Original article

Data mining using the Catalogue of Somatic Mutations in Cancer BioMart

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Catalogue of Somatic Mutations in Cancer (COSMIC) (http://www.sanger.ac.uk/cosmic) is a publicly available resource providing information on somatic mutations implicated in human cancer. Release v51 (January 2011) includes data from just over 19 000 genes, 161 787 coding mutations and 5573 gene fusions, described in more than 577 000 tumour samples. COSMICMart (COSMIC BioMart) provides a flexible way to mine these data and combine somatic mutations with other biological relevant data sets. This article describes the data available in COSMIC along with examples of how to successfully mine and integrate data sets using COSMICMart.

Database URL: http://www.sanger.ac.uk/genetics/CGP/cosmic/biomart/martview/

Project description

COSMIC is a repository for somatic mutations and associated phenotype/clinical information, combining data from a number of sources. Primarily, data are manually curated from the scientific literature for genes selected from the Cancer Gene Census (http://www.sanger.ac.uk/genetics/CGP/ Census/), a listing of genes that are known to be mutated in human cancer. The manual curation initially focused on known cancer genes that had a high proportion of coding point mutations. In recent years, this has been extended to include cancer genes that are also mutated by gene fusion events. With the advent of next-generation sequencing, COSMIC has been adapted to hold complete catalogues of somatic mutations for individual tumour samples.

The COSMIC website has been developed to overview the underlying database in a user-friendly manner. The website can be navigated by gene, cancer sample or tissue/histology type and has a series of graphical and tabular displays to summarize the content of the database.

However, the integration of biological data sets is still a major IT challenge. Although, the data in COSMIC can be

viewed and downloaded in a number of useful ways, it is still challenging for users to generate their own custom queries and also to link to related resources. In order to facilitate the integration of the COSMIC data set, we have used the BioMart data mining software. BioMart uses a federated data model to allow integration of biological data from diverse databases (1). There is now a significant number of data resources that have set up their own BioMarts including Ensembl, UniProt and HGNC. The BioMart software also has interfaces that allow users to easily generate custom queries to obtain subsets of data from one or more BioMart data resources. We have successfully set up an instance of a BioMart, COSMICMart, which holds a summary of the somatic mutations and associated phenotype data from the COSMIC database.

Data content

COSMIC version 51 (January 2011) contains a full curation of the scientific literature for 91 cancer genes, mostly point-mutated, as well as 53 curated fusion gene pairs, with over 11000 papers having been assessed. Weekly

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literature searches are carried out for each cancer gene and each abstract/article is manually evaluated. Checks are made to ensure a paper has not been curated already, whether the paper is a review or is reporting original data, if there are any inconsistencies in the data, and whether all the required information for full curation is present, e.g. information on the numbers of samples screened and sufficient mutation detail. Papers which pass the initial scan are submitted for detailed manual curation, while the remaining papers are 'listed' on the COSMIC website. Information is manually recorded on the actual mutation change, the detailed phenotype of the cancer samples and the original publication/study.

Mutation changes are checked for accuracy and recorded at the genome, transcript (to one reference transcript) and protein level using the mutation syntax standards developed by the Human Genome Variation Society (2). Negative results (sample screened does not have a mutation for a particular gene for a particular publication) are also recorded so prevalence statistics can be estimated for each gene. As part of the curation process, cancer sample tissue/histology descriptions from the original publication are recorded and then re-classified to a COSMIC standard tissue/histology ontology (See classification section of the COSMIC additional information page, http://www.sanger .ac.uk/genetics/CGP/cosmic/add_info/). The standardization of the main data types in COSMIC means the utility of the data is significantly enhanced and the data are easier to browse and incorporate into external data sets. When available, clinical and exposure data and therapeutic responses are also recorded. The full functionality of COSMIC has previously been described (3, 4).

