Deep Insight Section

General resources in Genetics and/or Oncology

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Abstract

This “Deep Insight” is a detailed subchapter of a general review article and summary on Internet databases for cytogeneticists: Internet databases and resources for cytogenetics and cytogenomics.

I- Bibliography

PubMed is a widely used and free search engine and database of biomedical citations and abstracts, based essentially on the MEDLINE database of references and abstracts on life sciences and biomedical topics. The database is maintained by the National Center for Biotechnology Information (NCBI), at the U.S. National Library of Medicine (NLM), located at the National Institutes of Health (NIH), as part of the Entrez system of information retrieval. From 1971 to 1997, the online version of MEDLINE through computerized database MEDLARS was mainly accessed through institutions, such as university libraries. In 1996, PubMed was launched but only as late as 1997 gave free access of MEDLINE to private home and office computers. PubMed Advanced Search Builder http://www.ncbi.nlm.nih.gov/pubmed/advanced uses keywords such as: Affiliation, All Fields, Author, Author First, Author Last, Journal, MeSH Major Topic, Title, Title/Abstract (Figure 1). It uses Booleans (AND, OR, NOT). You can query "(KMT2A[Title]) AND ((Acute myeloid leukemia) OR (Acute lymphoid leukemia))", you will get: Search results : Items 5. This only shows that the official name KMT2A remains totally ignored by scientists. If you replace KMT2A with MLL: "(MLL[Title]) AND ((Acute Myeloid Leukemia) OR (Acute lymphoid leukemia))", you get: Search results : Items 885. Which is what you were looking for. On the other hand, if you misuse the brackets in your query (e.g. "((KMT2A[Title]) AND Acute myeloid leukemia) OR Acute lymphoid leukemia"), you will have a huge amount of background noise! Search results: Items 36,244. PubMed comprises of more than 25 million citations of biomedical literature from MEDLINE, life science journals, and online books. Citations may include links to full-text content from PubMed Central and publisher web sites. As a broader research engine, PubMed also runs in several other databases like MEDLINE and Index Medicus, providing older references of the print versions as well as some journals not yet cited, like Science. The research engine also accesses entries for an article before it gets indexed by the Medical Subject Headings (MeSH) and added to MEDLINE.
Collections of full-text available books and other subsets of NLM records are available (https://www.nlm.nih.gov/pubs/factsheets/pubmed.html). The references catalogued in PubMed often contain links to the full text articles, some of them are free of access and more often in PubMed Central (http://www.ncbi.nlm.nih.gov/pmc/) and local mirrors like UK PubMed Central (http://www.jisc-content.ac.uk/node/52) or Europe PMC (https://europepmc.org/). NLM catalogue contains all the necessary information about the journals that are indexed in PubMed (http://www.ncbi.nlm.nih.gov/nlmcatalog). PubMed records back to 1966, selectively to the year 1865, and very selectively to 1809; about 500,000 new records are added each year. As of this date, 14,026,022 records are listed with their abstracts. Only journals achieving PubMed's scientific standards are indexed which, on the one hand, provides a way to control the quality of scientific publishing. PubMed, free of use, is an immense gift to the medical and scientific community. However, from the scientific editor's viewpoint, this quasi-monopoly position has an adverse aspect: to be referenced by PubMed is a terrifying verdict, in terms of recognition. This is all the more concerning, as the Literature Selection Technical Review Committee's decisions have been known to create controversy among scientific editor's and publisher's communities.

PubMed Central (http://www.ncbi.nlm.nih.gov/pmc/)
PubMed Central (PMC) is a free archive of biomedical and life sciences journal literature at the U.S. National Institutes of Health's National Library of Medicine (NIH/NLM) developed by the National Center for Biotechnology Information (NCBI). PubMed Central should not be confused with PubMed. As an archive, PMC is designed to provide permanent access to all of its content. Articles are deposited by participating journals, as well as for author manuscripts that have been submitted in compliance with the public access policies of participating research funding agencies.

Medline
Medline is the U.S. National Library of Medicine (NLM) premier bibliographic database that contains more than 22 million references to journal articles in life sciences with emphasis on biomedicine. A distinctive feature of MEDLINE is that the records
are indexed with NLM Medical Subject Headings (MeSH). MEDLINE is the primary component of PubMed.

Scopus (http://www.scopus.com/)
Scopus is a large abstract and citation database of peer-reviewed literature: scientific journals (more than 60 million records in Scopus, which includes over 21,500 peer-reviewed journals), books (more than 113,000 books) and conference proceedings. It is owned by Elsevier and it is available online by subscription.

II- Nomenclatures
Gene Nomenclature: HGNC (http://www.genenames.org/)
The HUGO Gene Nomenclature Committee (HGNC) is the worldwide authority assigning standardised nomenclature to human genes. HGNC approves unique names and symbols for human loci, including protein coding genes, ncRNA genes and pseudogenes, gene families and associated resources including links to genomic, proteomic and phenotypic information, to allow more unambiguously to scientific communication. This database contains 39,000 approved symbols (Gray KA et al., 2015).

