



Review

IMGT-ONTOLOGY and IMGT databases, tools and Web resources for immunogenetics and immunoinformaticsMarie-Paule Lefranc^{a,b,*}^a *Laboratoire d'ImmunoGénétique Moléculaire, LIGM, Institut de Génétique Humaine IGH, Université Montpellier II, UPR CNRS 1142, 141 rue de la Cardonille, 34396 Montpellier Cedex 5, France*^b *Institut Universitaire de France, France*

Received 18 June 2003; received in revised form 2 September 2003; accepted 16 September 2003

Abstract

The international ImMunoGeneTics information system® (IMGT; <http://imgt.cines.fr>), is a high quality integrated information system specialized in immunoglobulins (IG), T cell receptors (TR), major histocompatibility complex (MHC), and related proteins of the immune system (RPI) of human and other vertebrates, created in 1989, by the Laboratoire d'ImmunoGénétique Moléculaire (LIGM; Université Montpellier II and CNRS) at Montpellier, France. IMGT provides a common access to standardized data which include nucleotide and protein sequences, oligonucleotide primers, gene maps, genetic polymorphisms, specificities, 2D and 3D structures. IMGT consists of several sequence databases (IMGT/LIGM-DB, IMGT/MHC-DB, IMGT/PRIMER-DB), one genome database (IMGT/GENE-DB) and one 3D structure database (IMGT/3Dstructure-DB), interactive tools for sequence analysis (IMGT/V-QUEST, IMGT/JunctionAnalysis, IMGT/PhyloGene, IMGT/Allele-Align), for genome analysis (IMGT/GeneSearch, IMGT/GeneView, IMGT/LocusView) and for 3D structure analysis (IMGT/StructuralQuery), and Web resources ("IMGT Marie-Paule page") comprising 8000 HTML pages. IMGT other accesses include SRS, FTP, search by BLAST, etc. By its high quality and its easy data distribution, IMGT has important implications in medical research (repertoire in autoimmune diseases, AIDS, leukemias, lymphomas, myelomas), veterinary research, genome diversity and genome evolution studies of the adaptive immune responses, biotechnology related to antibody engineering (single chain Fragment variable (scFv), phage displays, combinatorial libraries) and therapeutical approaches (grafts, immunotherapy). IMGT is freely available at <http://imgt.cines.fr>.

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Keywords: IMGT; Database; Immunoglobulin; T cell receptor; MHC; HLA; Immunogenetics; 3D structure; Web resources; Leukemia; Lymphoma**1. Introduction**

The molecular synthesis and genetics of the immunoglobulin (IG) and T cell receptor (TR) chains is particularly complex and unique as it includes biological mechanisms such as DNA molecular rearrangements in multiple loci (three for IG and four for TR in humans) located on different chromosomes (four in humans), nucleotide deletions and insertions at the rearrangement junctions (or N-diversity), and somatic hypermutations in the IG loci (for review Lefranc and Lefranc, 2001a,b). The number of potential protein forms of IG and TR is almost unlimited. Owing to the complexity and high number of published sequences, data control and classification and

detailed annotations are a very difficult task for the generalist databanks such as EMBL (UK) (Stoesser et al., 2003), GenBank (USA) (Benson et al., 2003), and DDBJ (Japan) (Miyazaki et al., 2003). These observations were the starting point of the international ImMunoGeneTics information system® (IMGT; <http://imgt.cines.fr>) (Lefranc, 2003a), created in 1989, by the Laboratoire d'ImmunoGénétique Moléculaire (LIGM; Université Montpellier II and CNRS) at Montpellier, France. IMGT is a high quality integrated information system specialized in IG, TR, major histocompatibility complex (MHC) and related proteins of the immune system (RPI) of human and other vertebrate species (Giudicelli et al., 1997; Lefranc et al., 1998, 1999; Ruiz et al., 2000; Lefranc, 2001a, 2002, 2003a,b). IMGT consists of several sequence databases (IMGT/LIGM-DB, IMGT/MHC-DB, IMGT/PRIMER-DB, IMGT/PROTEIN-DB, this last one in development), one genome database (IMGT/GENE-DB)

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URL: <http://imgt.cines.fr>.

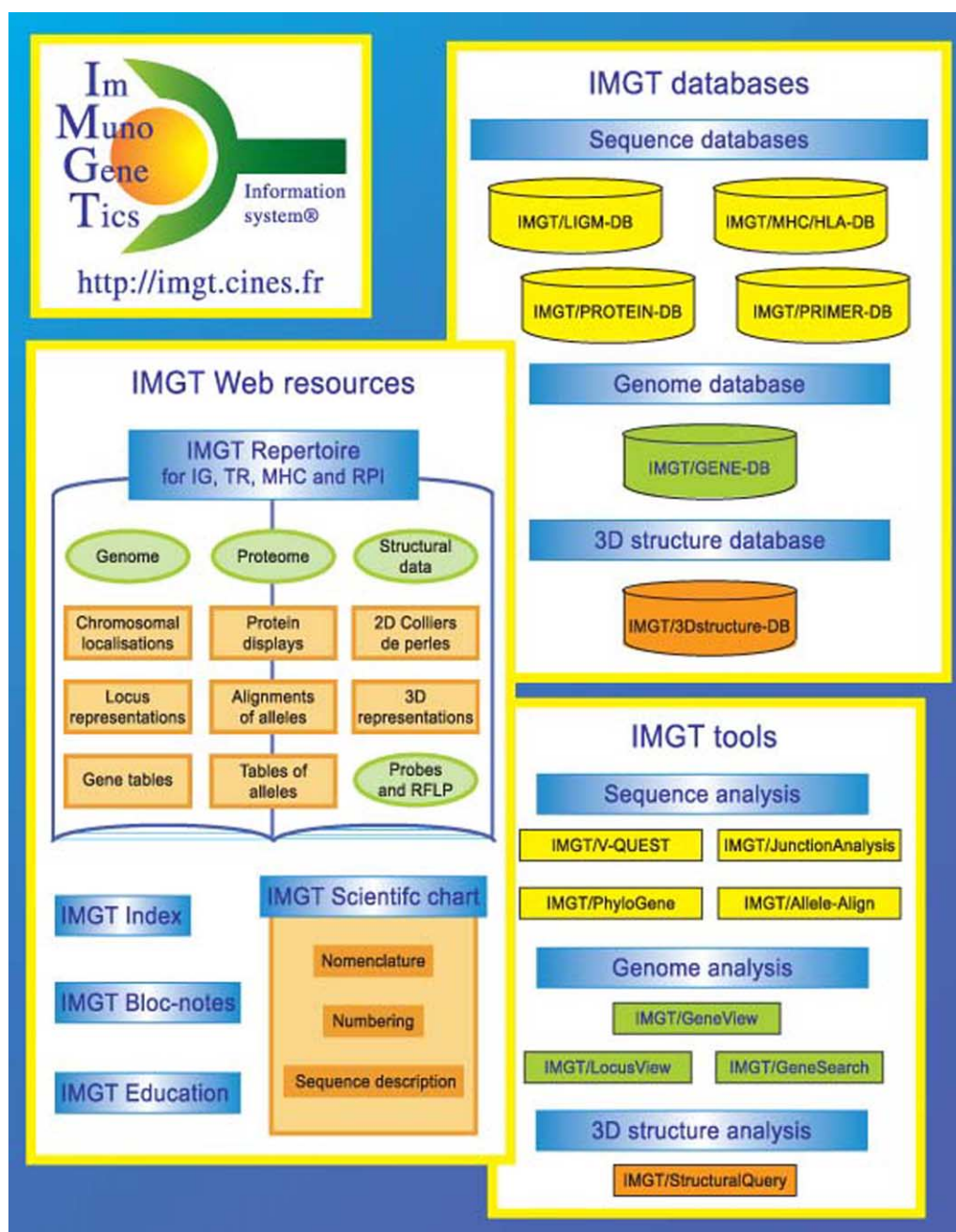


Fig. 1. Databases, interactive tools and Web resources of IMGT, the international ImMunoGeneTics information system® (<http://imgt.cines.fr>).

