is located in the middle of the long arm of chromosome 4. Note that *B7H6* mapped to the telomere of the short arm of chromosome 4 (Fig. 6).

In order to further study the evolutionary history of NKp30 and XMIV genes, we examined the synteny around NKp30 on scaffold-215. NKp30 genes are clustered at the 5' end of the scaffold, adjacent to beta-1,3-glucuronyltransferase 1 (B3GAT1) (human chr 11q25), followed by genes found in the telomeric region of human chromosome 3p21.3 (Online Resource 3). NKp30 genes are located at the 3'-end of scaffold-88 that contains genes mostly found in human chromosomes 3p21-25 and 22q13 (Online Resource 3) and these regions do not contain MHC paralogous genes and were never recognized as MHC paralogous regions (Flajnik and Kasahara 2010) (Fig. 7). All of the other scaffolds were short and only contain a single NKp30 gene. Inconsistent synteny around NKp30 genes between Xenopus and other species suggests that there has been a genomic modification at and around the locus, perhaps the result of translocation from the other end of the chromosome (Fig. 6); this scenario is consistent with the great expansion of genes in this species.

B7 family members contain V-IgSF domains that share a common ancestor with TAPBP (Greenwald et al. 2005), which maps to the MHC in all species so far examined. Based on the similar domain structure and the presence of "VJ"-type IgSF domains in both XMIV and NKp30, it is reasonable to predict that XMIV and NKp30 share a common ancestor that was linked in the primordial MHC. The close linkage of B7HXen and B2M and the mapping of B7H6 and NKp30 to the same chromosome further suggests that the precursor of B7 was also linked to the primordial MHC. Furthermore, the human chromosome region to which B7H6 maps (11p) has been considered to be an MHC-paralogous region (Du Pasquier 2009; Flajnik and Kasahara 2010) (Fig. 7). Thus, we hypothesize that XMIV and NKp30 were generated by a cis-duplication from an NKR precursor before the genome-wide duplication in the pre-adaptive immune complex (PIC), after which the MHC paralogous regions experienced differential silencing; this resulted in XMIV remaining with the MHC, and the movement of NKp30/B7H6 to another region. The chromosome containing NKp30 and B7H6 was further modified, presumably by translocation, leaving B7H6 at the telomere and NKp30 near the center. Since we detected no gene remnants of NKp30 or B7H6 near the large clusters, the major gene expansions of both NKp30 and B7H6 probably took place after the separation of these loci (Fig. 6). Interestingly, B3GAT1, which is immediately adjacent to NKp30 genes in Xenopus scaffold-215, is found on the telomeric end of the human chromosome 11q, whereas B7H6 is found in the center of the chromosome (Fig. 7); this suggests that the genome



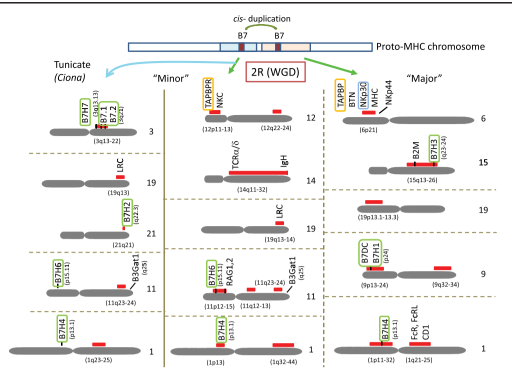


Fig. 7 Most B7 family members are distributed on the MHC paralogous regions inferred from the human and *Ciona* genomes. The four sets of "Major" and "Minor" paralogous regions were identified based on the distribution of homologous, duplicated genes related to the MHC and neurotrophins (Flajnik and Kasahara 2010) (*right* and *middle columns*). The additional "Minor" paralogues were identified from comparative genomic analysis of IgSF and CTX/JAM family members in the tunicate *Ciona* (a lower deuterostome whose ancestor arose before the emergence of the vertebrates and the 2R whole genome duplication (WGD)) and human genomes (Zucchetti et al. 2009) (*far left column*). There are other chromosomes that contain paralogous regions,

but we only show chromosomes relevant to the B7 family. Paralogous human chromosome numbers are indicated on the right side of the chromosomes. *Red bars* indicate the paralogous regions among the four sets of paralogous chromosomes. All B7 family members except B7H5 are found in the canonical paralogous regions and they are boxed in green. Since B7 family members are found in both "Major" and "Minor" paralogous sets, we predict that the B7 precursor was duplicated in the proto-MHC and then distributed to both paralogous regions as a result of WGD. Since V domains of TAPBP and B7 are similar, they may share the same common ancestor. TAPBP maps to MHC. TAPBP and related genes are boxed in *orange*

modification likely took place after the emergence of the amphibian lineage.

The phylogenetic tree was not informative regarding the affiliation of B7HXen to other B7 family members, but its linkage to B2M might be more revealing. The B2M translocation from its original location in the MHC (discussed in (Ohta et al. 2011)), the close linkage of B2M and B7HXen in Xenopus, and the wide distribution of most B7 families in MHC paralogous regions (see below, Fig. 7) support the hypothesis that B7 precursor ("B7p" in Fig. 6) was encoded in the proto MHC. Our hypothesis is also consistent with the previous observation of a connection of B7 to the MHC (Henry et al. 1999). In addition, B7H3 is on the same chromosome as B2M in many mammalian species, but Xenopus B7H3 is not closely linked to B2M (Fig. 6). We suggest that following the co-translocation of B2M and a B7 precursor early in vertebrate history, a secondary translocation occurred in *Xenopus* splitting *B7H3* and *B7HXen* onto two chromosomes. B7HXen subsequently expanded greatly, and thus it is difficult to ascribe its orthology to other B7 members; in contrast, the translocated *B7H3* remained a single-copy gene and evolved at a slower rate (Fig. 6; genomic data not shown).

Identification of Shark B7H6 and NKp30 genes

Since some B7 orthologues were found in the shark databases and based on the presence of *NKp30* genes in the *Xenopus* genome, we further examined the cartilaginous fish databases. We used the human B7H6 V domain to search the elephant shark (E-shark) genomic scaffolds (Venkatesh et al. 2007) and found 5 sequences hitting with E-values under e⁻⁵: four sequences contain only V domains (AAVX01166257.1, AAVX01015110.1, AAVX01166967.1, AAVX01076204.1) and one sequence contains a C domain (AAVX01231033.1). Phylogenetic tree analysis showed clustering with other B7H6, confirming the orthology (Fig. 2). All V sequences showed diversity distributed throughout the domain (Online

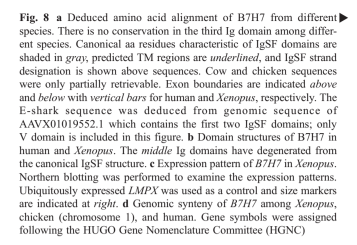


Resource 5), and thus shark B7H6 seems to belong to a multigene family. We also identified a cartilaginous fish (Squalus acanthias, spiny dogfish) EST sequence (ES788778) that showed high similarity to other species' NKp30 (E-values<e⁻⁹). Using dogfish NKp30 as bait, we identified five NKp30 genes from the E-shark genome (Online Resource 5; the V-IgSF domain of one representative sequence is shown in Fig. 4b). Phylogenetic tree analysis showed clear clustering of the shark NKp30 to other species' NKp30 (Fig. 4c). Like NKp30 in most vertebrates, a "VJ"-IgSF domain is present, but the C-terminal Tm domains were not identified by the gene models, perhaps due to the small exon size or the short scaffolds (Fig. 4b). Such conservation of NKp30 during vertebrate evolution was unexpected since NKp30 is not detected in any bony fish species and rapid expansion and contraction of NKR families in a speciesspecific manner is a general rule (Yoder and Litman 2011), especially the activating NKR. The presence of both B7H6 and NKp30 genes in cartilaginous fish species demonstrates that this system is evolutionary old, in fact the most ancient among NKR families. Thus, we suggest that the selection pressure on NKp30 is exceptional and quite different from most other NKR, perhaps a result of its co-evolution with the self ligand, B7H6. The expansion of both B7H6 and NKp30 in Xenopus and shark, and the absence of both genes in other species, correlate well with a co-evolving receptor/ligand relationship, perhaps dating back to when their precursor genes were encoded in the proto MHC.

