

# Gene Expression/Transcriptomics Resources

- [Arabidopsis Next-Gen Sequence Databases](#) —

<b>Resource Link</b>	<a href="#">Arabidopsis Next-Gen Sequence Databases</a>
<b>Description</b>	Small RNA, MetC and PARE data from Blake Meyers' group.
<b>Maintained by</b>	Blake Meyers

- [small\\_rna](#)
- [arabidopsis](#)
- [gene\\_expression](#)
- [methylation](#)

- [Arabidopsis Single Cell Developmental Atlas Viewer](#) —

<b>Resource Link</b>	<a href="#">Arabidopsis Single Cell Developmental Atlas Viewer</a>
<b>Description</b>	Browser and visualization of single nuclei transcriptome across development.

- [transcriptomics](#)
- [single\\_cell](#)

- [Array Express](#) —

<b>Resource Link</b>	<a href="#">Array Express</a>
<b>Description</b>	Repository for functional genomics data. Access and submit functional genomics data in MIAME compliant format.
<b>Maintained by</b>	EMBL-EBI

- [microarray](#)
- [gene\\_expression](#)
- [data\\_repository](#)
- [data\\_submission](#)

- [ARS-Arabidopsis RNA-seq Database](#) —

<b>Resource Link</b>	<a href="#">Arabidopsis RNA-seq Database (ARS)</a>
<b>Description</b>	ARS is a free, web-accessible, and user-friendly database, contributes to searching, filtering, visualizing, browsing, and downloading the RNA-seq data including over 28,000 publicly available datasets from Arabidopsis.

- [zhaijx@sustech.edu.cn](#) or [zhangh9@mail.sustech.edu.cn](#).

- [rna\\_seq](#)
- [gene](#)
- [gene\\_expression](#)
- [transcriptomics](#)

- [ARVEX-Arabidopsis Root Virtual Expression eXplorer](#) —

<b>Resource Link</b>	<a href="#">Arabidopsis Root Virtual Expression eXplorer</a>
<b>Description</b>	ARVEX is a Shiny app for exploring single cell gene expression of <i>Arabidopsis</i> root

- [Che-Wei Hsu](#)

- [single\\_cell](#)
- [gene\\_expression](#)

- [ATHENA \(Arabidopsis THaliana ExpressioN Atlas\)](#) —

<b>Resource Link</b>	<a href="#">ATHENA</a>
<b>Description</b>	ATHENA is a data exploration tool that provides access to a collection of > 18,000 protein and > 25,000 transcript expression profiles across a set of 30 matching tissues from <i>Arabidopsis thaliana</i> (Col-0). Data represent intensity-based absolute quantifications (iBAQ) for the proteome and transcripts per kilobase million (TPM) for the transcriptome.
<b>Maintained by</b>	Technical University of Munich

- [proteomics](#)
- [gene\\_expression](#)
- [transcriptomics](#)

• Bio-Analytic Resource for Plant Biology (BAR) —

<b>Resource Link</b>	<a href="#">Bio-Analytic Resource for Plant Biology (BAR)</a>
<b>Description</b>	ePlant tool allows for exploration of <i>Arabidopsis</i> data sets from the kilometre to nanometre scales. eFP Browser provides "gene expression anatograms" for depicting where and when a gene is expressed. <i>Arabidopsis</i> Interactions Viewer taps into the BAR's database of more than 100k protein-protein interactions and 2.8M protein-DNA interactions. Tools for exploring other plant data available too!
<b>Maintained by</b>	<a href="#">Provart Laboratory</a> at the University of Toronto

- [general](#)
- [gene\\_expression](#)

• CoNekT —

<b>Resource Link</b>	<a href="#">CoNekT</a>
<b>Description</b>	Platform to query and visualize plant co-expression and co-function networks. Compare expression profiles across sets of homologs.
<b>Maintained by</b>	<a href="#">Proost and Mutwil</a>

- [gene\\_expression](#)
- [gene\\_ontology](#)
- [homology](#)

• Diurnal —

<b>Resource Link</b>	<a href="#">Diurnal</a>
<b>Description</b>	Check if your favorite gene is cycling in different environmental conditions, locate orthologs between <i>Arabidopsis</i> , rice, <i>Brachypodium</i> , and check expression of orthologs in diel conditions (and sometimes circadian) in those species as well.
<b>Maintained by</b>	Todd Mockler

- gene\_expression

- eFP-SEQ Browser —

<b>Resource Link</b>	<a href="#">eFP-Seq Browser</a>
<b>Description</b>	Electronic pictograph browser for visualizing RNA seq data from existing sources or upload your own.
<b>Maintained by</b>	Provart Lab

- rna\_seq
- gene\_expression

- Eukaryotic Promoter Database —

<b>Resource Link</b>	<a href="#">Eukaryotic Promoter Database</a>
<b>Description</b>	Database of experimentally verified promoters from a variety model organisms including Arabidopsis. Genome annotation version is TAIR 10 not Araport 11

**Maintained by** Swiss Institute of Bioinformatics

- promoter
- gene\_expression
- genomics

- GemMaker —

<b>Resource Link</b>	GemMaker
	 <b>GEMmaker</b>
<b>Description</b>	GEMmaker is a <a href="#">Nextflow</a> workflow for large-scale gene expression sample processing, expression-level quantification and Gene Expression Matrix (GEM) construction. Results from GEMmaker are useful for differential gene expression (DGE) and gene co-expression network (GCN) analyses. The GEMmaker workflow currently supports Illumina RNA-seq datasets.
<b>Maintained by</b>	

