## **Resource Summary Report**

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# **Alternative Splicing Annotation Project II Database**

RRID:SCR\_000322

Type: Tool

### **Proper Citation**

Alternative Splicing Annotation Project II Database (RRID:SCR\_000322)

#### **Resource Information**

**URL:** <a href="http://www.bioinformatics.ucla.edu/ASAP2">http://www.bioinformatics.ucla.edu/ASAP2</a>

**Proper Citation:** Alternative Splicing Annotation Project II Database (RRID:SCR\_000322)

Description: THIS RESOURCE IS NO LONGER IN SERVICE, documented on 8/12/13. An expanded version of the Alternative Splicing Annotation Project (ASAP) database with a new interface and integration of comparative features using UCSC BLASTZ multiple alignments. It supports 9 vertebrate species, 4 insects, and nematodes, and provides with extensive alternative splicing analysis and their splicing variants. As for human alternative splicing data, newly added EST libraries were classified and included into previous tissue and cancer classification, and lists of tissue and cancer (normal) specific alternatively spliced genes are re-calculated and updated. They have created a novel orthologous exon and intron databases and their splice variants based on multiple alignment among several species. These orthologous exon and intron database can give more comprehensive homologous gene information than protein similarity based method. Furthermore, splice junction and exon identity among species can be valuable resources to elucidate species-specific genes. ASAP II database can be easily integrated with pygr (unpublished, the Python Graph Database Framework for Bioinformatics) and its powerful features such as graph query, multi-genome alignment query and etc. ASAP II can be searched by several different criteria such as gene symbol, gene name and ID (UniGene, GenBank etc.). The web interface provides 7 different kinds of views: (I) user guery, UniGene annotation, orthologous genes and genome browsers; (II) genome alignment; (III) exons and orthologous exons; (IV) introns and orthologous introns; (V) alternative splicing; (IV) isoform and protein sequences; (VII) tissue and cancer vs. normal specificity. ASAP II shows genome alignments of isoforms, exons, and introns in UCSC-like genome browser. All alternative splicing relationships with supporting evidence information, types of alternative splicing patterns, and inclusion rate for skipped exons are listed in separate tables. Users can also search human data for tissueand cancer-specific splice forms at the bottom of the gene summary page. The p-values for tissue-specificity as log-odds (LOD) scores, and highlight the results for LOD >= 3 and at

least 3 EST sequences are all also reported.

Abbreviations: ASAP II

Synonyms: ASAP II Database, Alternative Splicing Annotation Project II

Resource Type: data or information resource, database

**Defining Citation: PMID:17108355** 

**Keywords:** exon, gene structure, genome, alternative splicing, cancer genome alignment, intron, isoform, orthologous exon, orthologous gene, orthologous intron, protein sequence, splice site, tissue, genome alignment, cancer

Funding: NCRR U54 RR021813; NIDCR DE-FC02-02ER63421

Availability: THIS RESOURCE IS NO LONGER IN SERVICE

Resource Name: Alternative Splicing Annotation Project II Database

Resource ID: SCR\_000322

**Alternate IDs:** nif-0000-02572

**Record Creation Time:** 20220129T080200+0000

Record Last Update: 20250507T055851+0000

### Ratings and Alerts

No rating or validation information has been found for Alternative Splicing Annotation Project II Database.

No alerts have been found for Alternative Splicing Annotation Project II Database.

#### **Data and Source Information**

Source: SciCrunch Registry

## **Usage and Citation Metrics**

We found 2 mentions in open access literature.

**Listed below are recent publications.** The full list is available at dkNET.

Katsogiannou M, et al. (2014) The functional landscape of Hsp27 reveals new cellular

processes such as DNA repair and alternative splicing and proposes novel anticancer targets. Molecular & cellular proteomics : MCP, 13(12), 3585.

Lee Y, et al. (2007) ECgene: an alternative splicing database update. Nucleic acids research, 35(Database issue), D99.