and one three-dimensional 3D structure database (IMGT/3Dstructure-DB), interactive tools for sequence analysis (IMGT/V-QUEST, IMGT/JunctionAnalysis, IMGT/PhyloGene, IMGT/Allele-Align), for genome analysis (IMGT/GeneSearch, IMGT/GeneView, IMGT/LocusView) and 3D structure analysis (IMGT/StructuralQuery), and Web resources ("IMGT Marie-Paule page") comprising 8000 HTML pages which include IMGT Scientific chart, IMGT Repertoire (for IG and TR, MHC, RPI), IMGT Bloc-notes, IMGT Education and IMGT Index (Fig. 1). IMGT other accesses include SRS, FTP, search by BLAST,

etc. By its high quality and its easy data distribution, IMGT has important implications in medical research (repertoire in normal and pathological situations: autoimmune diseases, infectious diseases, AIDS, detection of residual diseases in leukemias, lymphomas, myelomas), veterinary research, genome diversity and genome evolution studies of the adaptive immune responses, biotechnology related to antibody engineering (single chain Fragment variable (scFv), phage displays, combinatorial libraries) and therapeutic approaches (grafts, immunotherapy). IMGT is freely available at <http://imgt.cines.fr>.

2. IMGT-ONTOLOGY

IMGT has developed a formal specification of the terms to be used in the domain of immunogenetics and immunoinformatics to ensure accuracy, consistency and coherence in IMGT. This has been the basis of IMGT-ONTOLOGY (Giudicelli and Lefranc, 1999), the first ontology in the domain, which allows the management of the immunogenetics knowledge for human and other vertebrate species. IMGT-ONTOLOGY comprises five main concepts: IDENTIFICATION, CLASSIFICATION, DESCRIPTION, NUMEROTATION and OBTENTION (Giudicelli and Lefranc, 1999). Standardized keywords, standardized sequence annotation, standardized IG and TR gene nomenclature, the IMGT unique numbering, and standardized origin/methodology were defined, respectively, based on these five main concepts. The controlled vocabulary and the annotation rules for data and knowledge management of the IG, TR, MHC and RPI of human and other vertebrate species constitute the IMGT Scientific chart. All IMGT data are expertly annotated according to the IMGT Scientific chart. IMGT is the global internationally acknowledged reference in immunogenetics and immunoinformatics.

IMGT standardized keywords, based on the IDENTIFICATION concept, have been assigned to all IMGT/LIGM-DB entries. They include (i) *general keywords*: indispensable for the sequence assignments, they are described in an exhaustive and non-redundant list, and are organized in a tree structure, and (ii) *specific keywords*: they are more specifically associated with particularities of the sequences (orphan, transgene, etc.) or with diseases (leukemia, lymphoma, myeloma, etc.) (Giudicelli et al., 1997). The list is not definitive and new specific keywords can easily be added if needed.

Two hundred fifteen feature labels, based on the DESCRIPTION concept, are necessary in IMGT/LIGM-DB to describe all structural and functional subregions that compose IG and TR sequences (Giudicelli et al., 1997), whereas only seven of them are available in EMBL, GenBank or DDBJ. Annotation of sequences with these labels constitutes the main part of the expertise. Levels of annotation have been defined, which allow the users to query sequences in IMGT/LIGM-DB even though they are not fully annotated (Giudicelli et al., 1997). An internal tool, IMGT/Automat, has been developed to automatically perform the annotation of the rearranged cDNA sequences in IMGT/LIGM-DB (Giudicelli et al., 2003). One hundred seventy two additional labels were defined for IG, TR, MHC, and RPI amino acid sequences and domain structures in IMGT/PROTEIN-DB and IMGT/3Dstructure-DB. Prototypes represent the organizational relationship between labels and give information on the order and expected length (in number of nucleotides) of the labels (Giudicelli et al., 1997; Lefranc et al., 1999). Prototypes can apply to general configuration of IG, TR or MHC, independently of the chain type, the species or any other parameters like functionality. However, prototypes

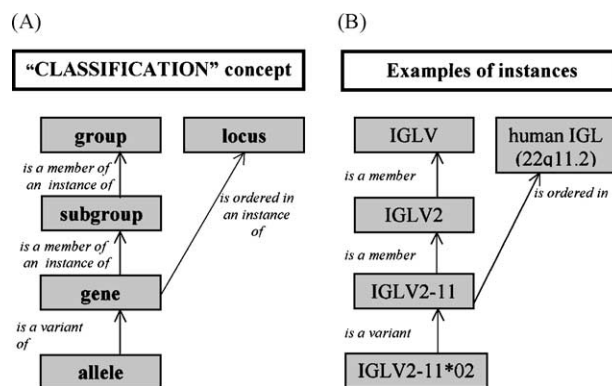


Fig. 2. The IMGT-ONTOLOGY CLASSIFICATION concept. The CLASSIFICATION concept organizes the immunogenetics knowledge useful to name and classify IG and TR genes in IMGT. (A) Concepts of classification and their relations. (B) Examples of instances: for each concept of classification is shown the instance which allows the classification of the human IGLV2-1 1*02 allele.

may also be established for very precise cases when sequence characteristics are clearly established.

IMGT provides immunologists and geneticists with a standardized nomenclature per locus and per species which allows extraction and comparison of data for the complex B and T cell antigen receptors. The CLASSIFICATION concept (Fig. 2) has been used to set up a unique nomenclature of the human IG and TR genes, which was approved by the Human Genome Organization (HUGO) Nomenclature Committee (HGNC) in 1999 (Wain et al., 2002a) and has become the community standard. The complete list of the human IG and TR gene names (Lefranc, 2000a,b,c,d; Lefranc, 2001b,c,d; Lefranc and Lefranc, 2001a,b) has been entered by the IMGT Nomenclature Committee in the genome database (GDB, Canada) (Letovsky et al., 1998), LocusLink at NCBI (USA) (Pruitt and Maglott, 2001), and GeneCards (Safran et al., 2002). The complete list of the mouse IG and TR gene names was sent by IMGT, in July 2002, to the mouse genome informatics (MGI) mouse genome database (MGD) (USA) (Blake et al., 2003), LocusLink at NCBI, and HGNC. Both lists are available from the IMGT site (Lefranc, 2003a) and queries on the human and mouse IG and TR gene classification and gene names can be made from IMGT/GENE-DB. IMGT reference sequences have been defined for each allele of each gene based on one or, whenever possible, several of the following criteria: germline sequence, first sequence published, longest sequence, mapped sequence (Lefranc et al., 1999).