Our phylogenetic tree analysis further showed an unexpected sister grouping of NKp30 with *NKp44*. NKp44 is expressed on the surface of NK cells and contains a V-IgSF domain (but is not of the "VJ"-type) (Cantoni et al. 1999) and non-mammalian orthologs were found only in some bony fish species including carp (Stet et al. 2005), trout (Kock and Fischer 2008; Ostergaard et al. 2009), and zebrafish (Ostergaard et al. 2009). *NKp44* maps in the TREM cluster region outside the MHC in the human and mouse genomes (Allcock et al. 2003). The clustering of the NKp30 and NKp44 V domains in phylogenetic analysis, their association with the MHC, and their presence in lower vertebrates identify these immune molecules as ancient and unique members of the NKR family.

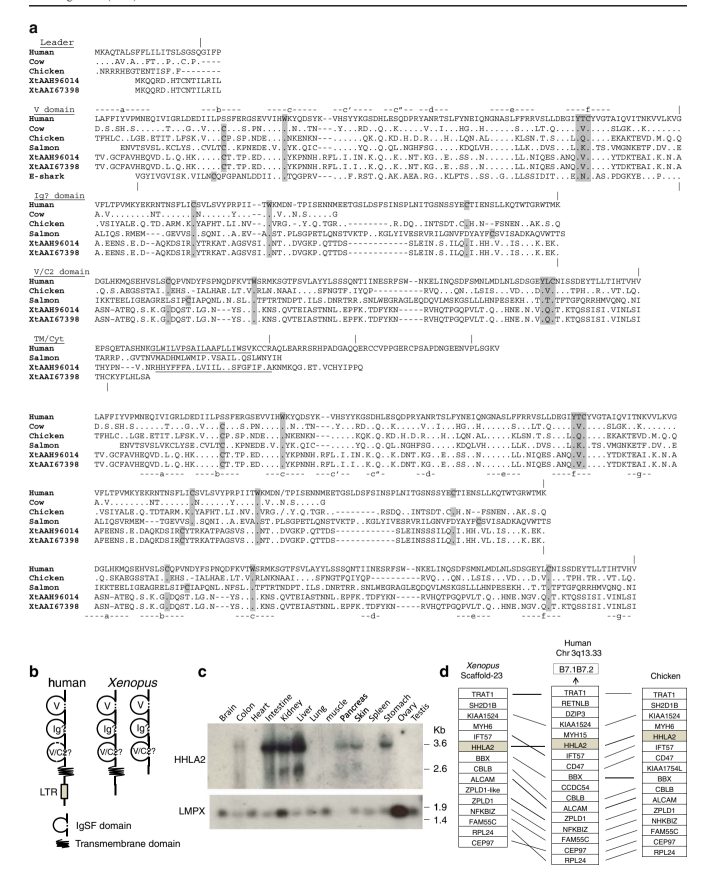
The new B7 family member: HHLA2

During the search for other B7 family members from the *Xenopus* databases, we uncovered an additional new sequence (AAH96014) that was most similar to B7H4. *Xenopus* AAH96014 matched with significant similarities (Evalue ~e⁻²⁰) to human <u>HERV-H</u> <u>LTR-associating 2</u> (HHLA2) having an unknown function (Mager et al. 1999). *HHLA2* was so-named because a sequence in the 3'-untranslated region has polyadenylated long terminal



repeats (LTR), but no further characterization has been done. Our phylogenetic tree analysis clearly shows that HHLA2 is a new B7 member, and we re-designate it B7H7. Our database searches revealed orthologs in hoofed mammals (e.g., horse, pig), chicken, opossum, salmon, and shark, but only pseudogenes in rodents (mouse and rat) (Table 1). The aa sequence alignment of HHLA2 showed a similar domain structures in all species, with conservation of the canonical aa residues for IgSF domains (Fig. 8a). HHLA2 contains three extracellular IgSF domains; one V-type, followed by a degenerate Ig-like domain that still was identified as a C1-type, one V/C2-IgSF domain, and a TM region (Fig. 8b). BLAST searches revealed that the V-IgSF domains from all species were most similar to either B7H3 or B7H4, depending on the species. The similarity of V-IgSF domain to B7H3 and B7H4 by BLAST was confirmed by the phylogenetic tree analysis, clearly demonstrating HHLA2's close relationship to other B7 family genes (Fig. 2). The other two IgSF domains matched to B7H4 $(\sim e^{-4})$, TCR/IgL $(\sim e^{-6})$, BTN $(\sim e^{-7})$, MHC class II $(\sim e^{-5})$, and CD4 (~e⁻⁷), along with other non-immune genes (ROR2 and NRCAM). Human B7H7 is expressed in gut, kidney, and lung, and it was speculated that B7H7 is expressed in epithelial cells. However, B7H7 was also detected in lymphocytes in the original analysis (Mager et al. 1999). Xenopus HHLA2 is expressed highly in intestine, kidney, liver, and stomach and weakly in pancreas, colon, and skin (Fig. 8c), suggesting that it may be involved in mucosal immunity. We identified two bands on the northern blot, and upon reexamination of the databases, we found another EST sequence (AAI67398) for Xenopus HHLA2. Both EST clones are splice variants, and AAI67398 lacks the TM region (Fig. 8a). Both clones are full-length (2,300 bp (AAH96014) and 5365 bp (AAI67398)), and it seems that the rapidly migrating band with the lower expression in northern blotting correlates with the TMcontaining splice variant, whereas the higher expression band correlates with the secreted variant. Xenopus HHLA2







is found in genomic scaffold-23 and genetic synteny between human, chicken, and *Xenopus* is well conserved (Fig. 8d). Human *HHLA2* is found on chromosome 3q13.33 near B7.1 and B7.2, which is a region rich in genes containing IgSF domains and one of the regions assigned to be a MHC paralog (Du Pasquier et al. 2004; Zucchetti et al. 2009) (Fig. 7).

Phylogenetic tree analysis of B7 family

Building on a previous study that centered on teleost fish (Hansen et al. 2009), we conducted updated phylogenetic tree analysis including B7H6 and B7H7. We also searched for B7 family members from shark databases and added them in the analyses. Our phylogenetic tree analysis showed that B7 family members were categorized into six major groups (Fig. 2), and we discuss each in turn.

B7.1/B7.2 group The CD28 ligands, B7.1 (Selvakumar et al. 1992) and B7.2 (Reeves et al. 1997), were the founding B7 molecules and are the best characterized B7 family members. Both B7.1 and B7.2 are present in mammals and their redundant roles in costimulation and their chromosomal positions had implied that they were recent duplicates. However, the presence of B7.2 in Xenopus and a B7.1 ortholog in all teleost fish (called B7R; (Hansen et al. 2009; Zhang et al. 2009)), as well as co-stimulation by trout B7R (Zhang et al. 2009), suggest that they are ancient. Interestingly, B7R formed a unique cluster within the B7 group in our phylogenetic tree analysis. Bony fish experienced a species-specific duplication of the genome (Meyer and Van de 2005) that generated separate clades for some genes and a general disruption of their genome (Postlethwait 2007). We could not detect opossum or *Xenopus B7.1* or both shark B7.1 and B7.2 (Table 1).

B7H2 group Phylogenetic analysis showed that B7H2 is most similar to B7.1 and B7.2. B7H2 was not found in teleost fish and genomic synteny is not well conserved between *Xenopus* and other species. However, B7H2 maps to human chromosomal region 21q22.3, which is likely to be paralogous to chromosomal region 3q13-22, where B7.1 and B7.2 are found (Du Pasquier et al. 2004) (Fig. 7). Thus, B7H2 and B7.1/B7.2 may be derived from an immediate common ancestor. Another area that is paralogous to these regions is chromosome 1q23-25, an MHC-paralog (Fig. 7). Therefore, the origin of these two B7 groups also has an MHC connection.