Nomenclature for the description of sequence variations (http://www.hgvs.org/mutnomen/)
The nomenclature for the description of sequence variations is maintained by the Human Genome Variation Society (HGVS). When describing a variation, first, i) Indicates the reference sequence (e.g. coding DNA: "c."); RNA: "r."); Protein: "p.").

For reasons of interoperability between different databases it is essential that a common language is found.

The WHO/OMS has established the ICD-O code for International Classification of Diseases - Oncology, first published in 1976. The third edition of ICD-O (ICD-O-3) contains an ICD-O-3-TOPO, which provides a topographical identifier for different organs (e.g. C220: Liver; C339: Trachea), and an ICD-O-3-MORPH, which provides basic and detailed description of pathology (e.g. respectively: 801: Carcinoma, NOS (not otherwise specified); 8013/3: Large cell neuroendocrine carcinoma; 922: Chondrosarcoma, NOS; 9221/3: Juxtacortical chondrosarcoma). A "0" means: benign tumor (e.g.: 9220/0: Chondroma); "1" means: borderline malignancy (e.g. 9751/1: Langerhans cell histiocytosis); "2" means: malignant tumor in situ (e.g. 8500/2: Intraductal carcinoma, noninfiltrating, NOS); and "3" means full malignancy.

III- Nucleic acid, genes and protein databases
III-1 Nucleic acid databases
The first database for DNA sequencing was The Los Alamos Sequence Database in 1979, which was consequently replaced by public GenBank (http://www.ncbi.nlm.nih.gov/genbank/) (Burks C et al., 1985) in 1982. The database was funded by the National Institutes of Health, the National Science Foundation, the Department of Energy, and the Department of Defense. Los Alamos National Laboratories (LANL) collaborated with several firms like Bolt, Beranek, and Newman to increase the size of the database. By the end of 1983 more than 2,000 sequences were stored in it.
Mid 1980s, the Intelligenetics bioinformatics company from Stanford University collaborated with LANL to manage the GenBank project (Burks et al., 1991). Since it was one of the earliest bioinformatics community projects on the Internet, BIOSCI/Bionet news groups was created to promote open access communications among bioscientists. From 1989 to 1992, the GenBank project transitioned to the newly created National Center for Biotechnology Information (Benton D, 1990).

From 1982 to present day, the number of bases in GenBank has doubled roughly every 1.5 years (Benson DA et al., 2015). As of February 2016, GenBank version 212.0 contains 190,250,235 loci, 207,018,196,067 bases, from 190,250,235 reported sequences (http://www.ncbi.nlm.nih.gov/genbank/statistics/).

The GenBank database includes additional data sets that are constructed mechanically from the main sequence data collection, and therefore are excluded from this count. In parallel, the EMBL database was created in 1981 and since this date there is an International Nucleotide Sequence Database Collaboration (INSDC) which is a long-standing foundational initiative that operates between DDBJ, EMBL-EBI and NCBI. INSDC covers the spectrum of data raw reads, through alignments and assemblies to functional annotation, enriched with contextual information relating to samples and experimental configurations. In particular there are numerous evolutions with the development of massive sequencing with creation of more integrated structures as ENA (European Nucleotide Archive at EBI, http://www.ebi.ac.uk/services/dna-rna) or SRA (Sequence read archive at NCBI, http://www.ncbi.nlm.nih.gov/sra/) (Cook CE et al., 2016).

In parallel with the genome projects, the need for the best representation of genomic and transcript sequences for diverse species has been the driver for creating consensus databases (as RefSeq, UCSC, Ensembl) with several methods of optimisation.

### III-2 Genes and Functions

#### Genomic sequences and transcripts

As mentioned in the general resources, several consensus nucleic sequence databases provide detailed structures of genes and isoforms. All the information can easily be visualized using different browsers (UCSC, Ensembl) or described in detail on the Entrez Gene (see above) page at NCBI. RefSeq (http://www.ncbi.nlm.nih.gov/refseq/) maintains and curates a database recording annotated genomic, transcript, and protein sequences. RefSeq release 71 provides sequences from over 55,000 organisms (more than 4,800 viruses, 40,000 prokaryotes and 10,000 eukaryotes) (OLeary NA et al., 2016). Ensembl (http://www.ensembl.org/) is a joint project between EMBL-EBI and the Wellcome Trust Sanger Institute to develop a software which develops and maintains automatic annotation of selected eukaryotic genomes (Gray KA et al., 2015).

The UCSC Genome Browser database is a large collection of 160 genome assemblies representing 91 species (Rosenbloom KR et al., 2015) (Figures 2 and 3: PAX5 at UCSC and at the Atlas site respectively).

Some standardisation within CCDS and GenCode http://www.gencodegenes.org/ gives an up-to-date information on them.

The nature of isoforms, expressed differently in normal tissues and in tumors, due to splicing variety, leads to protein product with different amino acid sequences.