The database coverage has recently been enhanced by the integration of somatic mutations from external data resources. A collaboration with the database curators at International Agency for Research on Cancer (IARC) has allowed the majority of mutations from IARC TP53 database R14 to be integrated into COSMIC (5). Somatic mutations from large-scale systematic cancer screens are curated in COSMIC to include studies on glioblastoma multiforme, breast, colorectal and lung cancer (6-9). Data are also directly curated from the data portals of large-scale projects to include the Cancer Genome Project (CGP) at the Sanger Institute and more recently validated somatic mutations from The Cancer Genome Atlas (TCGA) (6) and International Cancer Genome Consortium (ICGC) (10). COSMIC has been enhanced to accommodate somatic mutations from whole genome and exome screens. This includes all somatic and non-coding mutations, structural rearrangements and gene fusions. Currently, COSMIC has annotated 51 genomes and 332 exomes from a range of cancers to include lung, malignant melanoma, renal, AML, pancreatic and ovarian cancer. The current contents of the database (v51, January 2011) are displayed in Table 1.

Table 1. Total contents in v51 of the COSMIC database,January 2011 release

Curated data type	Curated data count			
Experiments	2 946 792			
Tumours	577 304			
Mutations	167 193			
References	11 062			
Genes	19 000			
Fusions	5573			
Structural variants	2729			
Whole-cancer genomes	51			
Whole-cancer exomes	332			

Query examples

COSMICMart allows data to be filtered on six different categories (Figure 1): cancer sample, gene, mutation, site of the tumour, histology and other (e.g. Ensembl Gene ID, Swissprot ID, Entrez Gene ID). The interface has a number of pre-selected filters and attributes; mutated samples are selected by default. Users can change these to suit their requirements. Results are displayed in tabulated form and are exportable in various formats for further analysis.

Query #1: 'Find all missense substitution mutations for BRAF in cell lines, and display sample, mutation, site, and histology information' (Figure 1, Table 2).

Missense mutations are the most common variant type in COSMIC; over 90% of mutations in BRAF are missense mutations at the p.V600 position. The results are returned as a tabular summary with links back to the COSMIC website. The sample name field links back to the COSMIC website. The sample name field links back to the COSMIC sample overview web page, and mutation ID (COSM ID) to the COSMIC mutation summary page (Figure 2). From the COSMIC mutation summary web page, there are links to the Ensembl contig view so the mutation can be viewed in a genomic context. There are also links to the GMOD's GBrowse where COSMIC coding and non-coding mutations, gene footprints, structural rearrangements and copy number variants can be viewed (11).

Query #2: 'Find all gene fusion mutations involving the FUS gene with a primary site of bone, and display mutation and sample information' (Table 3).

Gene fusions have been associated with a number of specific tumour types including prostate and blood tumours. These biomarkers can be useful in diagnosis and as targets for drug therapies. COSMIC has annotations, for an increasing number of gene fusion mutations, which are viewable using COSMICMart. The COSMIC fusion mutation ID links to the gene fusion summary pages, which give a graphical view of different fusion structures observed. Many of the papers describing gene fusions have identified