B7H1/B7DC group B7H1 and B7DC are adjacent to each other in all species that have both genes (e.g., human and Xenopus), and thus they were predicted to have arisen by a cis-duplication. B7H1 and B7DC genes were reported

previously in the chicken genome (Hansen et al. 2009): however, our analysis showed that the both clustered with mammalian B7H1, and thus are clearly duplicated B7H1 genes (Fig. 2). Two cis-duplicated genes were identified from the Xenopus genome and each picked up B7H1 and B7DC by Blast. Teleost fish have only one gene, which matches equally to B7H1 and B7DC in the Blast analysis (e.g., fugu CAF93166 in Fig. 2), suggesting the preduplicated state. There are two E-shark sequences (AAVX01047137.1 and AAVX01029406.1: similar to each other \sim 76 % at the aa level) that are most similar to B7H1. Phylogenetic analysis including all vertebrate species revealed that both Xenopus B7H1 and B7DC, the two shark B7H1, and fugu B7H1/DC clustered together, but did not group with mammalian B7H1 and B7DC. Thus, Xenopus B7DC may not be orthologous to mammalian B7DC. Taken together, the timing of the separation of B7H1 and B7DC may be relatively recent.

B7H3/B7H4/B7H7 group Here, we demonstrate that HHLA2 (Mager et al. 1999) is a new B7 family member, B7H7. B7H7 is found in an IgSF-rich genomic region on human chromosome 3q (Du Pasquier et al. 2004) along with B7.1 and B7.2, and it was likely generated by an ancient duplication from the common ancestor of B7.1 and B7.2 (Fig. 7). A previous report noticed a weak similarity of B7H7 (HHLA2) to the B7 family (Fahrer et al. 2001), but we have found that B7H7 clearly is related to B7H3 and B7H4 in the phylogenetic analysis (Fig. 2). A B7H3 ortholog was identified from skate (Hansen et al. 2009) (with E-values \sim e⁻⁷⁶ and \sim e⁻⁴⁵ to the human entire molecule and V-IgSF domains, respectively) and E-shark (AAVX01064573.1). Similarly, we found B7H4 (AAVX01079670.1) and B7H7 (AAVX01019552.1) from the E-shark genome, suggesting that this group of B7 family members is evolutionarily old (Fig. 2). The functions of these newer B7 family members are not fully known, but their phylogenetic conservation suggests important roles in immune regulation. There are several other B7-related genes in the shark and bony fish databases that clustered separately but belong to this group. However, we could not detect clear orthology to specific B7 family members (data not shown) and thus we did not include them in our analyses.

B7H5 group B7H5, also known as VISTA or Gi24, was recently discovered as an inhibitory B7 molecule expressed by APCs and T cells (Wang et al. 2011), and it is conserved among different species. We found dogfish shark EST sequence (e.g., DV500095) and E-shark sequence (AAVX01038049.1) with low E-values ~e⁻²⁷ and ~e⁻¹⁸ to *Xenopus* and human, respectively.

The V domain of B7H5 matches to the V domains of T cell receptors in Blastp analysis and B7H5 consistently



clustered distantly from other B7 members with long branch-lengths when the tree was made un-rooted, and thus it was used to root the trees. B7H5 maps to human chromosomal region 10q22.1, which has not been assigned to any MHC paralogous region (Fig. 7). Furthermore, B7H5 has a domain structure V-TM and lacks the C domain, which is distinctly different from other B7 family members (Table 1). All of the data to date suggest that B7H5 may be derived from a different precursor than all other B7 family members.

B7H6 group The presence of shark B7H6, the clustering of B7H6 with the TAPBP/TAPBPR group, and the presence of the unique C1 domain in B7H6 suggest that B7H6 arose early in vertebrate evolution and was lost in bony fish. In addition, the discovery of Xenopus and shark B7H6 has further highlighted the plastic features of B7 family genes: (1) The number of IgSF domains: Most B7 family members have two-domain structures (Table 1), but human B7H3 has duplicated its V- and C-IgSF domains, generating a longer form with 4 IgSF domains (Steinberger et al. 2004; Sun et al. 2002). Other members have three-domain structures (e.g., B7HXen and B7H7). Thus, it is clear that B7 family members need not conform to the two-domain paradigm. The cytoplasmic IgSF domains in Xenopus B7H6 and B7HXen are so far unique in vertebrates, and duplication of these domains has also been detected. These Xenopus cytoplasmic IgSF domains, which presumably play some role in signaling events, are puzzling and await biochemical analysis. Since receptor/ligand interactions occur via the extracellular domains in the B7 family, our assignation of orthology has been established with these domains. (2) Gene duplication: As we found for the Xenopus B7H6 and B7HXen genes, B7 genes are often duplicated. B7H1 and B7DC were generated by tandem duplications, and Xenopus B7H4 was duplicated in tandem, resulting in two genes with ~48 % similarities at the aa level (Table 1). Other species-specific B7 genes may have emerged by duplication as well. For example, one B7 gene model (Xt1303) from X. tropicalis scaffold-1303 is adjacent to the orthologous gene next to B7.1 (Fig. 2). Xt1303 has a predicted three-domain structure and BLAST searches matched closest to B7H1 with $\sim e^{-10}$ for all three domains, but it does not cluster in the tree with any particular family member. (3) B7 precursor candidates: BLAST searches and phylogenetic tree analysis further revealed a connection between Xenopus-specific B7HXen and hagfish IGSF3 genes (Haruta et al. 2006) (Fig. 2), suggesting that hagfish IGSF3 may be related to the common precursor of all B7 family members; if true, this would be the first B7 family member detected in taxa more primitive than gnathostomes. In the sea urchin, there are more than 50 genes encoding at least three types TM proteins with V- and C1-like Ig domains having low but intriguing sequence similarity to Ig/TCR/MHC, as well as a number of Ig-domain-containing proteins with similarity to the B7 co-stimulatory molecules (Sodergren et al. 2006). Our database searches did not detect any of these genes, but clearly they are worth examining in future expression/functional studies.

Genomic history of the B7 family

The chromosomal distribution of the B7 family members into paralogous regions in the human genome is consistent with the ancient genome-wide duplications (WGD) early in the history of vertebrates (Flajnik and Kasahara 2010; Ohno 1970) (right and middle columns in Fig. 7). All B7 members are distributed into the MHC paralogous regions; chromosomes 1 (B7H4) and 9 (B7H1, B7DC) as "major" paralogs, and 11 (B7H6) and 15 (B7H3) as "minor" paralogs (Flajnik and Kasahara 2010). Additionally, from the study on IgSF genes in the *Ciona* genome, chromosomes 3 (B7.1, B7.2, B7H7) and 21 (B7H2) have been identified for their connection to the MHC paralogous regions on chromosomes 1 (Du Pasquier et al. 2004) and 19q (Zucchetti et al. 2009) (left column in Fig. 7), suggesting that these chromosomal regions were associated with the proto-MHC and translocated out before WGD. Since B7 family members are found in MHC paralogous regions, the B7 precursor must have arisen before the emergence of the vertebrates (Proto-MHC in Fig. 7). B7H4 genes may have been differentially silenced in human and *Xenopus*, resulting in two paralogous regions; the Xenopus B7H4 gene is linked to genes found on human chromosome 3q13 in the vicinity of B7.1, while human B7H4 is on chromosome 1 (Fig. 7). Alternatively, Xenopus B7H4 could have been translocated from the equivalent location to human chromosome 1. Furthermore, the similarity of the B7 V domains to V domains found in MHC-linked BTN and MOG is also consistent with the B7 family's original presence in the proto MHC (Henry et al. 1999).

Our linkage analysis in Xenopus showed close linkage of B7HXen to B2M, XMIV mapping to MHC (Ohta et al. 2006), and NKp30 mapping to the same chromosome as B7H6. In conjunction with the mapping of NKp30 to the MHC in other species, all of these data suggest that the B7 precursor was likely encoded in the proto MHC along with B2M (shown from shark linkage study (Ohta et al. 2011)), NKRs (shown from linkage analysis in chicken (Kaufman et al. 1999) and opossum (Belov et al. 2007)), and perhaps antigen receptors (Flajnik and Kasahara 2010). Antigen receptors play the central role in adaptive immunity and their evolutionary origin has been speculated to be associated with the MHC. One theory for the origin of antigen receptors is that the C1-type IgSF domains became associated with "VJ" domains encoded within the same genetic region (Bartl et al. 1994; Du Pasquier 2002;



Du Pasquier et al. 2004). The presence of NKRs of the germline-joined "VJ"-type in the MHC (i.e., NKp30 and XMIV), and the fragments of synteny with C1-IgSF-containing B7 homologs (i.e., B7H6 and B7HXen) that are associated with the MHC certainly are consistent with this scenario (Fig. 6).