This reflects the variations in the structure in domains and in the 3D structure are the basis of the activity. On the other hand, the level of expression of transcript in different tissues can be obtained from SOURCE, GEO (Clough and Barett, 2016 ), Expression Atlas (Petryszak et al., 2016), Gene expression viewer (Firebrowse), BioGPS (http://biogps.org/#goto=welcome) (Wu C et al., 2016) (Figure 4).

#### III-3 Protein sequence databases

In parallel with the nucleic databases, the first protein database was established by M. Dayhoff as NBRF protein database in 1983, in continuity of the first comprehensive collection of macromolecular sequences in the Atlas of Protein Sequence and Structure, published from 1965-1978.

This was followed by the development of SwissProt, a curated dataset, by Amos Bairoch in 1986 (http://www.isb-sib.ch/sp30/the-history-of-swiss-prot).

With collaboration between the Swiss Institute of Bioinformatics and the EBI to lead in 2002 (in association with the PIR database) the SwissProt was extended to UniProt Knowledgebase (UniProtKB) in 1998, consisting in the curated UniProtKB/Swiss-Prot databank, its automatically annotated supplement TrEMBL, and the PIR protein database.

Today, UniProtKB represents the world’s most comprehensive catalogue of information on proteins. In the space of 30 years, the number of proteins entered in UniProtKB/Swiss-Prot has increased from 4,000 to 550,000 : 550,960 entries for the SwissProt part and 63,686,057 entries for the non-reviewed part for TrEMBL (Pundir S et al., 2015).
**Figure 2:** PAX5 gene with isoforms at UCSC (http://genome-euro.ucsc.edu/cgi-bin/hgGateway), Select Species: "Human"; Human Assembly: "Dec. 2013 (GRCh38/hg38)"; Position/Search Term: write "PAX5"; go!

**Figure 3:** PAX5 gene and protein in the Atlas (http://atlasgeneticsoncology.org//Genes/PAX5ID62.html)
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Figure 4: Expression of PAX3 in various tissues at BioGPS (http://biogps.org/#goto=genereport&id=5077)

UniProt (http://www.uniprot.org/)
The UniProt Knowledgebase (UniProt Consortium, 2015) (UniProtKB, http://www.uniprot.org/uniprot/) is a hub for the collection of information on proteins with annotation. In addition to amino acid sequences, protein names and domain descriptions, taxonomic data and citation information, it also provides brief annotation information (Figure 5). UniProtKB consists of two sections: computationally analyzed "TrEMBL" and manually annotated "Swiss-Prot", with information extracted from curator-evaluated computational analysis and literature. UniProt is a collaboration between the European Bioinformatics Institute (EMBL-EBI, http://www.ebi.ac.uk/), the SIB Swiss Institute of Bioinformatics (http://www.isb-sib.ch/) and the Protein Information Resource (PIR, http://pir.georgetown.edu/).

It is comprised of two separate tools: the Basic Local Alignment Search Tool (BLAST, http://www.uniprot.org/blast/), to find a region of local similarity between amino acid sequences used in identifying members of a gene family, and Align (http://www.uniprot.org/align/) to align two or more protein sequences.

neXtProt (http://www.nextprot.org/db/neXtProt (Gaudet P et al., 2015) is a resource for human proteins, including information on the exons, proteins sequences, function, subcellular localisation, expression, interactions and role in diseases (Figure 6). The major part of the information in neXtProt is obtained from the UniProt Swiss-Prot database but is gradually being complemented by original data. neXtProt contains 20,055 protein entries, and is maintained by Amos Bairoch at the Swiss Institute of Bioinformatics and GeneBio.
Figure 5: PAX5 at UniProtKB (http://www.uniprot.org/uniprot/Q02548)

Figure 6: PAX5 at neXtProt, tab "Function" (see on the left) (http://www.nextprot.org/db/entry/NX_Q02548)
Figure 7: PAX5 at PhosphoSitePlus (http://www.phosphosite.org/proteinAction.action?id=19058&showAllSites=true)

PhosphoSitePlus (http://www.phosphosite.org/homeAction.action)
PhosphoSitePlus (Hornbeck PV et al., 2015) is an excellent resource providing comprehensive information and tools for the study of protein post-translational modifications (PTMs) including phosphorylation, ubiquitination, acetylation and methylation (Figure 7). PhosphoSitePlus contains curated data on 53,219 human, mouse and to a lesser extent rat proteins, with protein name, protein type, domain, cellular component, and molecular weight. It is an excellent website. PhosphoSitePlus is based at Cell Signaling Technology, Danvers, Massachusetts.

PROSITE (http://prosite.expasy.org/)
PROSITE (Sigrist CJ et al., 2013) is one of the oldest catalogs of protein signatures, consisting of documentation entries describing protein domains, families and functional sites, via a specific pattern of conserved residues (manually defined). PROSITE contains 1756 documentation entries.

Pfam (http://pfam.xfam.org/)
Pfam (Finn RD et al., 2016) is a collection of multiple sequence alignments and hidden Markov models covering many common protein domains. The identification of domains that occur within proteins can provide insights into their function. Pfam contains 16295 entries. InterPro and Pfam are based at EMBL-EBI.