Α		— В
SAMPLE		Please select columns to be included in the output and hit 'Results' when ready
COSMIC Sample ID		B SAMPLE
Sample Name		COSMIC Sample ID Sample Source
Sample Source	blood-bone marrow cell-line cerebrospinal fluid fine meedle aspirate	a GENE
Mutated Sample	e Yes	Gene ID Accession Number Gene Name In Cancer Census
Tumour Source	NS hyperplasia adjacent to primary tumour metastasis primary recurrent	HUTATION GOSMIC Mutation ID CDS MIC Fusion Mutation ID COSMIC Fusion Mutation ID Genomic Coordinates (ROEI36) Genomic Coordinates (ROEI36) Genomic Coordinates (ROEI37) CAA Mutation Syntax
© GENE		CDS Mutation Type AA Mutation Stop
🗆 Gene ID		CDS Mutation Syntax Zygosity
i Gene Name	BRAF	B SITE ■ Primary Site Subtype 2 Site Subtype 1 Site Subtype 3
	Browse	E HISTOLOGY
Accession Number		
In Cancer Census	● Yes ○ No	Primary Histology Histology Histology Subtype 2 Histology Subtype 1 Histology Subtype 3
E MUTATION		BTUMOUR
COSMIC Mutation ID		I Tumour Source
COSMIC Fusion Mutation ID		BOTHER
Genomic Coordinates (NCBI36)(chr:start-end)		Entrez Gene ID Swissprot ID CSSMIC Study ID
Genomic Coordinates (GRCh37)(chr:start-end)		Ensembl Gene ID Whole Genome Screen
CDS Mutation Syntax		T and the second
CDS Mutation Type	Inferred Breakpoint Observed mRNA Complex - compound substitution - Complex - intronic	C Export all results to File STSV (2) Unique results only Co
CDS Mutation Start		Email notification to
CDS Mutation Stop		View 10 1 rows as HTML 1 Unique results only
I AA Mutation Type	Nonstop extension	Sample Name Game Source COSNIC COSNIC COS AA Mutation Primary Site Primary Histology Pubmed Tumour Source Sci 20 cell-me EAA 57 C17997A Pv600E tbin material Total Source 128/277 NS VM233 cell-me EAA 57 C17997A Pv600E tbin material Total/meterial 128/277 NS VM233 cell-me EAA 57 C17997A Pv600E tbin material/meterial 128/3721 primary VM233 cell-me EAA 57 C17997A Pv600E tbin material/meterial 128/3721 primary VM239 cell-me EAA 57 C17997A Pv600E tbin material/meterial 128/3721 primary
AA Mutation Syntax		451Lu cell-line BRAF 475 c.1799T>A p.V600E skin malignant melanoma 12581721 metastasis SKMet5 cell-line BRAF 475 c.1799T>A p.V600E skin malignant melanoma 12691721 metastasis
AA Mutation Start		NCH4508 cell-line BRAF 489 c.1796GoC p.G596R large integline carcinoma 12068308 NS KG-1-C cell-line BRAF 476 c.17997-XA p.V600E central nencus system gloma 12068308 NS
AA Mutation Stop		COLO-679 cell-Ine URAF 425 c.179915-A p.V600E stin malignant. melanoma 12068308 NS G-361 cell-Ine BRAF 426 c.179915-A p.V600E stin malignant. melanoma 12068308 NS
Zygosity	Heterozygous	
Systematic Screen	e Yes ⊙No	

Figure 1. Example of how COSMICMart can be queried. This query searches for all cell lines with missense substitution mutations in the BRAF gene (A). Attributes can be selected (B) to display in the results table (C).

Table 2. Data sets, filters and attributes selected for query #1

Data sets	Filters	Attributes
COSMIC51	Mutated sample: yes	Sample name
	Sample source: cell-line	Sample source
	Gene name: BRAF	Gene name
	AA mutation type: substitution—missense	Cosmic mutation ID (COSM ID)
		CDS mutation syntax
		AA mutation syntax
		Primary site
		Primary histology
		Tumour source
		Pubmed ID

Table 3.	Data se	ts, filters	and	attributes	selected	for query #	2
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Data sets	Filters	Attributes
COSMIC51	Mutated sample: yes Gene name: FUS	Cosmic sample ID Sample name
	CDS mutation type: inferred breakpoint, observed mRNA	Sample source
	Primary site: bone	Cosmic fusion mutation ID CDS mutation syntax Pubmed ID

more than one gene fusion product for the same genes in a single sample. Observed mRNAs are the actual expressed products reported in the results. However, to aid display

and website navigation, we have inferred the genomic breakpoint from the experimental data.

Query #3: 'Find variation information in Ensembl for all genes from mutated samples with a primary site of breast, and display COSMIC gene, mutation and sample information along with Ensembl variation information' (Table 4).