Conclusion

In this paper, we have shown: (1) the general scheme for evolution of the B7 family; (2) a new member of the B7 family, perhaps the last one to be discovered that is evolutionarily conserved; and (3) the evolutionary history of the B7H6 and NKp30 families, and we speculate that they persist over evolutionary time in a co-evolving receptor/ ligand unit. Furthermore, we perpetuate the theme that B7 family members and NKR (genes mostly in LRC and NKC) were encoded in the proto MHC. Various B7 family members have been identified from gnathostomes (Hansen et al. 2009; Zhang et al. 2009), but so far not from jawless vertebrates or invertebrates, strongly suggesting that the immune regulation associated with most B7 family members arose with the emergence of the Ig/TCR/MHC-based adaptive immune system. Indeed, the emergence of a large number of both B7 and TNF superfamily members (Collette et al. 2003), all involved in regulating adaptive immune responses, occurred rapidly in concert with the advent of adaptive immunity. When observing this system from an evolutionary perspective, it is fascinating how complicated networks of regulation must have fallen into place rapidly to harness the nascent Ig/TCR/MHC-based immune system. This also suggests that while the jawless fish have adaptive immunity complete with T and B cells, the gnathostome system must be regulated in a much more complicated fashion since the majority of costimulatory molecules, as well as most members of the cytokine and chemokine families arose after the split between agnathans and gnathostomes.

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Online resource 1a. Alignment of cytoplasmic Ig-like domains of *Xenopus* B7H6

Cytoplasmic	<pre>Ig-like domain-1 (Ig)</pre>
AAI66200	LKKEPPKLSPITGAEHLVHMNEATLSCQISGFRPKPIRITLCLQRKGGEVMEIYSWDSEAQTDHSIKRKELSVPIEL
AAH84632	YVARLW.K.NEDTGSSAPV.HH.E.QRVV.VP
AAI66200	EREGLVGNGE-TNLPPTQRALRVEMVPLIKTSKKHISNCQCTIRITPDIEEDDGAELTVRVTHSALRGPISVSCRLKVNGV
AAH84632	MSNGAP.QV.TSKSQRLHNLL
Cutonlasmia	domain-2 (C1)
AA166200	PPTISDIVPPLRILNEEPVTLTCPINGFKPKPLSVTWLKVDSGGRETELLSWDRGSTNIRDP
AAH84632	K.KLAK.ISHHGLYQRQ.TN.
AAII04032	к.кпак.10пп
AAI66200	KYSHOMLENEHGEHSNAFISVLAMKPTVKEDHGAKYICRTYHYATGSOADKTMEMLVS
AAH84632	LA.DER.KEEEDDI.Y.V.N.TLYO.VHT.EHT.E.R.K.O.L
Cytoplasmic	domain-3 (C1)
AAI66200	AVPELGPIQKSPEILYVGEEMTLSCRIHSFHPEPLKVRWYKGDELLESSTD
BX732750	.G.CA
AAH84632	SDENAVD.KRT.D.SENS.YN.
AAI66200	QAHRGASHLLDCTSRATYRPSVRDIGKSFRCEVSHESLQHPKCVSWELKHL
BX732750	N
AAH84632	PILHNN.G.FHFTT.CTPTNY.
a . 1	
	domain-4 (Ig)
AAI66200	VSEPRVSAIKCHPEAPECNQPATLSCAIENYYPKDLKIRWYRGLEVISSHN
BX732750	I
AAH84632	IDV.GTTDKFWK.EG
AAI66200	DTETKEDPPGSGLFSKMTEIHLTPTHSDHGTEFHMEIIHNMKITKKTFCLOFKG
BX732750	PKAA.EDOERTSOIIINYLK.T.GNFE.N.PMKLR.
AAH84632	P.GAKED.ERTOIIINKO.LFYSMKC
AAII04032	.F.GARED.ERIQIIINRQ.DF.IISMRC
Cytoplasmic	domain-5 (C1)
AAI66200	FPEVKDITCNKYPPKYGEALTLSCEVIGCSARDITAEWREGNNPIRGOMRAEPQI
BX732750	L.D. EVKRE.E.KT
AAH84632	
AAI66200	DRDSVTFLLTLTPTAEHYGKLLTCLIKHKDLPQPIKKNISLKLP
BX732750	I
AAH84632	ESFE

Online resource 1b. Deduced amino acid sequence of B7HXen.

Leader

AAH85212 MVSRLFLLLLLLRIAI

AAI61104 MGDPGGAAVLLRFLRIQGFLCLFLLPQT

Extracellular V domain

AAH85212 FSALEMTAPPTHRAEMGSNIIIPCTFRVDEPPVDLKFLAIIWHFQDKEVLNYTNTSLV
AAI61104 GSVLEVIARSPHVAQKGSNTLIPCTFRVNKFPADPKFLAIYWDFNGRRILTYDN--VV

AAH85212 STQNPRLSLNKDTTEDGVASLIISNVTISDGGLYRCRVLYSPKHMYKEVRLDIQ
AAI61104 STTDPRFSLDPSSAPRGEASLSVSNAQISDGGAYSCSVTYSPEQQQKEITVEIQ

Extracellular Ig domain

AAH85212 APPQITITNKVIIKDKESVLSASITRFYPVDIDIKWLRDGKILAGVILLTPQRNVDGT AAI61104 APPQITITNRVVTENKESALRASIIGFYPQDLDVTWLRDGAILRGVTMNKPQRDPDGT

AAH85212 YQVNNYMTIVPTEENRKQNFSIWVQHESLSSPLQEDFQLIYG AAI61104 HSTTSAVTIVPTEGNRNQNFSIRVQHESLSVPLQEDFQLLYA

Extracellular C domain

AAH85212 AAPSIHITHQLFHLHREQELVCQAWGFYPESIVMNWFLNGSHVETANRKTVNSSAVES
AAI61104 AAPSIHISHGSFYVNEEQTLMCRVWGYYPEDIAVSWFLNGSRVEPSEIKRINSSALEL

AAH85212 IYRILPTDQNWGMEISCEVEHQTLRSPLVEKLLILGN AAI61104 PYRFLPTAESQGMEISCVVEHRALTEPLVQSLKVELT

TM region

AAH85212 DGKVKHRAVVLSIAILSVIITFTLISLFLWRKFSSPK
AAI61104 DAQWSNTLIIPIAIVVVALVAVVAVLILKKKKRCCIRK

Cytoplasmic domain-1 (Ig)

AAH85212 APELLGPIEYSLCGAEEIQYSLNLONFYPRDINIKWTYGERNEKVNTSSNKFIRTAGE

AAH85212 KTFSVVSECRIPWHQYTMRVIWEHETLAEPQYREVRVT

Cytoplasmic domain-2 (C1)

AAH85212 DLPWRPVMSQILTPDVYVGTEAEIQCNISGYFPDKLTVTWYKKKENGKEELVNNGGRY
AAH85212 QIPDIQSQYQPDWTLTCTARLLFSPSLTEEHGTQFICRVKHPSLGEEIERRTGTLYVR

Cytoplasmic domain-3 (Ig)

AAH85212 ARPKVSKFRMAGEWGLSLEAEGFFPRDIGFSWEVVOGNKRDVTSLPCEFSISENTDGT

AAH85212 YKATSTCHCLKDQLNYLHTLRVSVQHEALGAPVCKDITQ

Cytoplasmic domain-4 (C1)

AAH85212 DFPWRPQVGEIVVPELIELEETQLTCNISNYFPDALTVTWFIKNKESEELTDVTQYIR AAH85212 YRHDTVSQYQPDQTLTCTTSLTFSPSLGMDLGAEFICRVKHPSLGEEIERRTGTLYVR

C-terminus domain?