InterPro (http://www.ebi.ac.uk/interpro/)
InterPro integrates PROSITE, Pfam and certain other resources in order to provide functional analysis of proteins by classifying them into families and predicting domains (with signatures) and important sites; InterProScan is the software package that allows sequences to be scanned against InterPro's signatures (Mitchell A et al., 2015).
Figure 8: and 9: JAK2 and SQSTM1 at Atlas: protein domains (http://atlasgeneticsoncology.org//Genes/JAKID98.html and http://atlasgeneticsoncology.org//Genes/GC_SQSTM1.html) There are also data and iconography on pathways (Figure 10).
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The Atlas presents highly curated paragraphs with the description of the protein listing domains and iconography, expression and localisation, function, homologs, and uniquely, a wide angle on cancers and other medical conditions where a gene or a protein is implicated (Figure 8 and 9).

IV- Cards


Entrez is NCBI’s primary text search and retrieval system integrating the PubMed database of biomedical literature with 39 other literature and molecular databases including DNA and protein sequences, structures, genes, genomes, genetic variation and gene expression. Entrez Gene, dedicated to gene information, integrates data from a wide range of species. A record can include nomenclature, Reference Sequences (RefSeqs), maps, pathways, variations, phenotypes, and links to genome-, phenotype- and locus-specific resources worldwide. Entrez Gene catalogs 59,941 human genes. Entrez Gene can be queried as a free text but also via a syntax with specific fields or filters (e.g. BRCA1[sym] ; 2[chr] AND adh*[sym]

..) with output in different formats. Once a result is obtained as a list of gene symbols, it is possible to link it to related data in another part of the Entrez database (e.g. list of publication in PubMed from a selected list of genes symbols) (NCBI Resource Coordinators, 2016).

IV-2 Genecards (http://www.genecards.org/)

Genecards is an integrative database that provides comprehensive, user-friendly information on all annotated and predicted human genes. It automatically integrates data from roughly 125 web sources and includes genomic, transcriptomic, proteomic, genetic, clinical and functional information.

There are some affiliated databases as MalaCards “The human disease database” (http://www.malacards.org/) which is an integrated database of human diseases and their annotations, modeled on the architecture and richness of the GeneCards database of human genes (Fishilevich S et al., 2016).

V- Genome cartography

The cartography of genes on a genome has been the favoured mean to represent genomic information. With the human Genome Project, several types of viewers have been developed. To date, two sites are of first interest for human genetics:
V-1 UCSC (http://genome.ucsc.edu/) and UCSC-Cancer (https://genome-cancer.ucsc.edu/)
The UCSC Genome Browser contains a reference sequence and working draft assemblies for a large collection of genomes. It also provides portals to ENCODE data at UCSC (2003 to 2012).
The Genome Browser zooms and scrolls over chromosomes, presenting the work of annotators worldwide. The "Gene Sorter" shows expression, homology and other information on groups of genes that can be related in many ways (with a chosen set of tracks). "Blat" maps sequences to the genome quickly. The Table Browser provides convenient access to the underlying database. "VisiGene" lets you browse through a large collection of in situ mouse and frog images to examine gene expression patterns. "Genome Graphs" allows you to upload and display genome-wide data sets. The UCSC Genome Browser is developed and maintained by the Genome Bioinformatics Group, a cross-departmental team within the UC Santa Cruz Genomics Institute at the University of California Santa Cruz (UCSC).
A parallel browser has been developed for visualizing and analysing cancer data. The UCSC Cancer Browser (https://genome-cancer.ucsc.edu/proj/site/help/) allows researchers to explore cancer genomics data and its associated clinical information in an interactive manner. Data can be viewed in several different ways, including by value, chromosome location, clinical features, biological pathways or genes of interest. It is also possible to quickly perform and easily view statistical analysis on subsets of the data. The data heatmap displays genome-wide data from copy number, transcriptome, protein, epigenetic, mutation, sh/siRNA, and PARADIGM pathway analysis studies as well as associated clinical information. The left column shows datasets that are currently in view along with a button to add more. Today the system has 720 datasets for an exploration (Goldman M et al., 2015).

V-2 Ensembl (http://www.ensembl.org)
Ensembl produces genomic datasets through a system that is designed to analyse, store and distribute data, and which enables interpretation through open data release. As a hub of reference and baseline data similar to UCSC Genome Browser and RefSeq, Ensembl also distributes created datasets and promotes standards and interoperability between genomic resources. In addition, Ensembl collaborates with and often plays active leadership roles in projects such as ENCODE, the "Genome Reference Consortium" (GRC), the "Global Alliance for Genomics and Health” (GA4GH) and GENCODE. Ensembl is updated 4-5 times annually with each release representing a data and software freeze. Ensembl provides two sets of human data based on the hg19 genome build (http://grch37.ensembl.org/Homo_sapiens/Info/Index) which has been updated by the data set based on the December 2013 Homo sapiens high coverage assembly GRCh38 from the Genome Reference Consortium. This assembly is used by UCSC to create their hg38 database. The data set consists of gene models built from the alignments (for comparison) of the human proteome as well as from alignments of human cDNAs. This release of the assembly has the following properties: assembly length with a total of 3.4 Gb, chromosome length total 3.1 Gb (excluding haplotypes). It also includes 261 alternate loci scaffolds, mainly in the LRC/KIR complex on chromosome 19 (35 alternate sequence representations) and the MHC region on chromosome 6 (7 alternate sequence representations) (Yates A et al., 2016).