A uniform numbering system for IG and TR sequences of all species, based on the NUMEROTATION concept, has been established to facilitate sequence comparison and cross-referencing between experiments from different laboratories whatever the antigen receptor (IG or TR), the chain type, the domain (variable V or constant C), or the species (Lefranc, 1997, 1999; Lefranc et al., 2003). This numbering results from the analysis of more than 5000 IG and TR

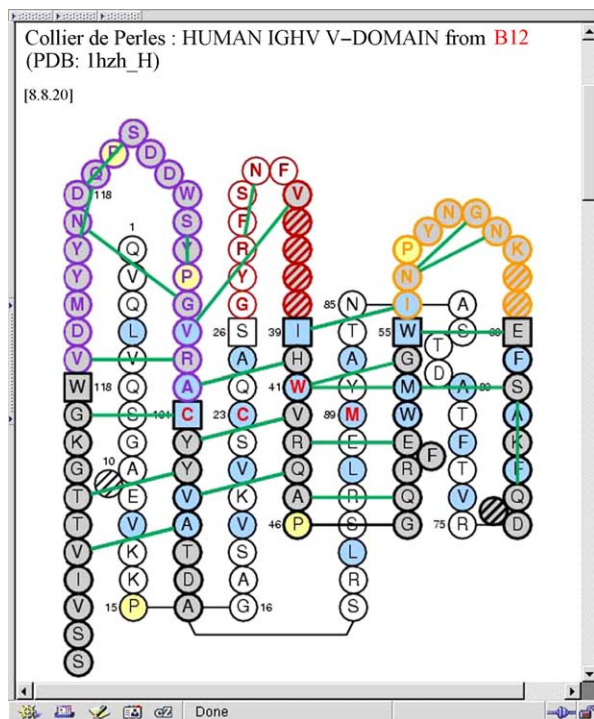


Fig. 3. Example of IMGT Collier de Perles on two layers. The IMGT Collier de Perles on two layers are generated dynamically in IMGT/3Dstructure-DB (<http://imgt.cines.fr>). CDR1-IMGT, CDR2-IMGT and CDR3-IMGT are in red, orange and purple, respectively. The outer A B D E beta strands (from left to right) are in white, whereas the inner G F C C' C'' beta strands (from left to right) are in grey. Hydrogen bonds between amino acids of the inner G F C C' C'' sheet, deduced from the PDB X-ray crystallographic data, are shown in green. Hatched circles correspond to missing positions according to the IMGT unique numbering. The disulfide bond between 1st-CYS 23 and 2nd-CYS 104 is not shown.

variable region sequences of vertebrate species from fish to human. It takes into account and combines the definition of the framework (FR) and complementarity determining region (CDR) (Kabat et al., 1991), structural data from X-ray diffraction studies (Satow et al., 1986) and the characterization of the hypervariable loops (Chothia and Lesk, 1987). In the IMGT numbering, conserved amino acids from frameworks always have the same number whatever the IG or TR variable sequence, and whatever the species they come from (as examples: cysteine 23 (in FR1), tryptophan 41 (in FR2), leucine 89 and cysteine 104 (in FR3). Based on the IMGT unique numbering, standardized 2D graphical representations, designated as IMGT Colliers de Perles (Lefranc et al., 1999) (Fig. 3), are available in IMGT Repertoire (<http://imgt.cines.fr>). This IMGT unique numbering has several advantages:

- It has allowed the redefinition of the limits of the FR and CDR of the IG and TR variable regions (Lefranc and Lefranc, 2001a,b) and domains (Ruiz and Lefranc, 2002). The FR-IMGT and CDR-IMGT lengths become in themselves crucial information which characterize variable regions belonging to a group, a subgroup and/or a gene.

- Framework amino acids (and codons) located at the same position in different sequences can be compared without requiring sequence alignments. This also holds for amino acids belonging to CDR-IMGT of same length.
- The unique numbering is used as the output of the IMGT/V-QUEST alignment tool. The aligned sequences are displayed according to the IMGT numbering and with the FR-IMGT and CDR-IMGT delimitations.
- The unique numbering has allowed a standardization of the description of mutations and the description of IG and TR allele polymorphisms (Lefranc and Lefranc, 2001a,b). These mutations and allelic polymorphisms are described by comparison to the IMGT reference sequences of the alleles *01 (Lefranc, 1998).
- The unique numbering allows the description and comparison of somatic hypermutations of the IG IMGT variable domains.

By facilitating the comparison between sequences and by allowing the description of alleles and mutations, the IMGT unique numbering represents a big step forward in the analysis of the IG and TR variable region (V-REGION) sequences of all vertebrate species (Pommié et al., *in press*). Moreover, it gives insight into the structural configuration of the variable domain (V-DOMAIN encoded by the V–J- or V–D–J-REGION) (Ruiz and Lefranc, 2002; Lefranc et al., 2003) (Fig. 3). The IMGT unique numbering opens interesting views on the evolution of the proteins belonging to the “immunoglobulin superfamily” (IgSF) (Williams and Barclay, 1988). It has been applied with success to all the sequences of domains belonging to the IgSF V-set, designated as V-LIKE-DOMAINS in IMGT, which include non-rearranging sequences in vertebrates (human CD4 [D1,D3], *Xenopus* CTXg1, etc.) and in invertebrates (*drosophila* amalgam, *drosophila* fasciclin II, etc.) (IMGT Repertoire (RPI), <http://imgt.cines.fr>) (Lefranc, 1997, 1999; Lefranc et al., 2003).

This standardized approach has been applied to the constant domain (C-DOMAIN) of the IG and TR, and to the C-LIKE-DOMAINS of proteins other than IG and TR (IMGT Repertoire (RPI), <http://imgt.cines.fr>). An IMGT unique numbering has also been implemented for the groove domain (G-DOMAIN) (Duprat and Lefranc, 2003) of the MHC class I and II chains (IMGT Repertoire (MHC), <http://imgt.cines.fr>), and for the G-LIKE-DOMAINS of proteins other than MHC (IMGT Repertoire (RPI), <http://imgt.cines.fr>).

The OBTENTION concept is a set of standardized terms that precise the origins of the sequence (the ‘origin’ concept) and the conditions in which the sequences have been obtained (the ‘methodology’ concept). The ‘origin’ concept comprises the subsets of ‘cell, tissue or organ’, ‘auto-immune diseases’, ‘clonal expansion diseases’ (such as leukemia, lymphoma, myeloma), whereas the ‘methodology’ concept comprises the subsets related to the ‘hybridoma’, to the experimental conditions (sequences

amplified by ‘PCR’), to the obtention from ‘libraries’ (genomic, cDNA, combinatorial, etc.) or from ‘transgenic’ organisms (animal, plant). At this stage of development, the exhaustive definition of the concepts of obtention and of their instances is still in progress.