AAH85212 GGPTSLSP*

Online resource 1c. Alignment of V-IgSF domain of Xenopus B7H6

AAI66200	DALGVKTSDVPIKSQRNQEAFIPCTVTDYGAGRVDPKLLSVEWKLKPLNGSEEQVYLYVS 60
1420-1	v
707-5	ITMGGSLVNAL.AHDVI.G.SEL.LTKT.T.SNKAATNQLKFL. 58
707-2	GS.RIEMGAE.VLESW.P.DIS.PKTQRT.D.R.G 60
707-4	GS.R.EMGAE.VLESW.P.DIS.PKAQ.TKG 60
AAH84632	.TFFL
AAH82995	E.VEVS.V.VLKD.DVISR.SSTEL.LQRQ.ID-PI.DG 55
AAI66200	GTPTPTRPGSYIS-RDLITGNVGLHLPRVQFTDEGEYTCTVIYTPNKAVGHSVLQVS 116
1420-1	H
707-5	QS.QDA
707-2	RQ117
707-4	HQ117
AAH84632	HDSEI.RA
AAH82995	.SHN.IOLD-IHKAY.AOVAET.DNEEDE.R.S.E 111

Online resource 1d. Alignment of V-IgSF domain of Xenopus B7HXen

AAI61104	GSVLEVIARSPHVAOKGSNTLIPCTFRVNKFPADPKFLAIYWDFNGRRILTYDNV 55
1657-1	T.PD
1146-1	T.PDIDPL.N.K.NOIR.SCTDSE 55
1146-2	VAS.PLTQISMDIDPL.D.K.HQIRESDKV 56
1119-1	.AALT.PPT.R.TL.ADETP.VRHF.Y.QDKENVTTT 55
1119-1	.AA.DLTVPPS.R.RVD.SLDRRSVL.T.S.Y.QD.ERFPGT 55
1119-2	
1119-3-2	-DA.LIGDEE.PT.TLDVILS.GQP-VSLQYL.T.QNKMLFRL.NK 54
1119-4	.AALM.PPT.R.TL.ADETP.VRH.G.F.S.QDKE.FNVIST 55
867-1	C.ATVPPNQSSPM.RDA.LDNP.MNL.H.GDKEV.RK 55
867-4	C.ATVPPDQSSPM.RDA.LDNP.MNL.H.GDKEV.RK 55
867-5	C.A.K.SDPPS.K.LLTASLSLG.S.I.HSA.S.I.T.RDKER.NKG 55
867-7	A.K.SDPPS.K.LLT.SLSLG.S.INHSA.S.I.T.RDKER.NKG 55
867-9	F.PPTQQTLLV.LQ.ENSQIL.NFVF.YIGDKE.VR.DNKG 54
3534-1	.AALM.PPT.R.TL.ADETP.V.SRHF.Y.QDKENVIST 55
1318-3	I.DIS.PQRVLL.NEAQLRSLA.P.IN.LVF.Y.QDMERK 55
1318-2	.AT.QLA.PPSRQVLVRAILMIS.GNASL.LNY.VVI.F.QNKERFK 55
1318-1	.TT.QLA.PPSRQVLVRAILMIS.GNASL.LNY.VVI.F.QNKERLK 55
1146-3	.AEQ.PPA.TVTLDVTLS.S.GPTQV.LQY.T.L.Y.QDTEFFNAH 55
1707-2	.AEQ.PPA.TVTLDVTLS.S.GPTQV.LQLL.Y.QDTEVFNAHGSI 58
AAH85212	F.AMT.PPT.R.EMIIDEP.V.LI.H.QDKEV.N.T.TSL- 57
Hagfish	ICG.K.ESP.EVGFVD.K.VSLI.TDHSEPIDLSKLELEWTNGHAILTYKNGKI 58
77TC1104	VALUE DE LA PROCES DE CESA DE CESA DE CESA DE CONTRA DE
AAI61104 1657-1	VSTTDPRFSLDPSSAPRGEASLSVSNAQISDGGAYSCSVTYSPEQQQKEITVEIQ 110
1146-1	
1146-2	EFNWTGTDW.E.KTIL111
1119-1	.GSSNLNRETIRA.VIA.VEL.RLDRRYVRLD. 110
1119-2	.GAPNLNKDTTKD.VLTGVRL.KGHGF.RSERLD 110
1119-3-1	.GSSNLNRDTIRA.VIA.VEL.RLDRRYVRLD. 110
1119-3-2	GKQLSVTFSDAD.MK.IVQLH.VSVA.A.I.M.KIIKKE.DLKV. 109
1119-4	.GSSNL.INRDTIRA.VIA.IEL.RLDMRYVRLD 110
867-1	GKVSSV.I.ERALLE.NLVTVT.RIT.KRLR.H 110
867-4	GKVSSV.I.ERALLE.NLVTVT.RIT.KRLR.H 110
867-5	RTLSQA.LAQAIEE.RVVVTVE.T.T.V.S.NM-K.EQGVKL.VA 110
867-7	RTLSQA.LAQAIEE.RVVVTVE.T.T.V.S.NM-K.EHVMELQVE 110
867-9	K.ISHS.VTA.EKGFPQ.NITLVAQ.T.R.LIIHSRKE.DKV. 109
3534-1	.GSSNLNRDTI.A.VIA.IEL.RLNRRYVRLD 110
1318-4	GLSLSV.FSKEA.NN.DV.V.LA.VSI.R.L.IKMEVLLF 110
1318-2	CFSSTS.R.FEQQEVDN.NFALVRSE.V.T.A.VREDR.IRLIVL 110
1318-1	CFSSTS.R.FEQQEVDN.NFALVRSE.V.T.A.VREDIRLNVL 110
1146-3	GKESEVTIREED.GK.IHLAQIRLA.L.K.L.I.I.RSHTVQLTVH 110
1707-2	KEESEVTIREED.GK.IHLAQIRLA.L.K.M.I.I.RSHTVQLTVH 113
AAH85212	ONLNKDTTEDGVIIVTL.R.R.LKHMYVRLD 112
61 1	
Hagfish	ETSSSFKGRVMMNEQNLRNGDA.LILNNVTEVDMLT.ING.TRRDT.ILQYK 113

Figure legend:

a) Alignment of cytoplasmic Ig-like domains of *Xenopus* B7H6 EST sequences. Cytoplasmic Ig-like domains are determined based on the exon-intron organization from genomic sequences. b) Deduced amino acid sequence of full-length EST sequences for *X. laevis* B7HXen. Only the extracellular domains of AAI61104 (*X. tropicalis*) are shown since there is no obvious cytoplasmic Ig domain in this sequence. Domains are predicted based on the sequence similarity to other sequence and exon assignment from *X. tropicalis* genome. c) The deduced amino acid sequences of V-IgSF domains of B7H6 are retrieved from the *Xenopus* genomic scaffolds and aligned with EST sequences. d) The deduced amino acid sequences of V-IgSF domains of *X. tropicalis* B7HXen are retrieved from scaffolds and aligned with EST sequences. The membrane distal V-IgSF of Hagfish IgSF3 sequence was aligned. Gaps and Identical aa residues are indicated with dashes and dots, respectively. Canonical amino acids characteristic for IgSF domains are highlighted in grey and predicted TM regions are underlined.

