Includes feature
Partial functionality
Uses old or insufficient methods
Unique to XPRESSyourself

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Tuned for ribo-seq data Input raw data Input alignment files Input count tables Input count tables Automated reference genome preparation Automated CDS reference modification Pre-processing Alignment to genome Alignment to transcriptome	
Input alignment files Input count tables Automated reference genome preparation Automated CDS reference modification	
Automated CDS reference modification Input count tables Automated reference genome preparation Automated CDS reference modification	
Automated CDS reference modification Input count tables Automated reference genome preparation Automated CDS reference modification	
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Alignment to genome Alignment to genome	
Alignment to transcriptome	
Read counts to variable features	
ion Isoform quantification	
Quantification Isoform quantification Isof	
Quart Translation Efficiency Calculation	
Differential Expression Analysis	
Novel uORF discovery/functional annotation	
Novel uORF discovery/functional annotation CDS and uORF visualization	
General quality control	
Read size distribution	
Pause prediction	
Read size distribution Pause prediction Periodicity	
Metagene profile (all genes)	
rRNA probe summary	
Other customizable analyses Full automation from pre-processing to analysis	