VI- Structural variation databases
Since the mid 2000's, there were several studies of copy number variation of DNA sequences to construct CNV map of the human genome through different populations using SNP genotypes and CGH (Iafrate AJ et al., 2004; Redon R et al., 2006). It is becoming clear that genomic structural variation (variation ranging from tens to millions of base pairs in size, and including insertions, deletions, inversions, translocations and locus copy number changes) accounts for individual differences at the DNA sequence level in humans and can play a major role in diseases. Many databases have integrated data produced in the literature.

VI-1 dbVar (http://www.ncbi.nlm.nih.gov/dbvar/)
dbVar is the NCBI's database of genomic structural variation. It contains data of insertions, deletions, duplications, inversions, multi-nucleotide substitutions, mobile element insertions, translocations, and complex chromosomal rearrangements (NCBI Resource Coordinators, 2016).

VI-2 DGV - Genomic Variants (http://dgv.tcag.ca/dgv/app/home)
DGV is a database with an objective to provide a comprehensive summary of structural variation in the human genome. Structural variation is defined as genomic alterations that involve segments of
DNA that are larger than 1kb. It also annotates InDels in 100bp-1kb range. The content of the database is only representing structural variations identified in healthy control samples (MacDonald JR et al., 2014).

**VI-3 DECIPHER**
(http://decipher.sanger.ac.uk/)

DECIPHER (DatabaSE of Genomic variants and Phenotype in Humans using Ensembl Resources) is an interactive web-based database which incorporates a series of tools designed to aid the interpretation of genomic variants. DECIPHER enhances clinical diagnosis by retrieving information from a variety of bioinformatics resources relevant to the variant found in a patient. The patient's variant is displayed in the context of both normal variation and pathogenic variation reported at that locus, thereby facilitating interpretation (Firth HV et al., 2009).

**VI-4 1000 Genomes**
(http://www.1000genomes.org/)

The 1000 Genomes Project benefitted from the progress in sequencing technology, which sharply reduced the cost of sequencing. It was the first project to sequence the genomes of a large number of people, to provide a comprehensive resource on human genetic variation. Data from the 1000 Genomes Project was quickly made available to the worldwide scientific community through freely accessible public databases.

In continuation of the 1000 Genome project (sequencing 1000 human genome as exomes or whole genomes), the International Genome Sample Resource (IGSR) aims to expand information to new populations, a better coverage for presenting a uniform analysis set. Data corresponds to both single nucleotide and structural variants (1000 Genomes Project Consortium et al., 2015). The 1000 Genomes Project operated between 2008 and 2015, creating the largest public catalogue of human variation and genotype data. As the project ended, the Data Coordination Centre at EMBL-EBI received continuous funding from the Wellcome Trust to maintain and expand the resource. The International Genome Sample Resource (IGSR) is maintaining and extending the 1000 Genomes Project data.

**VII- Polymorphism databases**

It is important to distinguish between polymorphisms due to a change in a single nucleotide (SNP) as the variability within a population and mutations acquired in a neoplastic process. The determination of variants was previously obtained by SNP arrays, but is nowadays performed by massive parallel sequencing. As a result, a huge quantity of polymorphisms and mutations in tumors are compared to controls. The landscape of the majority of recurrent mutations is now known and can be used for diagnosis.

**VII-1 dbSNP**

dbSNP is the main repository of Single Nucleotide Polymorphisms: A key aspect of research in genetics is associating sequence variations with heritable phenotypes. The most common variations are single nucleotide polymorphisms (SNPs), which occur approximately once every 100 to 300 bases. Because SNPs are expected to facilitate large-scale association genetics studies, there has recently been great interest in SNP discovery and detection. The database contains 164,986,514 SNPs for several species (NCBI Resource Coordinators, 2016).

**VII- 2 HAPMAP**

The International HapMap Project was a collaboration of scientists and funding agencies from Canada, China, Japan, Nigeria, the United Kingdom and the United States who wanted to develop a public resource that helps researchers find genes associated with human disease and consequently give response to pharmaceuticals. The goal of the project was to compare genetic sequences of different individuals in order to identify chromosomal regions where genetic variants are shared. An interface (http://hapmap.ncbi.nlm.nih.gov/cgi-perl/gbrowse/hapmap28B36/) permits to query all the data collected in phases 1, 2 and 3 of the project (International HapMap 3 Consortium et al., 2010).