B7H6

Scaffold-707 (535,221bp)

Gene ID	Position	Protein ID	number of IgSF domains	EST ¹
707-1	33133-40802	101494	5	BX745260 (tadpole embryo),DN082733 (lung)+DN055944 (kidney),
				BX732750 (tadpole stg 35-40) etc.
707-2	45969-58046	397915	6	BX745260 (tadpole embryo)
707-3	65268-68458	184972	1	EL657887 (small intestine)
707-4	71420-96658	397890	5	BX745260 (tadpole embryo), CX888209 (testes)
707-5	123639-178327	184975	15 (2 models?: 7+8)	CX917782 (testes)
707-6	194481-225600	184976	10	CX923745 (testes) etc.
707-7	243887-259814	397873	3	CF374995 (whole body) etc.
707-8	260674-294767	184978	4	CX996494 (intestine) etc.
707-9	314966-320370	184979	1	CF374995 (whole body) etc.
707-10	368340-390340	184981	4	CX923745 (intestine) etc.
707-11	415018-421374	397947	3	CX924635 (testes), CX923745 (intestine) etc.
PLEKHA7	507910-512028	397879	-	mapps to human chromosome 11p15
RPS13	530531-531763	88655	-	mapps to human chromosome 11p15

Scafold1420 (72,679bp)

SLC6A11	217-5750	190860	-	mapps to human chromosome 3p25.3
1420-1	34308-52814	461580	7	CX888807 (testes),CX913047 (testes),CX900575 (testes),CX888664 (testes),
				DN082435 (lung)

B7HXen

Scaffold-1119 (167,738bp)

Gene ID	Position	Protein ID	number of IgSF domains	EST ¹
1119-1	7713-27883	189479	9	CX923462 (testes),CX828489 (brain),EL662685 (small intestine),
				CX976768 (intestine)
1119-2	43003-82565	461214	13	CX805381 (brain),CX797364 (brain),CX798692 (brain),CX795849 (brain),
				CX798948 (brain),
1119-3	83902-140637	189481	17	CX812473 (brain), CX923462 (testes), CX888371 (testes), CX899439 (testes),
				CX820566 (brain), CX828489 (brain), CX900620 (testes),
				EG577962 (whole embryo stg 56-62), EB475845 (whole embryo stg 56-62),
				CX802600 (brain), EB484038 (spleen), CX820565 (brain),
				EL659223 (small intestine), CX905208 (testes), CX810694 (brain), etc.
1119-4	164692-167692	189482	2	CX923462 (testes), DN057478 (kidney), CX828489 (brain)

Scaffold-867 (337,695bp)

867-1	17572-57917	407771	8	CX842746 (brain), DN015523 (heart), EL661102(small intestine)
867-2	104292-110592	460532	4	EL655865 (small intestine),CX905130 (testes),
867-3	116712-125454	187169	4	EL699109 (Limb), CX854937 (brain), DN077598 (lung),
				EL661102 (small intestine)
867-4	127833-188870	460533	10	CX842746 (brain), AAI68128 (testes)
867-5	196341-211233	187171	6	DR836487 (fat body), DR834196 (fat body), CX999510 (heart),
				EG653999 (small intestine), DN015523 (heart), CX912262 (testes),
				DN059149 (fat body), etc.
867-6	265545-280100	187173	3	EG653999 (small intestine), CX912262 (testes), DN015523 (heart)
867-7	284326-296678	187174	3	DR836487 (fat body), DR834196 (fat body)
867-8	296678-312149	187174	3	CX924660 (testes), CX920068 (testes), CX915279 (testes), CX811079 (brain)
867-9	313339-322098	187174	2	CX903340 (testes), CX807560 (brain), CX886240 (testes), CX924661 (testes)
867-10	332423-336387	460534	1	CX924660 (testes), CX920068 (testes), CX915279 (testes),
				EL655892 (small intestine), CX811079 (brain)

Scaffold-3534 ((11,245bp)
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1707-2

20880-21915

191590

Scaffold-3	534 (11,245bp)			
3534-1	1894-8537	192957	3	CX828489 (brain)
Scaffold-1	318 (98,102bp)			
1318-1	10875-14000	190460	1	CX995964 (intestine), CX983522 (intestine), CX979602 (intestine),
				CX975809 (intestine), CX979601 (intestine), CX983521 (intestine),
				CX995963 (intestine), CX975808 (intestine)
1318-2	14000-19000	190460	2	all ESTs in 1318-1
				EL699109 (limb), DN077598 (lung), DN077598 (lung), CX854937 (brain)
1318-3	19000-34186	190460	7	EL659466, EL653181, EL663470 (small intestine)
1318-4	46075-70472	190461	5	all ESTs in 867-3; EG653270 (small intestine)
	•			
Scaffold-1	.146 (154,973bp)			
1146-1	10125-12391	257547	3	DR879120 (skin), BX691942 (neurula), AL788752 (neurula)
1146-2	64158-76640	461287	3	DT421997 (skin), CF590382 (whole body)
1146-3	84381-101615	189674	11	CX888371 (testes), CX899439 (testes), CX812473 (brain), CX820566 (brain)
				CX820565 (brain), CX905130 (testes), CX802601 (brain),
				EL655865 (small intestine), EL655864 (small intestine),
				EG651995 (small intestine), EL663892 (small intestine), CX977562 (intestine),
				DT401525 (oviduct), EL661154 (small intestine), CX978268 (intestine),
				CX978268 (intestine), EL661154 (small intestine), CX810609 (brain),
				EG650864(small intestine), CX974210 (intestine), EL663892(small intestine),
				EL653717 (small intestine), CX978267 (intestine), etc.
1146-4	148521-151681	189677	1	DN036288 (liver), DC144455 (embryo), DR879119 (skin), DT421998 (skin),
				DR879120 (skin) 97%, CF590382 (whole body) 94%
Scaffold-1	707 (36,085bp)			
1707-1	7687-19898	191589	11	EL661154 (small intestine), CX978268 (intestine), EG651995 (intestine),
				CX900915 (testes), CX810609 (brain), CX974210 (intestine),
				EL663892 (small intestine), EL653717 (intestine), CX978267 (intestine),
				EL707062 (limb), CX905615 (testes), CX888370 (testes), CX900914 (testes),
				EL707061 (limb), CX812472 (brain),

no EST

scaffold-16	557 (40,156bp)			
1657-1	13450-29812	191506	3	CX916757 (testes), DN036288 (liver), DC144455 (embryo), DR879119 (skin), DT421998 (skin), DR879120 (skin), CF590382 (whole body), CX979669 (intestine)
Scaffold-22	250 (20,541bp)			
2250-1	413-9768	192194	2	DN036288 (liver), DC144455 (embryo), DR879119 (skin), DT421998 (skin), DR879120 (skin), CF590382 (whole body), CX979669 (intestine), CX916757 (testes)
Scaffold-14	105 (75,973 bp)			
1405-1	9908-23896	190816	6	Same as 1318-2: EL653181 (small intestine), EL659466 (small intestine), EL663470 (small intestine), CX979602 (intestine), CX975809 (intestine), CX995963 (intestine), CX983521 (intestine), CX983522 (intestine), CX995964 (intestine), CX979601 (intestine)
1405-2	66602-75451	472810	2	EL699109 (limb), CX854937 (brain), CX854938 (brain), EL661102 (small intestine), DN077598 (lung), EG653270 (small intestine)
Scaffold-37	782 (10,288 bp)			<u> </u>
3782-1	4808-9885	193094	1	DR834196 (fat body), CX999510 (heart), DR836487 (fat body), EL674099 (thymus)
Scaffold-47	710 (8,229 bp)			
4710-1	2470-7801	193353	1	CX828489 (brain), CX923462 (testes)

¹ 'etc.' indicates there are many more sequences with lower than 90% nucleotide identities.

