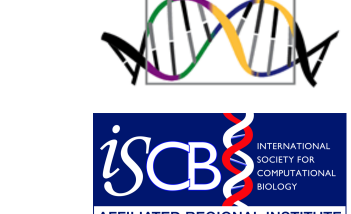


# SELECTION OF PRINCIPAL SPLICE ISOFORMS



http://appris.bioinfo.cnio.es

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### **ABSTRACT**

The cellular role of alternative protein isoforms is a topic of growing interest. We have developed the APPRIS database (1) and APPRIS Webserver and Web Services (2) to annotate splice variants with information relating to protein structure, function and cross-species conservation.

APPRIS makes use of the conservation of protein features to identify a single dominant (3) isoform for each gene. These principal isoforms are confirmed by orthogonal theoretical analyses (4) and by the results of multiple large-scale mass spectrometry experiments and databases (5,6).

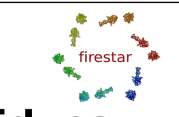
APPRIS is stable and is implemented as part of the GENCODE/Ensembl human genome annotation (7).

APPRIS has recently been expanded to *mouse*, *rat*, *pig* and *zebra fish*, and is being expanded to *chimpanzee*, *Drosophila* and *C. elegans*.

**Principal Isoform** 

### **METHODS**

### firestar<sup>(8)</sup>



### **Functionally Important Residues**

Functional residues are highly conserved, even across large evolutionary distances.

Since these residues are **unlikely to have** arisen by chance we can use this to help determine the principal isoform.

Seq. id

http://firedb.bioinfo.cnio.es/Php/FireStar.php

### Matador3D



#### **Protein structural information**

Codons not found | CCDS

Since protein structure is much more conserved than sequence variants with large inserts or deletions relative to their crystal structures are also not likely to be the principal isoform.

# **BLAST** against vertebrates

Transcripts **conserved** over greater evolutionary distances are more likely to be the principal variant. Good alignments with more distant relatives (Danio, Xenopus, Chicken) are regarded as more valuable.

CORSAIR

The more species that align correctly and without gaps, the better.

### SPADE

Pfam

#### **Conservation of protein** functional domains

Identifying the functional domains present in a variant can provide insights into the function.

The presence of protein domains is analysed with **Pfamscan** (9).

### SELECTION OF PRINCIPAL ISOFORM FOR *DNAJC5G*

	EN	ST00000296097	DNAJC5G-00	1 189	protein_	coding	-		CCDS1744.1	MINOR
	EN	ST00000402462	DNAJC5G-00	2 189	protein_	coding	-		CCDS1744.1	MINOR
	EN	ST00000404433	DNAJC5G-00	4 173	protein_	coding	-		-	PRINCIPAL:1
	ENST00000406962		DNAJC5G-00	3 104	protein_coding		-		-	MINOR
ENST		ST00000420191	DNAJC5G-00	7 62	protein_	coding	stop		-	MINOR
Seq. id		Seq. name	Length (aa)	No. Functional	Residues	3D Str	ucture Score	Whole	e Domains	
ENST0000029	6097	DNAJC5G-001	189	0		1.8		0		
ENST00000402462		DNAJC5G-002	189	0		1.8	1.8			
ENST00000404433		DNAJC5G-004	173	0		1.8		1		
ENST00000406962		DNAJC5G-003	104	2		0.8	).8			
ENST00000420191		DNAJC5G-007	62	0		0.8	0.8			
	<b>c</b>	4.00016								

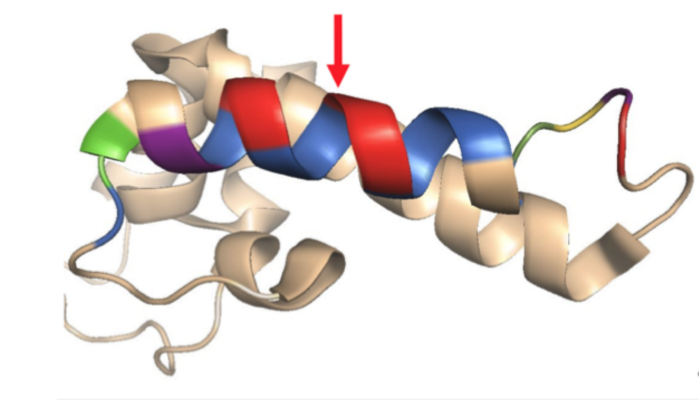
Length (aa) Biotype

Snapshot of the APPRIS web page, showing the five protein-coding transcripts annotated by GENCODE/Ensembl and the selection of the principal isoform by APPRIS (green). The variant selected by APPRIS (DNAJC5G-004) has a conserved Pfam domain.

The highlighted methods SPADE and Matador3D map Pfam functional domains, protein structure to the splice isoforms.