- gene\_expression
- software
- data\_analysis

- Gene Expression Omnibus (GEO) —

<b>Resource Link</b>	<a href="#">Gene Expression Omnibus (GEO)</a>
	 <b>GEO</b> Gene Expression Omnibus
<b>Description</b>	GEO is a public functional genomics data repository supporting MIAME-compliant data submissions. Array- and sequence-based data are accepted. Tools are provided to help users query and download experiments and curated gene expression profiles.
<b>Maintained by</b>	NCBI

- gene\_expression
- data\_repository
- data\_submission
- microarray

- Integrated System for Motif Activity Response Analysis (ISMARA) —

<b>Resource Link</b>	<a href="#">Integrated System for Motif Activity Response Analysis (ISMARA)</a>
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<b>Description</b>	The Integrated System for Motif Activity Response Analysis is an online tool for automatically inferring regulatory networks from gene expression (RNA-seq/ micro-array) or chromatin state (ChIP-seq) data.
<b>Maintained by</b>	van Nimwegen group of the Biozentrum, University of Basel and Swiss Institute of Bioinformatics.

- Jacobsen Epigenomics Browser —

<b>Resource Link</b>	<a href="#">Jacobsen Epigenomics Browser</a>
<b>Description</b>	DNA methylation, small RNA, ChIP data from Jacobsen lab publications, in searchable browser/
<b>Maintained by</b>	Steve Jacobsen

- epigenomics
- dna\_methylation
- small\_rna
- gene\_expression

- Medicinal Plants Genomics Resource —

<b>Resource Link</b>	<a href="#">Medicinal Plants Genomics Resource</a>
<b>Description</b>	Medicinal plant genome databases, genome browsers and genome annotation for 12 species.
<b>Maintained by</b>	Buell Lab

- gene\_annotation
- genomics
- gene\_expression
- metabolomics
- data\_sets

- MetaOmGraph (MOG) —

<b>Resource Link</b>	<a href="#">MetaOmGraph (MOG)</a>
<b>Description</b>	Java platform for rapid, interactive correlations and visualization of large data sets. Species and data-type agnostic; preformatted data supplied, or format your own.

- Eve Wurtele
- transcriptomics
- data\_visualization
- data\_analysis

- miROOT Browser —

<b>Resource Link</b>	<a href="#">MiROOT Browser</a>
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	<a href="https://doi.org/10.15252/embj.2018100754">https://doi.org/10.15252/embj.2018100754</a>
<b>Description</b>	Browser for examining root specific single cell miRNA expression and loading data from <a href="https://doi.org/10.15252/embj.2018100754">https://doi.org/10.15252/embj.2018100754</a>
<b>Maintained by</b>	Olivier Vionnet

- [mRNA](#)
- [single\\_cell](#)
- [gene\\_expression](#)
- [transcriptomics](#)

• [NCBI Bioinformatics Workshop Lessons](#) —

<b>Resource Link</b>	<a href="#">NCBI Bioinformatics Workshops</a>
<b>Description</b>	Recorded from Wash U/NCBI workshop. Lessons include: <b>NCBI Resources for Animal and Plant Genomics Research</b> , <b>Update on NCBI BLAST and Other Sequence Analysis Tools</b> and <b>NCBI Resources for Gene Expression and Genetic Variation Research</b>
<b>Maintained by</b>	NLM, NCBI

- [education\\_outreach](#)
- [bioinformatics](#)
- [analytics](#)
- [gene\\_expression](#)
- [genomics](#)

• [PlaNet](#) —

<b>Resource Link</b>	<a href="#">PlaNet</a>
<b>Description</b>	Co-expression network analysis across multiple plant species including Arabidopsis

- [interactome](#)
- [data\\_analysis](#)
- [transcriptomics](#)

• [Plant/Eukaryotic and Microbial Systems Resource \(PMR\)](#) —

<b>Resource Link</b>	<a href="#">Plant/Eukaryotic and Microbial Systems Resource (PMR)</a>
<b>Description</b>	Database with web-based visualization and analysis of processed metabolomics data and its metadata. Combined with transcriptomic and/or Mass Spec Imaging (MSI) data/metabolite imaging data from the same samples as the metabolomic data (as available).
<b>Maintained by</b>	Eve Wurtele

- [metabolomics](#)
- [transcriptomics](#)

• [Root Cell Atlas](#) —

<b>Resource Link</b>	<a href="#">Root Cell Atlas</a>
<b>Description</b>	Web service for visualizing single cell transcriptomic data in an AI model of a plant root.

- [single\\_cell](#)
- [transcriptomics](#)
- [gene\\_expression](#)

• [TF2Network](#) —

<b>Resource Link</b>	<a href="#">TF2Network</a>
	
<b>Description</b>	Predict gene regulatory networks in Arabidopsis using published TF binding sites.
<b>Maintained by</b>	Klaas Vandepoele <ul style="list-style-type: none"><li>• transcriptomics</li><li>• interactome</li><li>• data_analysis</li><li>• gene_expression</li></ul>