Online resource 3. Genomic coordinates of NKp30 in the Xenopus scaffolds

Scaffold-215: 1,772,981 bp

	Position (bp)	gene symbol	Protein ID	human chr	Type ¹	Gene description/EST for NKp30
215-1	29895-31362	NKp30	170752	6p21.3	DT	88% DT447366 (spleen)
215-2	36219-41502	NKp30	170753	6p21.3	ES	100% ES685492 (spleen); ES685490 (eye); DR850889 (ovary); DN098981 (Ovary); CX747707 (whole embryo);
						AL878396 (egg); AL961664 (gastrula); DR850888 (ovary); CX747706 (whole embryo);
						98% CX966132 (intestine); 95% CX445352 (whole embryo);
215-3	54224-56880	NKp30	170755	6p21.3	DT	95% DT447366 (spleen), 96% EL837225 (testes)
215-4	62790-67524	NKp30*	170756	6p21.3	DT, DT	91-6% DT447366 (spleen), 95-7% EL837225 (testes)
215-5	73213-83037	NKp30*	170757	6p21.3	DT, DT	91-2% DT447366 (spleen), 97-8% EL837225 (testes)
215-6	90642-97175	NKp30	170758	6p21.3	DT	95% DT447366 (spleen), 99% EL837225 (testes)
215-7	131570-137052	NKp30*	170763	6p21.3	DT, DT	92-3% DT447366 (spleen), 97-100% EL837225 (testes)
215-8	168948-174567	NKp30	170766	6p21.3	DT	91% DT447366(spleen), 99% EL837225 (testes)
215-9	205264-205539	NKp30	138657	6p21.3	DT	86% DT447366 (spleen)
215-10	209299-221397	NKp30	170769	6p21.3	DT	98% EL837335 (testes), 91% DT447366 (spleen)
215-11	224116-248685	NKp30	170770	6p21.3	DT	99% DT447366 (spleen)
215-12	269377-275917	NKp30	466773	6p21.3	ВХ	100% match to BX773925 (egg)
13	283772-288419	B3GAT1-like	350405	11q25		beta-1,3-glucuronyltransferase 1
14	432001-504608	BSN	170777	3p21.31		bassoon (presynaptic cytomatrix protein)
15	526523-528650	BSN-like	466774			protein basoon-like
16	855898-891267	DAG1	170780	3p21.31		dystroglycan 1 (dystrophin-associated glycoprotein 1)
17	952064-980011	USP4	170783	3p21.3		ubiquitin specific peptidase 4 (proto-oncogene)
18	995022-999682	GPX1	454592	3p21.3		glutathione peroxidase 1
19	1004872-1020027	RHOA	149865	3p21.3		ras homolog gene family, member A
20	1163158-1174921	GMPPB	454601	3p21.31		GDP-mannose pyrophosphorylase B
21	1180654-1198431	IP6K1	170800	3p21.31		inositol hexakisphosphate kinase 1
22	1268580-1287897	CDHR4	170804	3p21.31		cadherin-related family member 4
23	1296340-1299408	TLR9	350411	3p21.3		toll-like receptor 9
24	1303105-1313373	TRAIP	350388	3p21.31		TRAF interacting protein
25	1385332-1390848	C3orf54	293751	3p21.31		

Scaffold-88: 3,020,139bp

	Position (bp)	gene symbol	Protein ID	human chr	Type ¹	Gene description/EST for NKp30
1	55318-84426	TNRC6B	163058	22q13.1		trinucleotide repeat containing 6B
2	107991-116671	ADSL	451771	22q13.2		adenylosuccinate lyase
3	123789-145333	SGSM3	271451	22q13.1-q13.2		small G protein signaling modulator 3
4	149696-155729	CHAD	271472	17q21.33		chondroadherin
5	156637-167119	RANGAP1	163063	22q13		Ran GTPase activating protein 1
6	173261-202155	ZC3H7B	271466	22q13.2		zinc finger CCCH-type containing 7B
7	217888-223875	TEF	208802	22q13.2		thyrotrophic embryonic factor
8	244085-246726	PKN2	327775	1p22.2		protein kinase N2
9	280977-293299	GALR2	327802	17q25.3		galanin receptor 2
10	349153-400242	TOM1	208803	22q13.1		target of myb1
11	402220-412313	HMGXB4	163074	22q13.1		HMGXB4 and Name: HMG box domain containing 4
12	418993-426545	ANKRD54	327650	22q13.1		ankyrin repeat domain 54
13	431466-441809	EIF3L	451784	22q		eukaryotic translation initiation factor 3, subunit L
14	443205-486752	MCALL1	451786	22q13.1		MICAL-like 1
15	492687-499703	C22orf23	292767	22q13.1		chromosome 22 open reading frame 23; EVG1
16	499702-504092	POLR2F	271486	22q13.1		polymerase (RNA) II (DNA directed) polypeptide F
17	512058-518986	SOX10	451789	22q13.1		SRY (sex determining region Y)-box 10
18	559579-580477	BAIAP2L2	327814	22q13.1		BAI1-associated protein 2-like 2
19	589335-609781	PLA2G6	451790	22q13.1		phospholipase A2, group VI
20	621288-626606	MAFF	163085	22q13.1		v-maf musculoaponeurotic fibrosarcoma oncogene homolog F
21	638353-678699	TMEM184B	163086	22q12		transmembrane protein 184B
22	692261-732951	CSNK1E	163088	22q13.1		casein kinase 1, epsilon
23	755336-756936	KCNJ4	451791	22q13.1		potassium inwardly-rectifying channel, subfamily J, member 4
24	811924-817996	KDELR3	148351	22q13.1		KDEL (Lys-Asp-Glu-Leu) endoplasmic reticulum protein retention receptor 3
25	845791-849259	TNIK	163095	3p26.31		TRAF2 and NCK interacting kinase
26	853890-854186	nog4	18973	17q22 paralog?		noggin4
27	855476-871597	GCAT	327659	22q13.1		glycine C-acetyltransferase
28	878989-879330	H1F0	109293	22q13.1		H1 histone family, member 0
29	897679-908475	TRIOBP	327579	22q13.1		TRIO and F-actin binding protein
30	922457-943491	SRRM2?	163100	16p13.3		serine/arginine repetitive matrix 2
31	944169-954045	NOL12	148353	22q13.1		nucleolar protein 12
32	954084-972241	ADAP1	225007	7p22.3		ArfGAP with dual PH domains 1

Online resource 3. Genomic coordinates of NKp30 in the Xenopus scaffolds

33	979371-987199	LGALS1	148356	22q13.1	lectin, galactoside-binding, soluble, 1
	994368-1009091	LGALS1	148357	22q13.1	lectin, galactoside-binding, soluble, 1
		LGALS1	292772	22q13.1	lectin, galactoside-binding, soluble, 1
		LGALS1	163108	22q13.1	lectin, galactoside-binding, soluble, 1
37	1068659-1080149	SH3BP1	327728	22q13.1	SH3-domain binding protein 1
	1108125-1124159		327829	22q13.31	golgi-associated, gamma adaptin ear containing, ARF binding protein 1
39	1136061-1138851	LGALS2	163110	22q13.1	lectin, galactoside-binding, soluble, 2
40	1269635-1272443	ELFN2	451800	22q13.1	extracellular leucine-rich repeat and fibronectin type III domain containing 2
41	1314787-1336565	CYTH2	327670	19q13.33	cytohesin 2
42	1358871-1384147	RAC2	475981	22q13.1	ras-related C3 botulinum toxin substrate 2
43	1405028-1406239	SSTR3	327701	22q13.1	somatostatin receptor 3
44	1470189-1495484	TMPRSS6	464718	22q12.3	transmembrane protease, serine 6
45	1498089-1503584	KCTD17	14752	22q12.3	potassium channel tetramerisation domain containing 17
46	1505535-1512725	MPST	163119	22q13.1	mercaptopyruvate sulfurtransferase
47	1556398-1564872	TST	163123	22q13.1	thiosulfate sulfurtransferase (rhodanese)
48	1578379-1605830	CSF2RB	451803	22q13.1	colony stimulating factor 2 receptor, beta, low-affinity
49	1615529-1638662	NCF4	148362	22q13.1	neutrophil cytosolic factor 4, 40kDa
50	1658714-1677150	PVALV	148364	22q12-q13.1	parvalbumin
51	1688532-1706043	IFT27	451804	22q13.1	intraflagellar transport 27 homolog (Chlamydomonas)
52	1729296-1782366	CACNG2	271449	22q13.1	calcium channel, voltage-dependent, gamma subunit 2
53	1809705-1822472	EIF3D	148366	22q13.1	eukaryotic translation initiation factor 3, subunit D
54	1825333-1831291	MUSTN1	451808	3	musculoskeletal, embryonic nuclear protein 1
55	1840257-1853219	TMEM110-MUS	199241	3	TMEM110-MUSTN1 readthrough
56	1860469-1891127	SFMBT1	327822	3p21.1	Scm-like with four mbt domains 1
57	1939871-1956718	RFT1	475987	3p21.1	RFT1 homolog (S. cerevisiae)
58	2020934-2033178	PRKCD	271492	3p21.31	protein kinase C, delta
59	2034703-2055023	PFKFB4	451813	3p22-p21	6-phosphofructo-2-kinase/fructose-2,6-biphosphatase 4
60	2118529-2130605	WASL	451814	7q31.3	Wiskott-Aldrich syndrome-like
61	2158165-2175889	ATRIP	451816	3p21.31	ATR interacting protein
62	2185665-2188143	BTG1	292779	12q22	B-cell translocation gene 1, anti-proliferative
63	2235401-2260694	IPPK	271516	9q22.31	inositol 1,3,4,5,6-pentakisphosphate 2-kinase
64	2289075-2302416	ECM2	163153	9q22.3	extracellular matrix protein 2, female organ and adipocyte specific
65	2321556-2332848	ASPN	327585	9q22	asporin
66	2350549-2353158	OMD	199249	9q22.31	osteomodulin