1

1							
Sample Name	S64003						
COSMIC id	685178						
Source	Cultured Samp	le (NS; cell-line)					
Tumour Classification	Martin Color	ary Site; Subtype1 ary Histology; Sub		and the second second		Entry NS; NS; NS mant melanoma;	NS; NS; NS
Mutations	Sample Gene		DS ation <u>99T>A</u> Repo	427 	Somatic Status	24	Zygosity Validated
References	melano Dong J,	Phelps RG, Qiao Res. 2003;63;388	R, Yao S, B	Benard O, Ro			n of human
ndividual				Sample: 6	85178		
		Age:	Unknown				
		Gen	der: Unknow	/n			
		Pare	nts tested: L	Jnknown			
		Fam	ly: Unknown	n			
Mutation id	476						
AA Mutation	p.V600E (Substitution - Misse	nse)				
CDS Mutation	c.1799T>/	A (Substitution)					
Mutation Loca	tion Length (A	AD 🔫		- 70	57.00 AA		•
Genomic View	Clickhere to st GRCh37 : <u>Clickhere</u> to st	A)	w 7:14 notpreviously use w 7:14 notpreviously use	G D F GL A 1009960514 10009800 DAS 1000800 DAS 10045313614	0099605 10453136	er Genome Browse	
Gene	BRAF						
Tissue Distribu (Top 5)	tion Tissues scale (ftu	L large L skin L NS (2 L ovary	24)	2,000 265) 2,000	2,666	4,011	5,000
Associated	Sample	Name Primary	Tissue	Zygosity	Sample Nam	e Primary Tiss	ue Zygosity
Samples	10	skin		Jnknown	1011-mel	NS	Unknown
	and a state of the	large intestir		Jnknown	1020356	large intestine	Unknown
	1020355	large intestir	-	JURIOWIT	1020300	range intestine	Contra to the
	<u>1020355</u> <u>1020357</u>	large intestin	10	Jnknown	1020330	thyroid	Unknown

Figure 2. The COSMIC sample (A) and mutation (B) summary pages are linked directly from COSMICMart output table.

Table 4. Data sets, filters and attributes selected for query #3

Data sets	Filters	Attributes
COSMIC51	Mutated sample: yes	Cosmic sample ID
	Primary site: breast	Sample name
		Sample source
		Cosmic mutation ID (COSM ID)
		CDS mutation syntax
		AA mutation syntax
Ensembl: <i>Homo</i> sapiens genes		Features: Ensembl gene ID
		Features: Ensembl transcript ID
		Variations: variation source
		Variations: source description
		Variations: reference ID
		Variations: allele

COSMICMart is federated with Ensembl (12), which allows Biomart gueries to return and integrate data from both resources. For instance, the linking of the two resources can allow the retrieval of variation data from both resources (somatic mutations from COSMIC and germline polymorphisms from Ensembl) for a particular gene or set of genes or cancer type. There is an increasing awareness of how genomic variation can affect a tumour's sensitivity or resistance to anti-cancer agents. While this genetic variation can be familial or somatic, an understanding of common genetic variation around known cancer genes can be of much value to investigations searching for loci modifying a tumour's response to drug therapy (13-15). This query is achieved by first selecting the filters/attributes in the COSMIC BioMart and then clicking the 'Dataset' link at the bottom of the left hand margin of the BioMart interface. An additional data set can then be selected from the drop down list, in this instance Ensembl, which allows a federated query between COSMIC and Ensembl. The filters/attributes are then set in the usual way using the Ensembl BioMart to produce an integrated guery.

Future directions

COSMIC will continue to curate newly discovered cancer genes and is committed to update existing cancer genes with a data release every 2 months. This will ensure that the scientific community has an up-to-date catalogue of somatic mutations implicated in human cancer. COSMICMart is also automatically updated with each new COSMIC release, which allows the data set to be easily mined and integrated with other resources. COSMIC has been successfully adapted to hold complete catalogues of somatic mutations for individual cancer samples. Currently COSMIC holds genome-wide data on 383 tumour samples and we expect this to increase in the near future.

It is intended to federate COSMICMart with further BioMart-driven data resources in addition to the current link with Ensembl. Linking our data to PRIDE (16), UniProt (17) and InterPro (18) will allow COSMIC somatic mutation data to be linked to protein and peptide annotation, while the addition of the Reactome (19) database will allow the incorporation of pathway data. We also intend to create direct links between COSMIC and the ICGC Data Portal (http://dcc.icgc.org/) so somatic mutation data can be integrated between the two data resources.

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