3. IMGT databases

IMGT/LIGM-DB contains three sequence databases, IMGT/LIGM-DB, IMGT/MHC-DB, and IMGT/PRIMER-DB, one genome database, IMGT/GENE-DB, and one 3D structure database, IMGT/3Dstructure-DB. IMGT/LIGM-DB, the first and the largest database, is the comprehensive IMGT database of IG and TR nucleotide sequences from human and other vertebrate species, with translation for fully annotated sequences, created in 1989 by LIGM, at Montpellier, France, on the Web since July 1995 (Giudicelli et al., 1997; Lefranc et al., 1998, 1999; Ruiz et al., 2000; Lefranc, 2001a, 2002, 2003a,b). In October 2003, IMGT/LIGM-DB contained 77,810 nucleotide sequences of IG and TR from human and 150 other vertebrate species.

IMGT/LIGM-DB sequence data are identified by the EMBL/GenBank/DDBJ accession number. The unique source of data for IMGT/LIGM-DB is EMBL which shares data with the other two generalist databases GenBank and DDBJ. Once the sequences are allowed by the authors to be made public, LIGM automatically receives IG and TR sequences by e-mail from EMBL. After control by LIGM curators, data are scanned to store sequences, bibliographical references and taxonomic data, and standardized IMGT/LIGM-DB keywords are assigned to all entries. Based on expert analysis, specific detail annotations are added to IMGT flat files in a second step (Giudicelli et al., 1997). Since August 1996, the IMGT/LIGM-DB content has closely followed that of the EMBL for the IG and TR, with the following advantages: IMGT/LIGM-DB contains IG and TR entries which have disappeared from the generalist databases (as examples: the L36092 accession number which encompasses the complete human TRB locus is still present in IMGT/LIGM-DB, whereas it has been deleted from EMBL/GenBank/DDBJ due to its too large size (684,973 bp); in 1999, IMGT detected the disappearance of 20 IG and TR sequences which inadvertently had been lost by GenBank, and allowed the recuperation of these sequences in the generalist databases); conversely, IMGT/LIGM-DB does not contain sequences which have previously been wrongly assigned to IG and TR.

The IMGT/LIGM-DB data are provided with a user friendly interface. The Web interface allows searches according to immunogenetic specific criteria and is easy to use without any knowledge in a computing language. The interface allows the users to get easily connected from any type of platform (PC, Macintosh, workstation) using freeware such as Explorer, Netscape. All IMGT/LIGM-DB information is available through five modules of search:

catalogue, taxonomy and characteristics, keywords, annotation labels and references (Fig. 4). Selection is displayed at the top of the “results of your search” page, so the users can check their own queries (Lefranc et al., 1999). Users have the possibility to modify their request or consult the results. They can (i) add new conditions to increase or decrease the number of resulting sequences, (ii) view details: selecting this “View” option provides a list of resulting sequences; selection of one sequence in the list offers nine possibilities: annotations, IMGT flat file, coding regions with protein translation, catalogue and external references, sequence in dump format, sequence in FASTA format, sequence with three reading frames, EMBL flat file, IMGT/V-QUEST, or (iii) search for sequence fragments: selecting this “Subsequences” option allows to search for sequence fragments (subsequences) corresponding to a particular label for the resulting sequences (available for fully annotated sequences (Lefranc et al., 1999).

IMGT/LIGM-DB data are distributed by anonymous FTP servers at the Centre Informatique National de l’Enseignement Supérieur (CINES), Montpellier, France (<ftp://ftp.cines.fr/IMGT/>), at the European Bioinformatics Institute (EBI), Hinxton, UK (<ftp://ftp.ebi.ac.uk/pub/databases/imgt/>) and at the Institut de Génétique Humaine (IGH), Montpellier, France (<ftp://ftp.igh.cnrs.fr/pub/IMGT/>), and from many sequence retrieval system (SRS) sites. IMGT/LIGM-DB releases are produced weekly. Users can also compare their own sequences against IMGT/LIGM-DB data using BLAST or FASTA on different servers (EBI, IGH, INFOBIOGEN, Institut Pasteur Paris, etc.). Links are provided from the IMGT Home page, <http://imgt.cines.fr>, in “IMGT other accesses”).

IMGT/MHC-DB, hosted on the EBI server at Hinxton (UK), comprises a database of the human MHC allele sequences (IMGT/MHC-HLA, developed by Cancer Research, UK and Anthony Nolan Research Institute (ANRI), London, UK, on the Web since December 1998 (1738 entries in October 2003) (Robinson et al., 2003), databases of the MHC class II sequences from non-human primates (IMGT/MHC-NHP, curated by the Biomedical Primate Research Centre (BPRC), Rijswijk, The Netherlands) and from felines and canines (IMGT/MHC-FLA and IMGT/MHC-DLA, curated by the Centre for Integrated Genomic Medical Research, Manchester, UK), on the Web since April 2002 (Robinson et al., 2003).

IMGT/PRIMER-DB (Fig. 5) is the IMGT oligonucleotide (primer) database for the IG and TR, created by LIGM (Montpellier, France) in collaboration with EUROGEN-TEC S.A. (Seraing, Belgium), on the Web since February 2002. IMGT/PRIMER-DB provides information on IG and TR primers which are useful for the analysis of the IG and TR gene repertoire and expression, the detection of minimal residual diseases in B and T cell malignancies, the construction of antibody combinatorial libraries, scFv, phage display or microarray technologies. In October 2003, IMGT/PRIMER-DB contained 292 entries from

Help IMGT Home page IMGT Marie-Paule page New search

Created by Marie-Paule Lefranc (CNRS, Montpellier II University, France)
http://imgt.cines.fr

IMGT/LIGM-DB ON LINE, HERE YOU ARE !

Five types of search are available : select one by clicking on the button

Catalogue	accession number, mnemonic, definition, creation date, length, <u>annotation level</u>
Taxonomy, ...	<u>taxonomy</u> (species and classification level), nucleic acid type, " <u>Loci, genes or chains</u> ", <u>functionality</u> , <u>structure</u> , <u>specificity</u> , group and subgroup
Keywords	<u>standardized keywords</u> , selection of IMGT reference sequences (for human and mouse IG and TR) NEW!
Annotation labels	<u>standardized LIGM-DB labels</u>
References	authors, publication type, journal, year, title, Medline reference number

Short path selection

Enter enter an accession number and click on the "Enter" button.

Transferring data from imgt.cines.fr...

Fig. 4. IMGT/LIGM-DB search page (<http://imgt.cines.fr>). Five modules of search are available: Catalogue, Taxonomy and Characteristics, Keywords, Annotation labels, and References. These modules allow extensive and complex queries on IG and TR sequences from human and other vertebrates. In October 2003, IMGT/LIGM-DB contained 77,810 sequences of IG and TR from 150 species. A short path selection allows a direct query with an accession number or with a part of it. For example, "AJ012264" will retrieve that sequence, whereas "AJ012" will retrieve all sequences beginning with AJ012.