**VII-3 1000 Genomes Project (see above)**

As the Phase3, 1000 Genomes variants are in the process of being archived at dbSNP and DGVa and a version of the Ensembl databases has been created, containing the phase3 autosomal variants. This is presented alongside the v80 GRCh37 Ensembl core and regulatory databases. This release represents more than 80M short variants with genotypes for 2,504 individuals across 26 populations. The latest major update was released to the 1000 Genomes Website in February 2016 (1000 Genomes Project Consortium et al., 2015).

**VII- 4 Exome Variant server (EVS)**
(http://evs.gs.washington.edu/EVS/)

The goal of the NHLBI GO Exome Sequencing Project (ESP) is to discover novel genes and mechanisms contributing to heart, lung, and blood
disorders by pioneering the application of next-generation sequencing of the protein coding regions of the human genome across diverse, richly-phenotyped populations and to share these datasets and findings with the scientific community to extend and enrich the diagnosis, management and treatment of the aforementioned disorders. Two categories of populations are considered: European-American and African-American. Some criteria or impact scores of the variation on the gene function are also presented (Tennessen JA et al., 2012).

**VIII- Portals/Working consortium**

**VIII-1 TCGA**

(https://cancergenome.nih.gov/)

Since 2005 TCGA (The Cancer Genome Atlas) has indexed genetic mutations responsible for cancer, using genome sequencing and bioinformatics. TCGA applies high-throughput genome analysis to progress our ability to diagnose, treat, and prevent cancer.

TCGA is administered by the National Cancer Institute’s Center for Cancer Genomics and the National Human Genome Research Institute funded by the US government. A pilot project, initiated in 2006, focused on analysing three types of human cancers: Glioblastoma multiforme, lung cancer, and Ovarian cancer (Cancer Genome Atlas Research Network, 2011).

In 2009, a second phase started, 20-25 different tumor types were included to complete the genomic characterization and sequence analysis (Figure 11). TCGA surpassed that goal, characterizing 33 different cancer types including 10 rare cancers (http://cancergenome.nih.gov/abouttcga/overview). Funding is split between genome characterization centers (GCCs), which perform the sequencing, and genome data analysis centers (GDACs), which perform the bioinformatic analyses.

The project scheduled 500 patient samples using several analysing techniques: Gene expression profiling, copy number variation profiling, SNP genotyping, genome wide DNA methylation profiling, microRNA profiling, and exon sequencing of 1,200 or more genes (Figure 12). TCGA is sequencing some tumors, including at least 6,000 candidate genes and microRNA sequences.

This targeted sequencing is being performed by all three sequencing centers using hybrid-capture technology. In phase II, TCGA is performing whole exon sequencing on 80% of the cases and whole genome sequencing on 80% of the cases used in the project.

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**Figure 11:** Acute Myeloid Leukemia query in TCGA datasets with the Data Matrix option (https://tcga-data.nci.nih.gov/tcga/dataAccessMatrix.htm?mode=ApplyFilter)
ICGC (The International Cancer Genome Consortium) was organized to launch and coordinate a large number of research projects with the common aim of comprehensively elucidating the genomic changes present in many forms of cancers. Funding and Research members proposing a project must agree to the ICGC’s policies (Figure 13). ICGC’s primary objectives are to generate comprehensive catalogues of genomic abnormalities (somatic mutations, abnormal expression of genes, epigenetic modifications) in tumors representing 50 different cancer types and/or subtypes which are of clinical and societal importance across the globe and make the data available to the entire research community. Each of the 50 projects will generate the genomic analyses on approximately 500 cancer samples of each class. This will cover the various types and subtypes but cannot exhaustively cover the full spectrum of cancer types. The ICGC facilitates communication among the members and provides a forum for coordination with the objective of maximizing efficiency among the scientists working to understand, treat, and prevent these diseases.


OASIS (http://www.oasis-genomics.org/) OASIS, which was created by Pfizer Oncology Research Computational Biology in collaboration with Research Business Technology (RBT), is an open-access web portal that provides the possibility to run exploratory and integrative analyses of somatic mutations, copy number variation (CNV) and gene expression data (Figure14). This data originates from thousands of different tissues of tumour samples, normal tissues and cell lines thus representing a broad spectrum of malignancies. This portal contains 30 datasets, mainly from TCGA, with access to mutations, copy number variation, expression (microarrays) and expression (RNA-Seq).
Figure 13: ICGC International Cancer Genome consortium: Home page (https://icgc.org/)
VIII-4 Firebrowse (http://firebrowse.org/)
This portal developed at the Broad Institute presents 38 cancer cohorts and 14,729 samples, mainly from the TCGA program, and provides an option to browse reports, clinical analysis, copy number variation, mutation, expression, and to download data for further analysis (Figure 15). See the tutorial for a complete view of possibilities (http://firebrowse.org/tutorial/FireBrowse-Tutorial.pdf).