The principal isoform for *DNAJC5G* has 16 fewer residues than the **longest isoform**, which has an inserted exon that would compromise Pfam domains and 3D structure

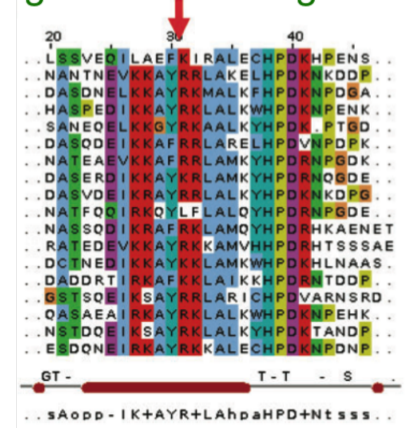
### Homologue showing CCDS insertion



The 3D structure of mouse DNAJ subfamily C2 member 5 (PDB:2CTW), to DNAJC5G-004 has 56% identity with no gaps.

The large red arrow shows that the 16 extra residues in the alternative isoform would insert into an important helix.

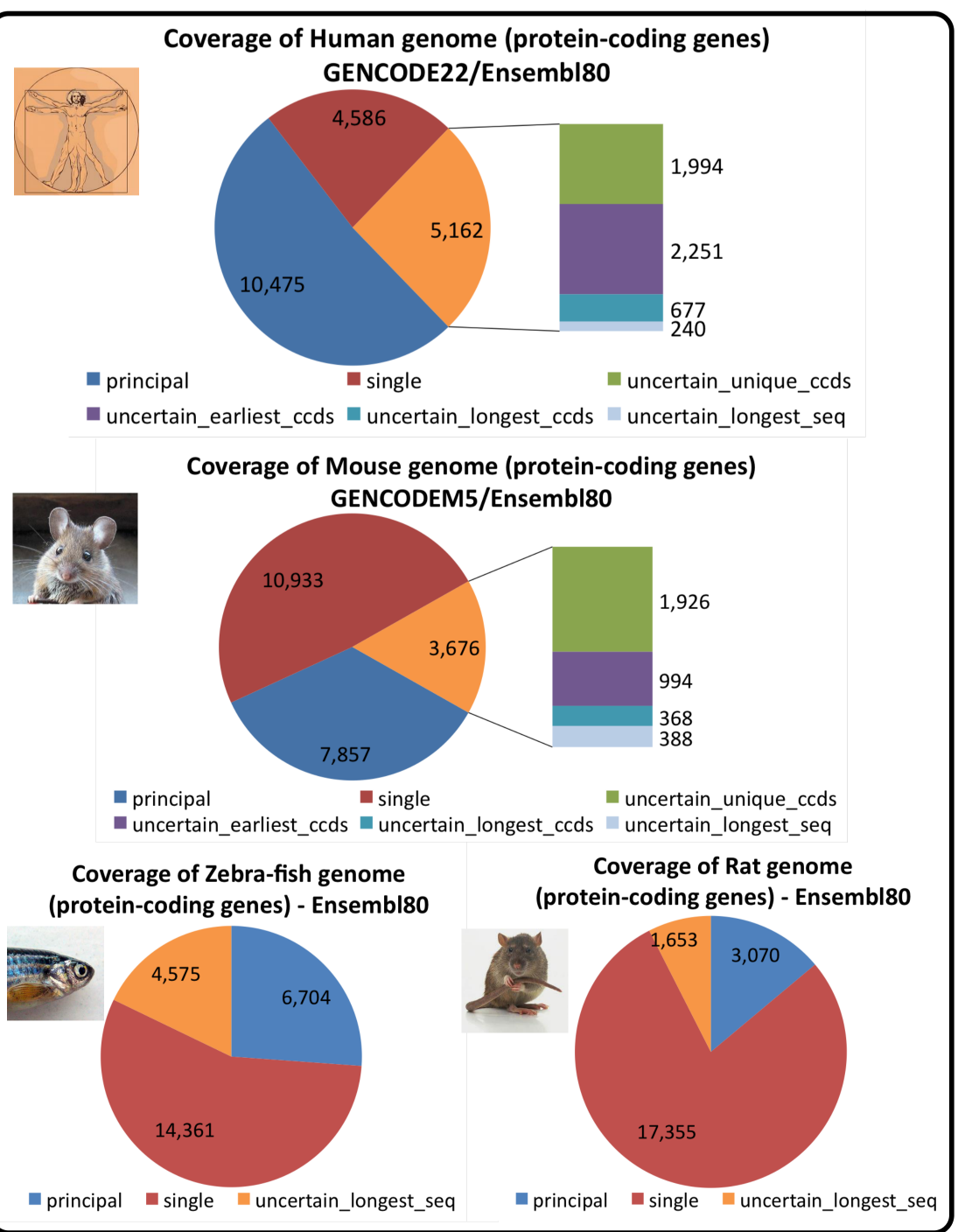
Pfam alignment showing CCDS insertion



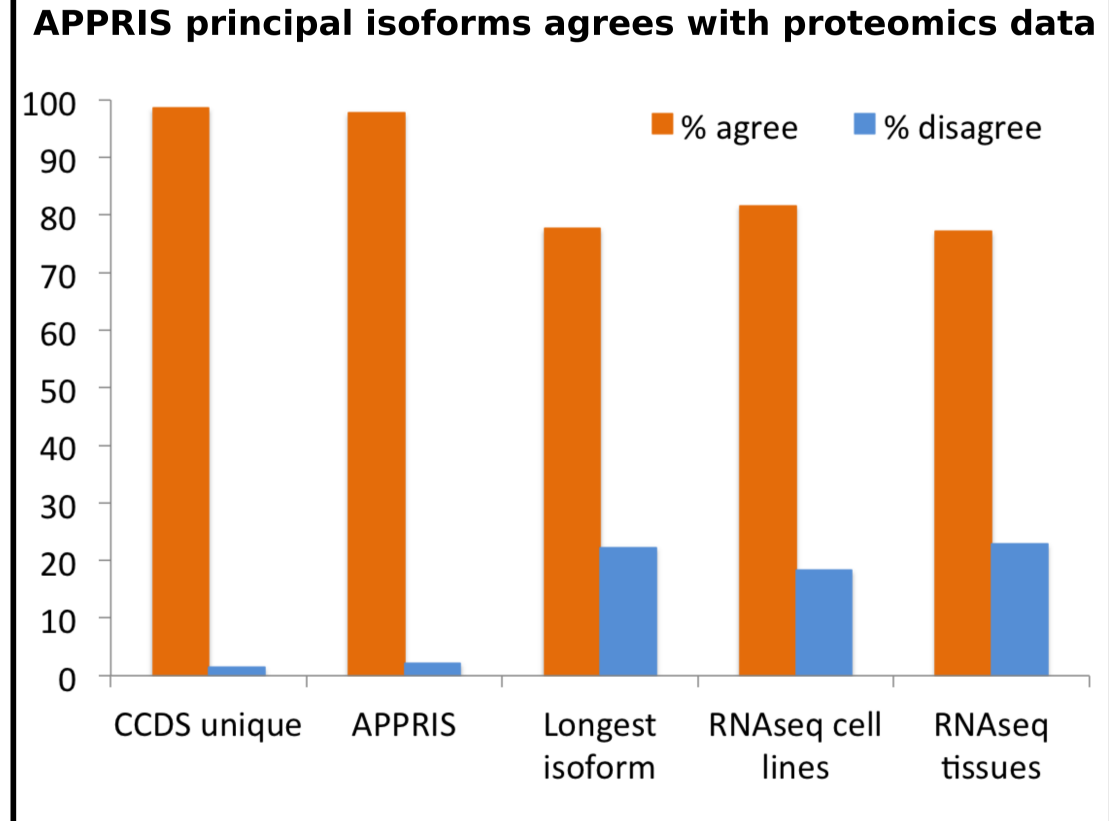
The multiple alignment for a section of the Pfam DNAJ family of sequences.

The red arrow shows that the 16 extra residues in the alternative isoform would insert into a critical region of the functional domain of DNAJC5G.

## **GENOME COVERAGE**



### **VALIDATION**



Here we compared the **main proteomics isoform** from multiple proteomics experiments (6) with (from left to right) the CCDS (4) variants from genes with a unique CCDS variant, APPRIS principal isoforms,

the **longest annotated isoform**, and the dominant transcripts carried out using **RNAseq data** by Gonzalez-Porta et al. (4) on cell lines and tissues.

APPRIS principal isoforms, the main isoforms from proteomics experiments and the unique CCDS isoforms have an exceptionally high level of agreement (99.5% over those genes where each selects a reference isoform).

## CONCLUSIONS

APPRIS principal isoforms have a wide range of uses and are applicable in all fields of research.

Determining a principal isoform is important for research groups **studying individual genes**, and the designation of a single variant as the principal isoform is a critical first step for any genome analysis, for example studies of cancer mutations would be able to use APPRIS data to determine whether the mutations are in principal or alternative exons.

We believe that the principal isoforms identified by APPRIS are a significant advance on the current practice of selecting the longest variants as the reference isoform.

The **APPRIS WebServer** allows for the **annotation** of splice isoforms for individual genes, and provides a range of visual representations and tools to allow researchers to identify the likely effect of splicing events.

The **APPRIS WebServices** have been implemented using REST architecture that permit users to generate annotations automatically in high throughput mode.

At present the APPRIS Database houses annotations for five Ensembl species (human, mouse, rat, pig and zebra fish), the APPRIS WebServer allows users to check Ensembl annotations for six other species, dog, cat, cow, opossum, chicken and fugu.

### REFERENCES

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- lakes Ezkurdia and Paolo Maietta: CNIO, Spain. Adam Frankish, and Jennifer Harrow: Sanger, Cambridge.
Amonida Zadissa, and Fergal Martin: Ensembl, Cambridge.

- Mark Diekhans: UCSC, California. here you are the QR code

If you want this amazing POSTER :-)



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