Online resource 3. Genomic coordinates of NKp30 in the Xenopus scaffolds

67	2371700-2375287	OGN	327661	9q22		osteoglycin
68	2403512-2429465	NOL8	512865	9q22.31		nucleolar protein 8
69	2462656-2511391	IARS	451824	9q21		isoleucyl-tRNA synthetase
70	2761382-2851564	ATP2B2	199252	3p25.3		ATPase, Ca++ transporting, plasma membrane 2
71	2872484-2911770	FANCD2	327776	3p26		Fanconi anemia, complementation group D2
72	2914797-2915547	C3orf24	271423	3p25.3		
73	2917929-2926887	TMEM111	148375	3		transmembrane protein 111
88-2	2955788-2958378	NKp30	163171	6p21.3	New (88-2)	no match
88-1	3005933-3014759	NKp30	163176	6p21.3	ES	91% ES685492 (spleen); ES685490 (eye); DR850889 (ovary); DN098981 (Ovary); CX747707 (whole embryo);
						89% CX445352 (whole embryo); CX966132 (intestine); AL878396 (egg); 93% CX747706 (whole embryo);
						92% DR850888 (ovary); AL961664 (gastrula);

Scaffold-3675: 10,658bp

	Position (bp)	gene symbol	Protein ID	human chr	Type ¹	Gene description/EST for NKp30
3675-1	2198-4041	NKp30	193035	6p21.3	DT	90% DT447366 (spleen)

Scaffold-15172: 3,739bp

15172-1	40-595	NKp30	195782	6p21.3	ES	98% ES685492 (eye); 96% ES685490 (eye), DR850889 (ovary), DN098981 (ovary),
						CX747707 (whole embryo), AL878396 (egg); 94% CX747706 (whole embryo), CX966132 (intestine);
						93% CX445352 (whole embryo), DR850888 (Ovary)

Scaffold-3771: 10,314bp

3771-1	683-2204	NKp30	193089	6p21.3	ES	95% ES685492 (eye), ES685490 (eye), DR850889 (ovary), DN098981 (ovary), CX747707 (whole embryo),
						AL878396 (egg), CX445352 (whole embryo), CX966132 (intestine); 94% C7447706 (whole embryo);
						93% DR850888 (ovary), AL961664 (gastrula)

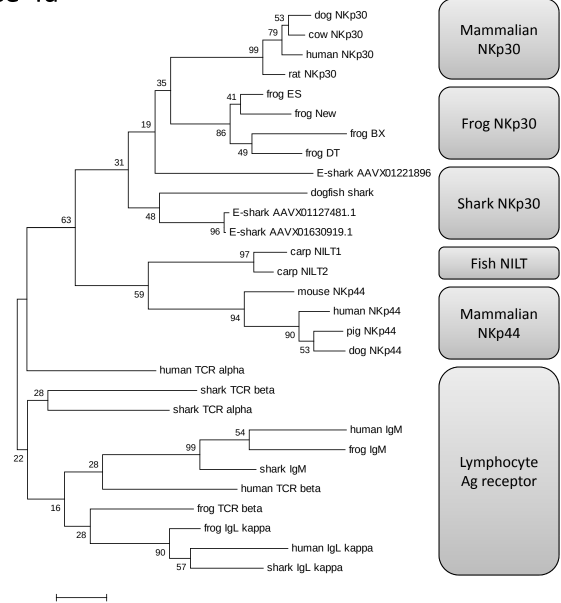
^{*} indicate these gene models contain two IgSF exons, presumably assembly error.

¹: Nkp30 types based on the amino acid sequences

Online Resource 4a

ML tree

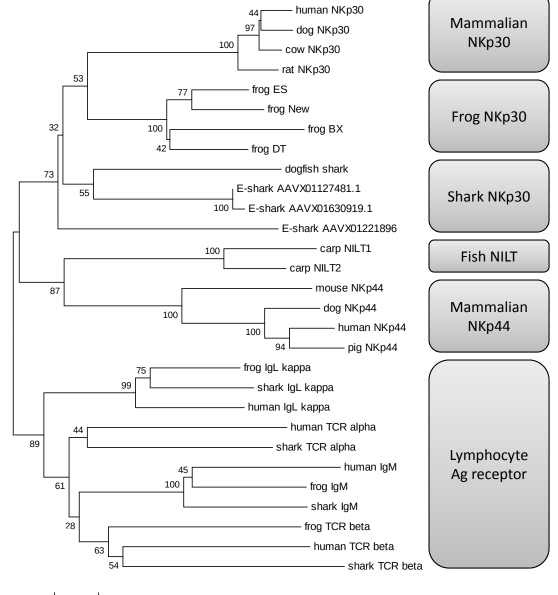
Using nucleotide sequences of V domains



Online Resource 4b

ME tree

Using nucleotide sequences of V domains



Online Resource 5.

a) Divergent V-IgSF domains of Elephant shark B7H6 genes

```
AAVX01166257.1
                    DALEMTIFKDQLEVFQGQDVVLDCKVTGYTPAVLNLSFVAVEWRFVS----TQQLVYSFN 56
AAVX01015110.1
AAVX01166967.1
                    S....VL.HSPMKGQVNET.K.E...S..SSPQ.Q.KT.G.Q.I.T.----.RKE...YI 56
                    .V.NVLMDSGPVR.AVNK.TL.K.Q.P..ESRTID.AS.S...Y.SKPPDV.RK...K.V 60
AAVX01076204.1
                    -.AR..LRTSPA.CLVNE..S.Q.EIADL.GIT.DKNHLG.L.S.TD-----DIFKYI 52
AAVX01166257.1 GGFIDALRPGATVTIENLKRGDASLHLPKVQIEEEGEYKCIVYVTP-SKAEKSSSVRVL 114
AAVX01015110.1 ..KAS.S.N...ISED..RT.....Y..NI...D..D.T.V.I...-E.V..T.TLQ.S 114
AAVX01166967.1 NSAHTPH.K..KISDDK..K.I...F.LS.GFQD..Q.E.V.FI..-DSGRGNAEMF.G 118
AAVX01076204.1 N.ALFK.GK.VELLESSILK.NM..L.KN.TMKQM....T.F.P.NEQ.TGTVRLE.- 110
b) Divergent V-IgSF domains of Elephant shark NKp30 genes
                    ----VVQSPSRVEAAEGERVSLSCTYTGDVGSGIGGYQWLKHS--PHGVPVSNDTEGYRG 54
AAVX01127481
AAVX01545093.1
                   ---- 53
AAVX01630919
                    ----. H....-. 53
AAVX01169427
                    ----.T..SLN.T..QT.L.N..FD.PS--TN..FS.VRDLGGERERE...T.GP.E. 52
AAVX01221896
                    DPPG.T.W.PAIQ.LS.QT.T.N.SFAAR---.VTVKT...DG----AALDLSSPR.S. 52
AAVX01545093.1
                    RVHRKHGRSFIETRDASIAISDVRQSDSGVYYCAIDLMGDVRGYGSGTTLT---- 105
                   .....QN..G......N.....EA------ 94
                    AAVX01630919
AAVX01169427
AAVX01169427
AAVX01221896
                    ...ISRSSAE.KTLK....S.R.LTGR...L...RVE.VSQTESV.P..R.QVV--- 105
                    ..AQADAQT.RARG..T.HL.NLSVC...R.V.RVQPL.SAAAL.R..QV.VTRES 108
```

Figure legend:

Deduced amino acid alignment of V domains screened from E-shark genome using ClustalW2. Gaps and Identical aa residues are indicated with dashes and dots, respectively. Canonical amino acids characteristic for IgSF domains are highlighted in grey. a) B7H6, b) NKp30.