VIII-5 GDC (https://gdc.nci.nih.gov/)
The NCI's Genomic Data Commons (GDC) provides the cancer research community with a unified data repository that enables data sharing across cancer genomic studies in support of precision medicine. (note added in proof, June 6, 2016). The GDC supports several cancer genome programs at the NCI Center for Cancer Genomics (CCG), including The Cancer Genome Atlas (TCGA), Therapeutically Applicable Research to Generate Effective Treatments (TARGET), and the Cancer Genome Characterization Initiative (CGCI). The GDC Data Portal provides a platform for efficiently querying and downloading high quality and complete data. The GDC also provides a GDC Data Transfer Tool and a GDC API for programmatic access.

IX- Impact on diseases
IX-1 OMIM
"Mendelian Inheritance in Man: Catalogs of Autosomal Dominant, Autosomal Recessive and X-linked Phenotypes" was first published in 1966 by Victor A. McKusick (Johns Hopkins University Press), after a catalog of X-linked traits, published in 1962. In parallel, the "Human Gene Mapping" was first organized in New Haven in 1973, and mapped 119 and 100 loci respectively to confirmed or provisional/tentative chromosome assignments (Birth Defect, 1974).
The first edition of the "Mendelian Inheritance in Man: Catalogs of Autosomal Dominant, Autosomal Recessive and X-linked Phenotypes" had 1487 entries and no mapped autosomal loci. Victor A. McKusick published 12 editions, the last one in 1998, of his catalog. "Online Mendelian Inheritance in Man" (OMIM, http://omim.org/) was consequently published online. OMIM is a continuously updated catalog of human genes and genetic disorders and traits, with particular focus on the molecular relationship between genetic variation and phenotypic expression (Amberger JS et al., 2015).
As of April 2016, it consists of 23,460 entries: 15,237 gene descriptions, 4,705 phenotypes with known molecular basis, an additional 1,626 phenotypes with unknown molecular basis, and 1892 other entries. Gene entries start at: * 100640. Aldehyde dehydrogenase 1 family, member A1; ALDH1A1 Cytogenetic location: 9q21.13, Genomic coordinates (GRCh38): 9:72,900,661-72,953,316, and ends with * 616906. Cancer susceptibility candidate 1; CASC1. Phenotypes with known molecular basis entries start at: # 100100. Prune belly syndrome; PBS, Cytogenetic location: 1q43, and ends with # 616903. Nucleoside diphosphate-linked moiety X motif 15 deficiency; NUDT15D. OMIM describes somatic mutations in genes (11,139 entries for the term "mutation"). It is a very well curated database, with excellent reliability. Unfortunately the addition process of data as literature is published, by successive layerings/sedimentation makes it sometimes a laborious consultation. OMIM is authored and edited at the McKusick-Nathans Institute of Genetic Medicine, Johns Hopkins University School of Medicine.


ClinVar is designed to provide a public archive of reports of the relationships among human variations and phenotypes, with supporting evidence. By doing so, ClinVar facilitates access to and communication about the relationships asserted between human variation and observed health status, and the history of that interpretation. ClinVar collects reports of variants found in patient samples, and assertions made regarding their clinical significance. The alleles described in submissions are mapped to reference sequences, and reported according to the HGVS standard. ClinVar then presents the data for interactive users in daily workflow and other local applications. ClinVar works in collaboration with interested organizations to meet the needs of the medical genetics community as efficiently and effectively as possible (Harrison SM et al., 2016).


MedGen is an NCBI portal of information about human disorders and other phenotypes having a genetic component. MedGen is structured to serve health care professionals, medical genetics community and other interested parties by providing centralized access to diverse content. MedGen aggregates the plethora of terms used for particular disorders into a specific concept, providing a "Rosetta stone" for stakeholders who may use different names for the same disorder.
Maintaining a clearly defined set of concepts and terms for phenotypes is essential in supporting characterization of genetic variation by its specific phenotypes effect. The assignment of identifiers for those concepts allows computational access to phenotypic information, an essential requirement for the large-scale analysis of genomic data. (NCBI Resource Coordinators, 2016).

The database of Genotypes and Phenotypes (dbGaP) was developed to archive and distribute the data and results from studies where the interaction of genotype and phenotype in Humans has been investigated.

IX-5 SNPs3D (http://www.snps3d.org/)
SNPs3D is a website which assigns molecular functional effects of non-synonymous SNPs based on structure and sequence analysis. The site presents a data mining method to infer candidate SNP for 16 types of cancer (e.g. more than 1,000 genes potentially implicated in breast cancer: http://www.snps3d.org/modules.php?name=Candidate&disease=BREAST%20CANCER) (Yue P and Moulit J, 2006).

IX-6 GTR (http://www.ncbi.nlm.nih.gov/gtr/)
The Genetic Testing Registry (GTR) provides a central location for voluntary submission of genetic test information by providers. The scope includes purpose of the test, methodology, validity, evidence of its usefulness and laboratory contacts and credentials. The overarching goal of the GTR is to advance public health research to include the genetic basis of health and disease (Rubinstein WS et al., 2013).

IX-7 ClinGen (https://www.clinicalgenome.org/)
ClinGen is a National Institutes of Health (NIH)-funded resource dedicated to building an authoritative central resource that defines the clinical relevance of genes and variants for use in precision medicine and research. This resource has several goals for building a genomic knowledge base to improve patients care.