Homo sapiens and *Mus musculus*. IMGT/PRIMER-DB contains information on primers and combinations of primers described as "sets" (primers sharing identical properties (species, group and orientation)) and "couples" (sets of opposite orientation for which IMGT/LIGM-DB sequences are known (or expected)). Primers, Sets and Couples are described in IMGT Primer cards, IMGT Set cards and IMGT Couple cards, respectively. An IMGT Primer is an oligonucleotide described by comparison to an IMGT/LIGM-DB reference sequence, according to the standardized rules of the IMGT Scientific chart, based on IMGT-ONTOLOGY (Giudicelli and Lefranc, 1999). Taxonomy species and IMGT classification (group, subgroup, gene, allele) of a primer are those of the IMGT/LIGM-DB reference sequence, and not those of the PCR amplification products. This provides the following advantages for the data standardization: IMGT/PRIMER-DB primer definition, classification, and description are independent from the experimental conditions, from DNA sources and from the different combinations (sets and couples) in which the primer can be used. That means that (i) the specificity of a primer (subgroup, gene or allele specific) which is either described experimentally or deduced from sequence

comparison is not used for the primer classification, although these data are provided in the IMGT/PRIMER-DB Primer card in "Classification comments and specificity", (ii) the sequences resulting from the PCR amplifications are uniquely associated to the couples. The IMGT Primer cards are linked to IMGT/LIGM-DB flat files, IMGT Colliers de Perles and IMGT Repertoire > Alignments of alleles of the IMGT/LIGM-DB reference sequence used for the primer description.

IMGT/GENE-DB (Fig. 6) is the comprehensive IMGT genome database for IG and TR genes from human and mouse, and in development, from other vertebrates, created by LIGM, on the Web since January 2003. IMGT/GENE-DB annotated data are extracted from IMGT Repertoire, the global ImMunoGeneTics Web resource, and from the IMGT/LIGM-DB database. All the human IMGT gene names (Lefranc and Lefranc, 2001a,b) were approved by HGNC in 1999 (Wain et al., 2002a), and entered in GDB (Canada), LocusLink at NCBI (USA) and GeneCards. In October 2003, IMGT/GENE-DB contained 1375 genes and 2201 alleles (673 IG and TR genes and 1024 alleles from *H. sapiens*, and 702 IG and TR genes and 1177 alleles from *M. musculus*, *Mus cookii*, *Mus pahari*, *Mus*

WELCOME !
on **IMGT/PRIMER-DB**
THE
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Gene
Tics
Information
system®
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IMGT/PRIMER-DB Query page

IMGT/PRIMER-DB contains **292** entries:
(13/06/2003)

SEARCH BY :

Just enter the number

IMGT/PRIMER-DB Primer accession number (ex: IPP000201) : IPP

or IMGT/PRIMER-DB Set accession number (ex: IPS000018) : IPS

or IMGT/PRIMER-DB Couple accession number (ex: IPC000001) : IPC

OR SELECT IMGT/PRIMER-DB PRIMERS FOR A GROUP:

Species

IMGT group

Primer orientation :

or display, from the group list, only the primers which correspond to:

an IMGT subgroup ☐ select IMGT subgroup:

or an IMGT gene name ☐ select IMGT gene name:

or an IMGT allele name ☐ select IMGT allele name:

OR SELECT IMGT/PRIMER-DB SETS

Species

Fig. 5. IMGT/PRIMER-DB Query page (<http://imgt.cines.fr>). IMGT group, subgroup, gene and allele name used for the query and display are according to the IMGT gene nomenclature, based on the CLASSIFICATION concept of IMGT-ONTOLOGY.

spretus, *Mus saxicola*, *Mus minutoides*). IMGT/GENE-DB allows a search of IG and TR gene entries by locus, group, subgroup, based on the CLASSIFICATION concept of IMGT-ONTOLOGY (Giudicelli and Lefranc, 1999). A short cut allows to search genes by a selection on gene name (according to the IMGT nomenclature) or on clone name(s) (data from the “Reference sequences” and “Sequences from the literature” columns in IMGT Repertoire > Gene tables). The selection is displayed at the top of the resulting genes page. The users can select the genes to view their detailed entries. Each IMGT/GENE-DB entry corresponds to one gene and provides, for each gene, the chromosomal localization, the gene name and definition, the number of alleles, links to IMGT Repertoire and to external sequence databases (EMBL, GenBank, DDBJ), genome databases (LocusLink, OMIM), and gene nomenclature database (Genew (Wain et al., 2002b)). Reciprocally, LocusLink, GDB, GeneCards and HGNC Genew have direct links to the IMGT/GENE-DB entries. IMGT/GENE-DB provides for each allele, the functionality, the clone names, the IMGT/LIGM-DB reference sequence accession numbers

(with link to the flat files), and the “IMGT/GENE-DB reference sequences in FASTA format” (nucleotide and amino acid sequences of the coding regions extracted from the IMGT/LIGM-DB reference sequence), with gaps according to the IMGT unique numbering (Lefranc et al., 2003).

IMGT/3Dstructure-DB is the IMGT 3D structure database for IG, TR, MHC, and RPI of human and other vertebrate species, created by LIGM, on the Web since November 2001 (Kaas and Lefranc, 2002; Kaas et al., 2004). IMGT/3Dstructure-DB comprises IG, TR, MHC, and RPI with known 3D structures. In October 2003, IMGT/3Dstructure-DB managed 634 atomic coordinate files which correspond to 422 different proteins (260 IG, 18 TR and 144 MHC). Coordinate files are extracted from the Protein Data Bank (PDB) (Berman et al., 2000), and IMGT annotations are added according to the IMGT Scientific chart rules, based on the IMGT-ONTOLOGY concepts (Giudicelli and Lefranc, 1999). An IMGT/3Dstructure-DB card provides the IMGT gene and allele identification (based on the CLASSIFICATION concept), domain delimitations (based on the DESCRIPTION concept), amino

Fig. 6. IMGT/GENE-DB Query page (<http://imgt.cines.fr>). IMGT locus, group, subgroup are according to the IMGT nomenclature, based on the CLASSIFICATION concept of IMGT-ONTOLOGY. In October 2003, IMGT/GENE-DB contained 1375 IG and TR genes and 2201 alleles, from human and mouse.

acid positions according to the IMGT unique numbering (Lefranc et al., 2003) (based on the NUMEROTATION concept). Domains that are analyzed in IMGT/3Dstructure-DB include V-DOMAIN (variable) and C-DOMAIN (constant) found in IG and TR, V-LIKE- and C-LIKE-DOMAINS found in proteins other than IG and TR, G-DOMAIN (groove) found in MHC, and G-LIKE-DOMAINS found in proteins other than MHC (Duprat and Lefranc, 2003). Moreover, IMGT/3Dstructure-DB provides renumbered coordinate flat files, Colliers de Perles (Fig. 3) (standard and two-layer 2D graphical representations) (Ruiz and Lefranc, 2002; Kaas and Lefranc, 2002; Kaas et al., 2004), and results of contact analysis. The IMGT unique numbering and gene standardization will provide a great help in large scale sequence–structure studies and more generally in protein engineering.

