

The topological shape of gene expression across the evolution of flowering plants

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Interactive “Intro to Python” JupyterBook:

<https://plantsandpython.github.io/PlantsAndPython/>

Introduction

Grand Challenge: Relating genotype to phenotype: development, environment, evolution, . . .

- **Gene expression:** Common currency across scales.
- **Molecular level:** DNA, -omics. Responsible for gene expression complexity.
- **Organism level:** Cell-specific expression, development orchestrated by gene expression.
- **Population level:** Life history, evolution.
- **Ecological level:** Climate, Global distribution of species, etc.

Grand Challenge: Relating genotype to phenotype: evolution, development, environment.

- 900 mil. years of evolution, Over 300K gene expression data sets.
- Typically, 1-specie or 1-gene studies.
- **Goal:** A meta-study of gene expression across all species.
- **Challenge:** Tremendous biological complexity, data set heterogeneity.
- **Approach:** Reduce heterogeneity, use Mapper for visualization.
- **Observations:**
 - Core, conserved backbone defining plant form and function.
 - Patterns differentiating plant tissues, biotic and abiotic stresses.

Data

- Selected 16 plant families, 54 distinct species.
 - Broad phylogenetic diversity within angiosperms.
 - High quality reference genome.
 - Breadth of tissue and stress types.
- Sample expression, metadata: NCBI BioProject, SRA, primary publications.
- Raw RNAseq data processed through common analytical pipeline.
- \approx 3200 samples, 2671 left after processing.
- 8 tissue types.
- 9 biotic and abiotic stresses (+ healthy samples!)

Frequency Plots

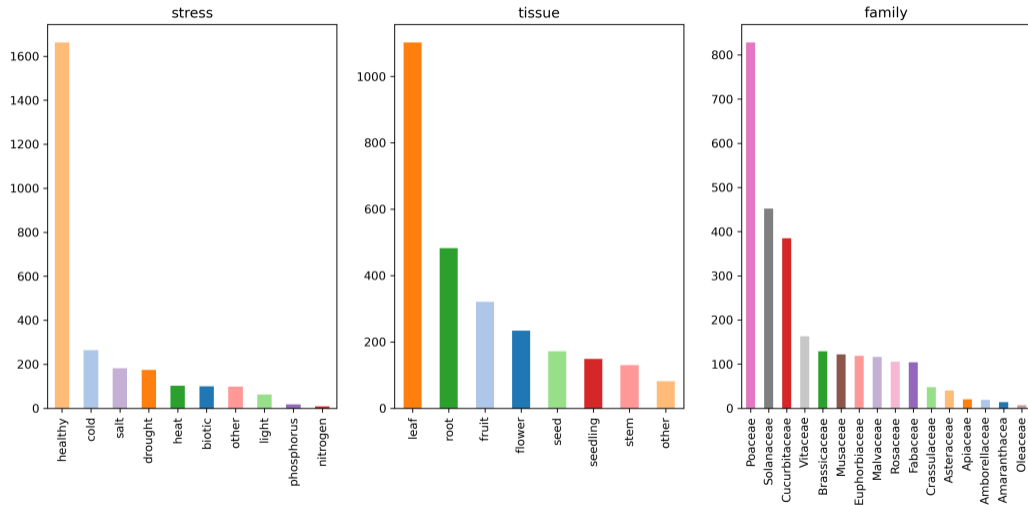


Figure: Factor Frequency Plots

Finding Orthogroups

- Cross-species comparisons: Need correspondences.
- *Orthogroups*: Groups of genes with similar function across species.
- Orthofinder: sequence alignment and clustering
- Excluded multi-gene families with diverse functions.
- Excluded genes with high copy number.
- 2 million genes \rightarrow 6328 orthogroups.
- TPM counts summed for genes in an orthogroup.
- Highly diverse, heterogeneous data combined into single expression matrix.