X- Pathology

X-1 Authoritative books in pathology are the following:
The "Rosai and Ackerman's Surgical Pathology" was first published in 1953. The tenth edition was published in 2011 by Elsevier, and contains 2892 pages. It includes clinical features, morphologic, immunohistochemical and molecular genetic features and prognosis, with a very large iconography. "WHO/IARC Classification of Tumours series" (http://publications.iarc.fr/Book-And-Report-Series/Who-Iarc-Classification-Of-Tumours) is not on free access, except editions prior to 2006, which are on free access in pdf format. The Armed Forces Institute of Pathology (AFIP) publishes series of the "AFIP Atlas of Tumor Pathology". The WHO/OMS code, the ICD, genetic registries are organizations seeking to collect, store, analyze, and report data on various cancers for epidemiological purposes, for providing statistics on the occurrence of cancer in a defined population, and for obtaining a framework to assess the impact of cancer in a given population. Cancer registries are crucial for healthcare policy planning. They are key data source for clinical research, (epidemiology, study of carcinogens, evaluation of
treatments), providing the assessment of the care structures and care pathways, and research tools for social sciences and humanities (see https://www.iarc.fr/en/publications/pdfs-online/epi/cancerepi/CancerEpi-17.pdf)


The IARC is the outcome of an initiative by a group of leading French public figures; it was created on 20 May 1965, by a resolution of the World Health Assembly (http://www.iarc.fr/en/about/iarc-history.php). IARC is the specialized cancer agency of the World Health Organization (WHO/OMS). The objective of the IARC is to promote international collaboration in cancer research. The Agency is inter-disciplinary. Emphasis is placed on elucidating the role of environmental and lifestyle risk factors and studying their interplay with genetic background. IARC publishes the “Cancer Incidence in Five Continents” series and GLOBOCAN (Figure 16). The aim of the GLOBCAN project (http://globocan.iarc.fr/Default.aspx) is to provide contemporary estimates of the incidences of, mortality and prevalence of major types of cancer, at national level, for 184 countries of the world.

**XI-2 International Association of Cancer Registries (IACR, http://www.iacr.com.fr/)**. The IACR (not to be confused with the IARC) was founded in 1966 as a professional society dedicated to fostering the aims and activities of cancer registries. It is a non-governmental organization which has had official relation with the World Health Organization since January 1979. With IACR IARC has developed with CanReg5, an open source tool to input, store, check and analyze cancer registry data. IACR has developed classifications (the successive editions of the International Classification of Diseases for Oncology, published by WHO), guidelines for registry practices and standard definitions, quality control, consistency checks and basic analysis of data, making data comparable between registries.

**XI-3 Examples: The European Network of Cancer Registries (ENCR, http://www.enccr.eu/)** has the same role in Europe as IACR has worldwide. The National Program of Cancer Registries (NPC, http://www.cdc.gov/cancer/), maintained by the Centers for disease control and prevention (CDC), collects data on cancer occurrence (including the type, extent, and location of the cancer), the type of initial treatment, and outcomes in the USA. The Surveillance, Epidemiology, and End Results (SEER, http://seer.cancer.gov/) program of the National Cancer Institute provides information on cancer. Research is supported by grants from the SEER. Quality improvement is another part of the SEER activities and it is dedicated to improving data quality by performing rigorous quality control studies and various data assessments. Union for International Cancer Control (UICC, http://www.uicc.org/). Founded in 1933, UICC brings together 900 organisations (cancer societies, ministries of health, research institutes and patient groups) over across 155 countries.

**XII- Patient associations and interfaces between science and patients - freely accessible services**

**XII-1 Associations of parents and friends of patients**

These associations of parents of patients with a rare disease are precious. They give moral support and help, and offer practical guidances and information about social benefits, subsidies and day-to-day life to families affected by illness. They often establish a program of grants for research (e.g. Xeroderma Pigmentosum Society (http://www.xps.org/), Sarcoma Foundation of America (http://www.curesarcoma.org/), Union for International Cancer Control (UICC (http://www.uicc.org/)).

**XII-2 interfaces between science and patients**

**GeneTests** (https://www.genetests.org)


**NORD** (http://rarediseases.org)

NORD provides information on more than 1,300 rare diseases (e.g. Carcinoid syndrome http://rarediseases.org/rare-diseases/carcinoid-syndrome/), state health insurance information, guides for physicians, and patient assistance programs. They also provide grants to academic scientists for translational or clinical studies to help patients obtain life-saving or life-sustaining medication they could not otherwise afford.

**Orphanet** (http://www.orpha.net/)

Orphanet (Rath et al., 2012) offers an inventory of rare diseases with data on 5,833 diseases (e.g. Fanconi anaemia), an inventory of orphan drugs, list of 6,636 expert centres and 3,280 laboratories,
19,894 professionals for genetic counselling and medical management. Orphanet does not provide gene annotations. They hold a large partnership from 38 countries participating in the Orphanet consortium. They maintain large disease registries in Europe.

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