4. IMGT interactive tools

IMGT interactive immunoinformatics tools rely on the DESCRIPTION and NUMEROTATION concepts of IMGT-ONTOLOGY, and are available for sequence anal-

ysis: IMGT/V-QUEST, IMGT/JunctionAnalysis, IMGT/PhyloGene and IMGT/Allele-Align, for genome analysis: IMGT/GeneSearch, IMGT GeneView, IMGT LocusView, and for 3D structural analysis: IMGT/StructuralQuery. IMGT/V-QUEST (V-QUery and standardization) is an integrated software for IG and TR (Lefranc, 2001). This tool, easy to use, analyses an input IG or TR germline or rearranged variable nucleotide sequence. IMGT/V-QUEST results (Fig. 7) comprise the identification of the V, D, and J genes and alleles and the nucleotide alignment by comparison with sequences from the IMGT reference directory, the delimitations of the FR-IMGT and CDR-IMGT based on the IMGT unique numbering, the protein translation of the input sequence, the identification of the JUNCTION and the 2D Collier de Perles representation of the V-REGION. IMGT/V-QUEST is particularly useful for the analysis of the rearranged variable genes: it allows to identify the V-GENE and J-GENE and alleles involved in the IG and TR rearrangements, and to delimit the JUNCTION. Searches can be done related to IG and TR of human and mouse, and of other species (non-human primates, sheep, teleostei, and chondrichthyes). IMGT/V-QUEST can also be used for the analysis of functional or ORF

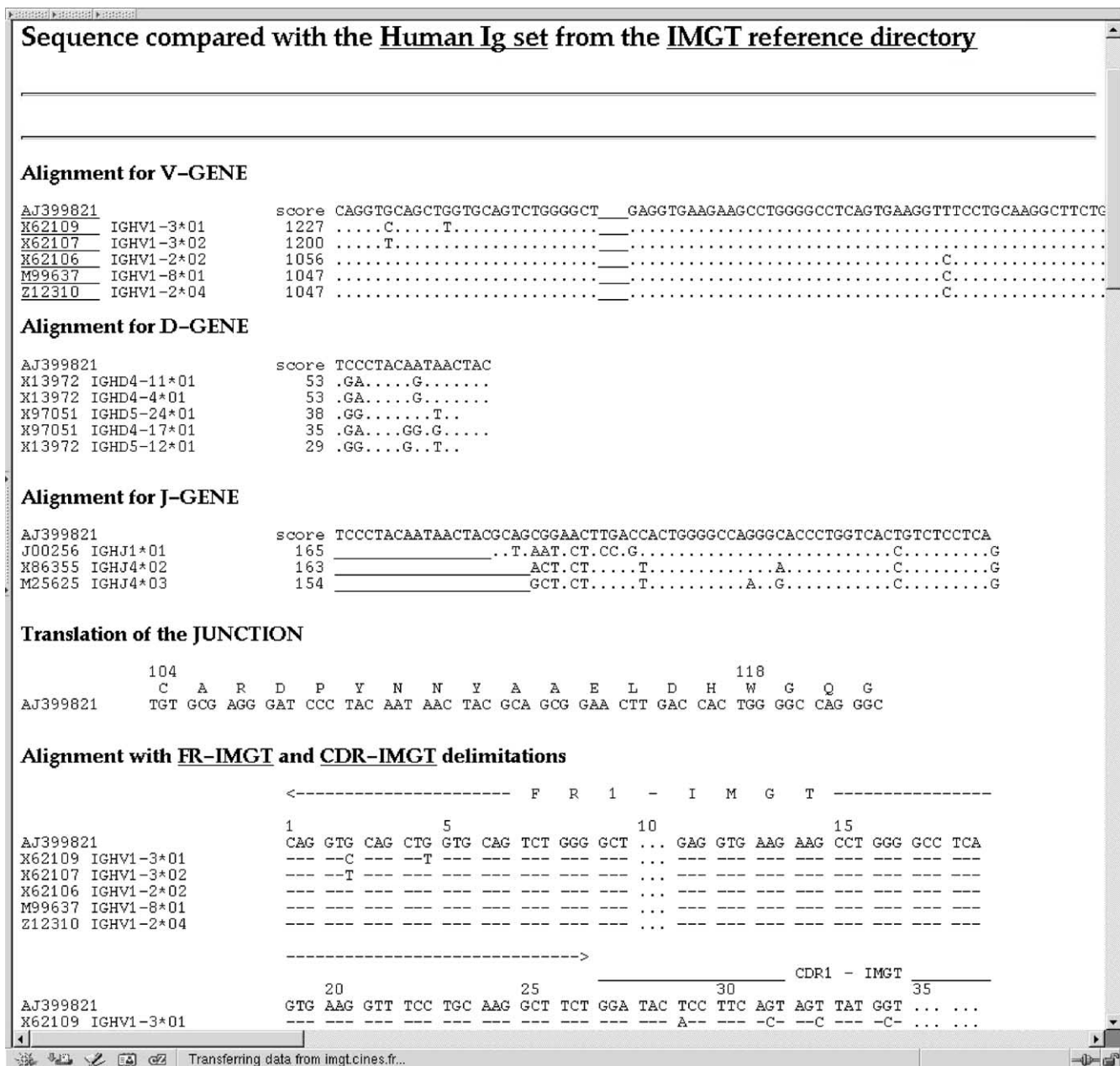


Fig. 7. IMGT/V-QUEST (<http://imgt.cines.fr>) results on gene and allele identification. IMGT/V-QUEST compares the input germline or rearranged IG or TR variable sequences with the IMGT/V-QUEST reference directory sets. For example, the highest scores for the input AJ399821 rearranged sequence allow to identify IGHV1-3*01, IGHD4-4*01 or IGHD4-11*01, IGHJ1*01 as being the genes and alleles most probably involved in the V-D-J rearrangement. The IMGT/V-QUEST results comprise the translation of the JUNCTION for rearranged sequences, the delimitations of the FR-IMGT and CDR-IMGT, and also, not shown in the figure, the protein translation and the two-dimensional representation or Collier de Perles of the V-REGION. Information provided by IMGT/V-QUEST (V and J gene and allele names, sequence of the JUNCTION (from 2nd-CYS 104 to J-PHE or J-TRP 118)) can then be used in IMGT/JunctionAnalysis for a confirmation of the D gene and allele identification and a more accurate analysis of the junction (see Fig. 8).

germline variable genes from other species, to delimit the FR-IMGT and CDR-IMGT, provided that the similarity with sequences of the IMGT/V-QUEST reference directory sets is sufficiently high. The set of sequences from the IMGT reference directory, used for IMGT/V-QUEST, can be downloaded in FASTA format from the IMGT site. IMGT/JunctionAnalysis is a tool, complementary to IMGT/V-QUEST, which provides a thorough analysis of

the V-J and V-D-J junction of IG and TR rearranged genes (Fig. 8). IMGT/JunctionAnalysis identifies the D-GENES and alleles involved in the IGH, TRB and TRD V-D-J rearrangements by comparison with the IMGT reference directory, and delimits precisely the P, N and D regions (Fig. 8). Results from IMGT/JunctionAnalysis are more accurate than those given by IMGT/V-QUEST regarding the D-GENE identification. Indeed, IMGT/JunctionAnalysis

