.PAR Parameters

Name	Tool (s)	Long	Short	Value	Since	Description
COMPLET E3	asta	-cp3		Boolean	2.2	Require 3' complete transcripts DEPRECATED, by (3.1) refactored to OUTOPTIONS.CP3
GENOME CHR_SEQ	all	- genome chr	-g -c	File	2.2 3.1	Path to a folder with genomic sequences, one per chromosome (.FASTA format). The header tags of each chromosome have to coincide with the <seqname> field (\$1) of the annotation (GTF file).</seqname>
EVENTS	asta	ev	-e	{ASE,ASI,DSP, VST}	3.1	Types of events that are considered: • ASE: external AS events • ASI: internal AS events • DSP: adDitional SPlicing events • VST: Variable SiTe events
EVENTS_ ATR	asta	ea	-a	{CP3,CSS,FLT, IOK,NMD,SEQ}	3.1	CP3: predict 3'-complete CSS: consider only introns with canonical splice sites FLT: output flank type, 'constitutive' or 'alternative' IOK: consider only introns that are acceptable considering basic splicing rules NMD: predict NMD SEQ: output splice site sequences of event flanks
EVENTS_ DIMENSION	asta	ed	-k -d	Integer	0.0	Dimension of the AS events to be extracted, Default is 2 (i.e. 'pairwise events'), and values < 2 stand for 'complete' events <tm></tm>
EVENTS_ FILE	asta	out eo	-0	File stdout	0.0	Keyword 'stdout' for writing results to the standard output stream, or a fully qualified path to the output file. The parameter is optionalif nothing is specified, the output will be written to a file ' <input file=""/> _astalavista.gtf.gz'.
EVENTS_ OPT	asta	ер	-p	{FLT,NMD,CP3}	3.1	Flags to control output options for events: CP3: predict 3'-complete CSS: consider canonical splice sites only FLT: output flank type, i.e., 'constitutive' or 'alternative' site IOK: acceptable introns NMD: predict NMD SEQ: output splice site sequences
EDGE_CO NFIDENCE	asta	ec		Integer	2.2	Level of confidence for edges (i.e., annotated transcription starts/poly-adenylation sites). The default is to trust no annotated edge and to extend overlapping first/last exons of a transcript to their most extreme position: • 0 if 'RefSeq' appears in the source field of the annotation • 1 if 'mRNA' appears in the source field of the annotation • 2 if 'EST' appears in the source field of the annotation • 3 if if none of the above applies All transcript edges of confidence level > edgeConfidence are extended in case the annotation shows another exon with the same adjacent splice site and an earlier/later start /end. DEPRECATED, to be refactored (soon)
FLANK_T YPE	asta	 flankTy pe		Boolean	2.2	Output the type of the event flanks, i.e., 'constitutive' or 'alternative'. DEPRECATED, by (3.1) refactored to OUTOPTIONS.FTY
GENE_ID	scorer	gid	-g	File	3.1	Name and path of the GeneID parameter file with models for splice sites
HELP	all		-h	Boolean	3.0	Help on usage, list of parameters and corresponding descriptions
INPUT	all	in	-i	File	0.0	Input file with reference annotation (.GTF format), MANDATORY
IN_FILE					3.1	
INTRON_ CONFIDE NCE	asta	ic		Integer	2.2	Level of intron confidence, below which introns are trusted without checks. The default is to trust all introns (i.e., ic= 255). Introns are assigned a confidency class: • 0 for 'RefSeq' appears in the source field of the annotation • 1 for 'mRNA' appears in the source field of the annotation • 2 for 'EST' appears in the source field of the annotation All introns in transcripts of confidence level > threshold are discarded. DEPRECATED, to be refactored (soon)

NMD	asta	-nmd			2.2	Check nonsense-mediated decay conditions. DEPRECATED, by (3.1) refactored to OUTOPTIONS.NMD
SITES	scorer	SS	-S	{SSD,SSA,TSS, CLV,SST,SND, AUG,STP}	3.1	Types of sites that are output: SSD: Splice Site Donor SSA: Splice Site Acceptor TSS: Transcription Start Site CLV: Cleavage Site SST: Soft Start (Unconfirmed 5'-end of transcribed sequence) SND: Soft End (Unconfirmed 3'-end of transcribed sequence) AUG: Start Codon TTP: Stop Codon
SITE_FILE	scorer	so	-f	File stdout	3.1	Keyword 'stdout' for writing results to the standard output stream, or a fully qualified path to the output file.
SITE_OPT	asta	sp	-t	{SSS}	3.1	Flags to control optional output for sites: • SSS: Splice Site Score
SEQ_SITE	asta	_ seqsite		Boolean	2.2	Output splice site sequences with events. DEPRECATED, by (3.1) refactored to EVENTS_ATR
TMP_DIR	all	tmp		File	0.0	The temporary directory, reads the environment variable when set to 'null'.
VARIANT_ FILE	scorer	vcf	V	File	3.1	Name and path of a file with the variant information (.VCF format)