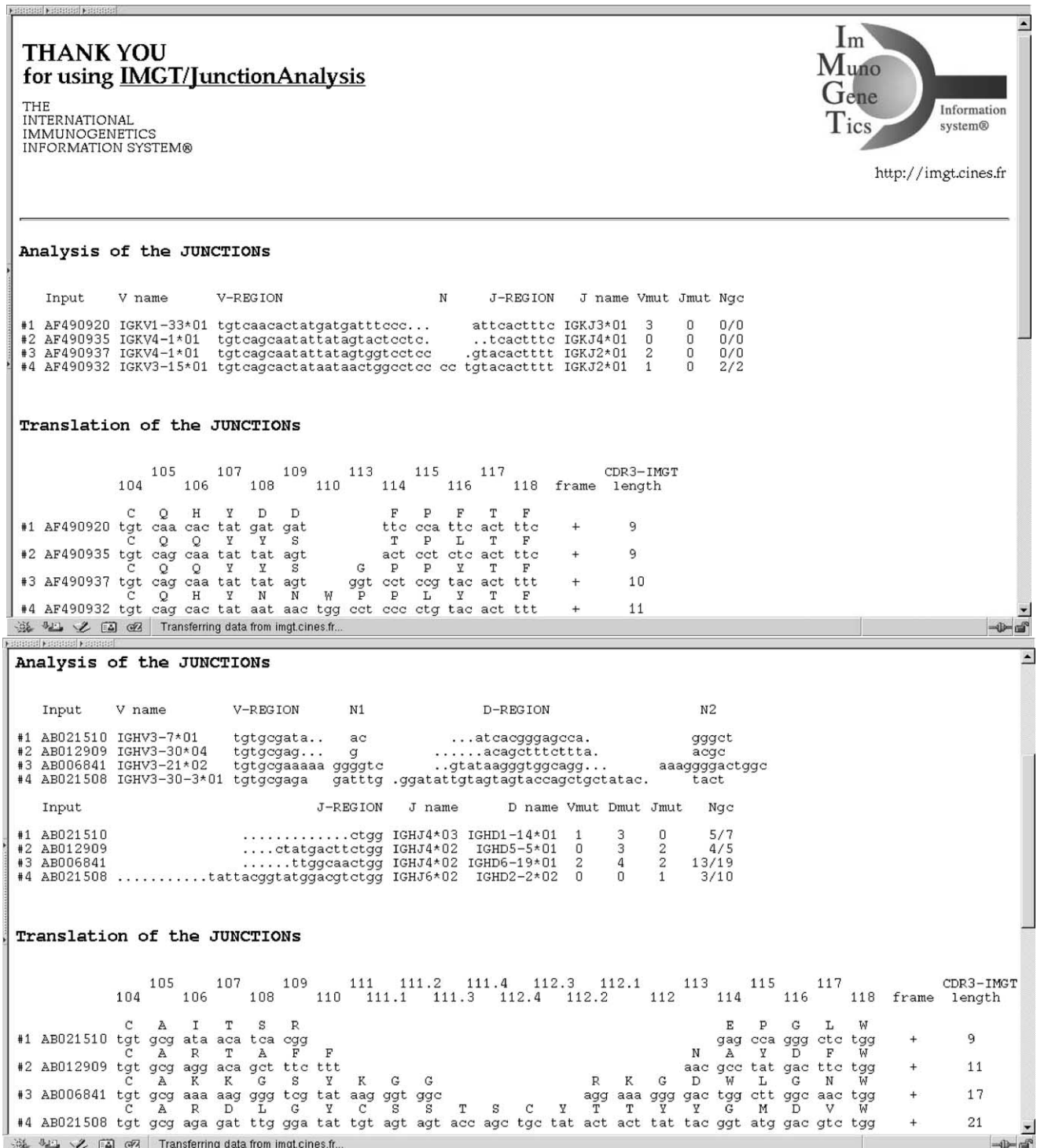


Fig. 8. IMGT/Junction-Analysis (<http://imgt.cines.fr>) results. (A) Examples of IGH V-D-J junctions. (B) Examples of IGH V-D-J junctions. The IMGT/JunctionAnalysis results comprise the junction translation, the identification of the D-GENE and allele (for each V-D-J junction), the identification of the P and N regions (N1, N2, etc.) and their precise delimitations, the identification of the deleted nucleotides by comparison to the germline allele sequences. The CDR3-IMGT numbering is according to the IMGT unique numbering for V-DOMAIN. Vmut, Dmut and Jmut correspond to the number of mutations in the input junction sequence by comparison to the germline allele sequences. Ngc is the ratio of the number of g + c nucleotides to the total number of nucleotides in the N regions. IMGT/JunctionAnalysis analyses, in a single search, an unlimited number of junctions provided that the V-GENE and J-GENE allele IMGT names are identified.

works on shorter sequences (JUNCTION), and with a higher constraint since the identification of the V-GENE and J-GENE and alleles is a prerequisite to perform the analysis. Several hundreds of junction sequences can be analyzed simultaneously.

IMGT/PhyloGene is an easy to use tool for phylogenetic analysis of variable region (V-REGION) and constant domain (C-DOMAIN) sequences (Elemento and Lefranc, 2003). This tool is particularly useful in developmental and comparative immunology. The users can analyze their own sequences by comparison with the IMGT standardized reference sequences for human and mouse IG and TR. IMGT/Allele-Align allows the comparison of two alleles highlighting the nucleotide and amino acid differences. IMGT/GeneSearch, IMGT GeneView and IMGT LocusView are tools which provide the display of physical maps for the human IG, TR, and MHC loci. The mouse TRA/TRD locus is also available. IMGT/StructuralQuery is an interactive tool which allows to retrieve the IMGT/3Dstructure-DB entries based on specific structural characteristics: phi (ϕ) and psi (ψ) angles, accessible surface area ASA, amino acid type, distance in angstrom between amino acids, CDR-IMGT lengths (Kaas and Lefranc, 2003; Kaas et al., 2004). IMGT/StructuralQuery is currently available for the V-DOMAINS.

5. IMGT Web resources

The IMGT Web resources (“IMGT Marie-Paule page”) (Lefranc, 2003a) consist of 8000 HTML pages and comprise the following sections: “IMGT Scientific chart”, “IMGT Repertoire”, “IMGT Bloc-notes”, “IMGT Education” and “IMGT Index”. IMGT Scientific chart provides the controlled vocabulary and the annotation rules and concepts defined by IMGT-ONTOLOGY (Giudicelli and Lefranc, 1999) for the identification, the description, the classification and the numerotation of the IG, TR, MHC, and RPI data of human and other vertebrates. The IMGT Scientific chart rules are described in the corresponding sections: sequence and 3D structure identification and description, nomenclature, numbering (<http://imgt.cines.fr/textes/IMGTScientificChart>).

IMGT Repertoire is the global Web resource in ImMunoGeneTics for the IG, TR, MHC, and RPI of human and other vertebrates, based on the “IMGT Scientific chart”. IMGT Repertoire provides an easy-to-use interface to carefully and expertly annotated data on the genome, proteome, polymorphism, and structural data, organized in three major sections: IMGT Repertoire (IG and TR), IMGT Repertoire (MHC), and IMGT Repertoire (RPI) (Lefranc, 2001a). Only titles of this large section are quoted here. Genome data (“Locus and genes”) include chromosomal localizations, locus representations, locus description, gene tables, potential germline repertoires, lists of IG and TR genes and links between IMGT, HUGO, GDB, LocusLink and

OMIM, correspondence between nomenclatures (Lefranc and Lefranc, 2001a,b), reference sequences (Barbié and Lefranc, 1998; Pallarès et al., 1998, 1999; Ruiz et al., 1999; Folch and Lefranc, 2000a,b; Scaviner and Lefranc, 2000a,b; Martinez-Jean et al., 2001; Bosc and Lefranc, 2003). Proteome and polymorphism data (“Proteins and alleles”) are represented by protein displays which show translated sequences of the allele *01 of each functional or ORF gene (Lefranc and Lefranc, 2001a,b; Scaviner et al., 1999; Folch et al., 2000), alignments of alleles, tables of alleles, allotypes, particularities in protein designations, IMGT reference directory in FASTA format, correspondence between IG and TR chain and receptor IMGT designations (Lefranc and Lefranc, 2001a,b). Structural data (“2D and 3D structures”) comprise 2D graphical representations designated as Colliers de Perles (Lefranc et al., 1998, 1999), FR-IMGT and CDR-IMGT lengths, and 3D representations of IG and TR variable domains (Ruiz and Lefranc, 2002; Lefranc et al., 2003). This visualization permits rapid correlation between protein sequences and 3D data retrieved from the PDB. Other data comprise: “Probes and RFLP” with phages, probes used for the analysis of IG and TR gene rearrangements and expression, and restriction fragment length polymorphism (RFLP) studies, “Taxonomy” of vertebrate species present in IMGT/LIGM-DB, “Gene regulation and expression” with data on promoters, primers, cDNAs, reagent monoclonal antibodies, and “Genes and clinical entities”: translocations and inversions, humanized antibodies, monoclonal antibodies with clinical indications.

IMGT Bloc-notes is a selection of useful links for immunoinformatics, immunogenetics, immunology, genetics, molecular biology, and bioinformatics. IMGT Bloc-notes is organized in several sections. The IMGT immunoinformatics page comprises links to databases, tools, and resources on IG, TR, MHC, and RPI. Interesting links provide numerous hyperlinks towards the Web servers specializing in immunology, genetics, molecular biology, and bioinformatics (associations, biopharmaceuticals, collections, companies, databases, immunology themes, journals, molecular biology servers, resources, societies, tools, etc.) (Lefranc, 2000e). Other sections are meeting announcements, postdoctoral positions, etc.

IMGT Education is a section which provides useful biological resources for students. It includes IMGT Aide-mémoire which provides an easy access to information such as genetic code, splicing sites, amino acid structures, restriction enzyme sites, etc., questions and answers, tutorials (in English and/or in French) on 3D structure, immunoglobulins and B cells, T cell receptors and T cells, NK receptors, pathologies of the immune system, cancer, AIDS, etc. IMGT Index is a referential index which provides a fast way to access data when information has to be retrieved from different parts of the IMGT site. For example, “allele” provides links to the IMGT Scientific chart rules for the allele description, and to the IMGT Repertoire Alignments of alleles and Tables of alleles.

6. Conclusion

Since July 1995, IMGT has been available on the Web at <http://imgt.cines.fr>. IMGT provides the biologists with an easy to use and friendly interface. Since January 2000, the IMGT www server at Montpellier has been accessed by more than 250,000 sites. IMGT has an exceptional response with more than 140,000 requests a month. Two-thirds of the visitors are equally distributed between the European Union and the United States. IMGT distributes high quality data with an important incremental value added by the IMGT expert annotations, according to the rules described in the IMGT Scientific chart. Control of coherence in IMGT combines data integrity control and biological data evaluation (Giudicelli et al., 1998a,b). The information provided by IMGT is of much value to clinicians and biological scientists in general (Lefranc, 2002, 2003b; Chardès et al., 2002). Further developments include the implementation of the IMGT/GeneInfo (Baum et al., 2004) and IMGT/GeneFrequency tools, and that of IMGT/PROTEIN-DB, a protein database for IG and TR, which will contain translations of potentially functional and ORF sequences from IMGT/LIGM-DB, and protein data from Kabat et al. (1991) and PDB. To facilitate the integration of IMGT data into applications developed by other laboratories, we have built an application programming interface (API) to access the database (Giudicelli et al., 1998a). This API includes: a set of URL links to access biological knowledge data (keywords, labels, functionalities, list of gene names, etc.), a set of URL links to access all data related to one given sequence. To increase interoperability with other ontologies and information systems, IMGT-ONTOLOGY is currently being written using extensible markup language (XML) approach, in IMGT-ML (Chaume et al., 2001, 2003). IMGT-ML will also be used for implementing Web services and developing IMGT-Chorégraphie, a long-term project aimed at creating dynamic interactions between the IMGT databases, tools, and Web resources. IMGT is designed to allow a common access to all immunogenetics data, and particular attention is given to the establishment of cross-referencing links to other databases pertinent to the users of IMGT.

7. Citing IMGT

If you use IMGT databases, tools and/or Web resources, please cite Lefranc (2003) and this paper as references, and quote the IMGT Home page URL address, <http://imgt.cines.fr>.

Acknowledgements

I am deeply grateful to the IMGT team for its hard work, expertise, constant motivation and enthusiasm. “Mille

mercis” to Nathalie Bosc, Denys Chaume, Kora Combres, Elodie Duprat, Géraldine Folch, Chantal Ginestoux, David Girod, Véronique Giudicelli, Delphine Guiraudou, Joumana Jabado-Michaloud, Stéphanie Jeanjean, Quentin Kaas, Gérard Lefranc, Séverine Magris, Christelle Pommié, Céline Protat, Dominique Scaviner, Valérie Thouvenin, and Mehdi Yousfi Monod. I thank Marc Lemaitre, Marie-Claire Beckers, Géraldine Folch, and Delphine Valette from EUROGENTEC S.A., Belgium, for their scientific contribution to IMGT/PRIMER-DB, Julien Bertrand, Olivier Elemento, Christèle Jean-Martinez, and Manuel Ruiz for their previous work at LIGM, Peter Stoehr (EBI), Denis Pugnère (IGH), Nicolas Joly (Institut Pasteur Paris), and Jean-Marc Plaza (INFOBIOGEN) for the IMGT/LIGM-DB flat file distribution. IMGT is funded by the European Union's fifth PCRDT programme (QLG2-2000-01287), the Centre National de la Recherche Scientifique (CNRS), the Ministère de l'Éducation Nationale, and the Ministère de la Recherche. Subventions have been received from Association pour la Recherche sur le Cancer (ARC) and the Région Languedoc-Roussillon.

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