Dimension Reduction 1

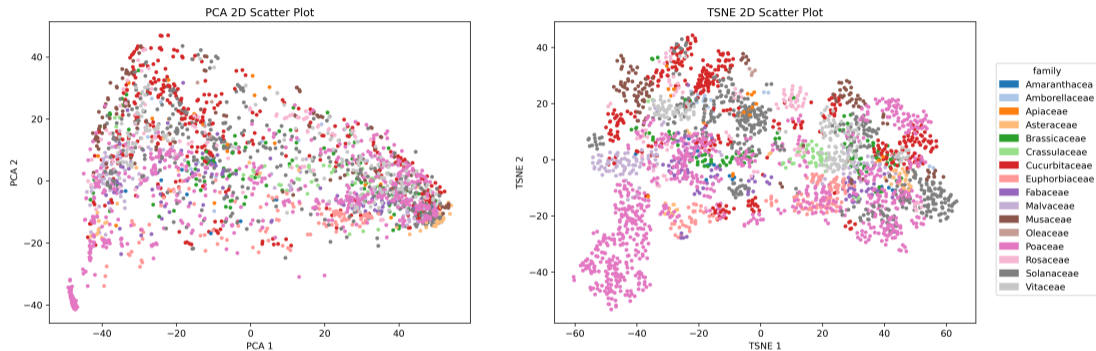


Figure: Dimension Reduction: Points colored by Family

Dimension Reduction 2

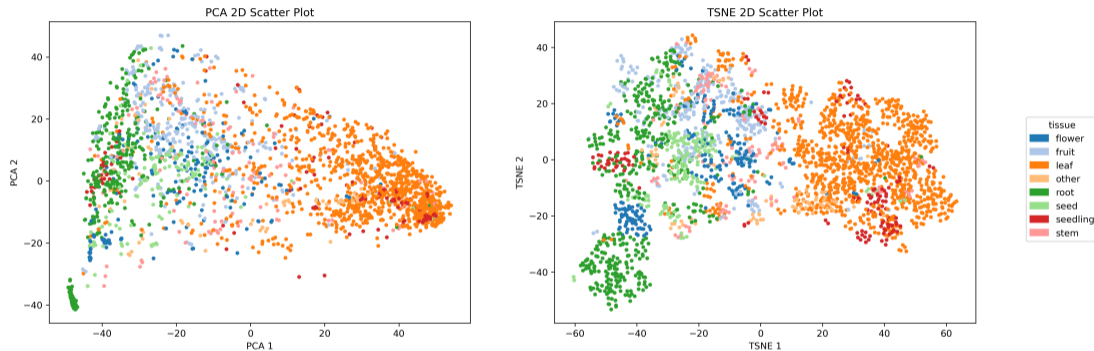


Figure: Dimension Reduction: Points colored by Tissue type

Dimension Reduction 3

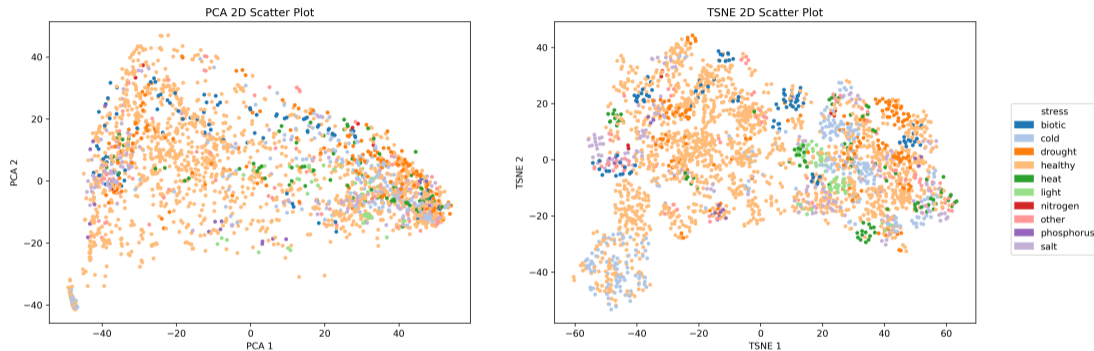


Figure: Dimension Reduction: Points colored by Stress type

Mapper

Mapper Algorithm

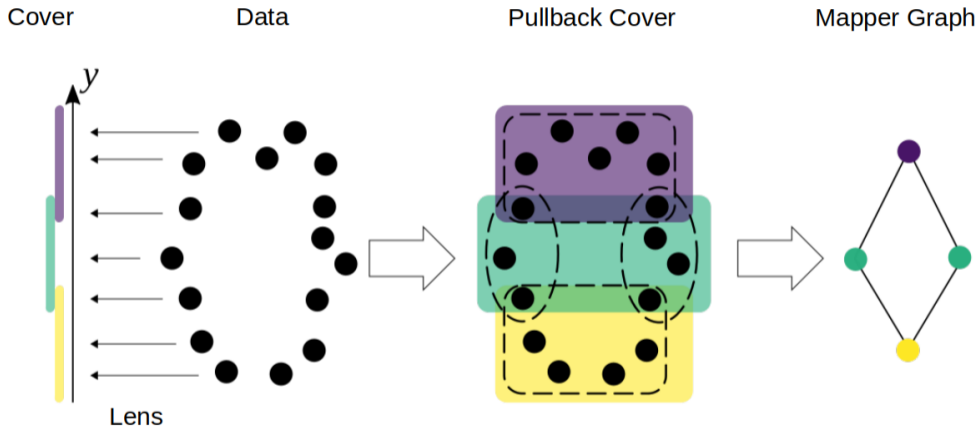


Figure: Mapper Algorithm

Mapper: Key Components

- Choice of lens: Domain / application dependent.
 - Only observe structure visible through specified lens.
 - Induce priors, domain knowledge.
 - We'll focus on creating good lens functions.
- Choice of cover
 - Determines connectivity, density of output graph.
 - Heuristics for optimal cover choice available.
 - Usually: trial and error!
- Clustering algorithm
 - Pick your favorite!
 - We stick to the default: DBSCAN

Creating Lenses

- Two lenses^a: Tissue lens, Stress lens.
- Pick a base class: *healthy vs stressed*, *leaf vs other*.
- Fit a linear model: *ideal* expression for base class.
- Project all samples on to the linear model.
- Residuals: Deviation from *ideal* expression.
- Use norm of the residual as lens.

^aNicolau, Levine, and Carlsson 2011.

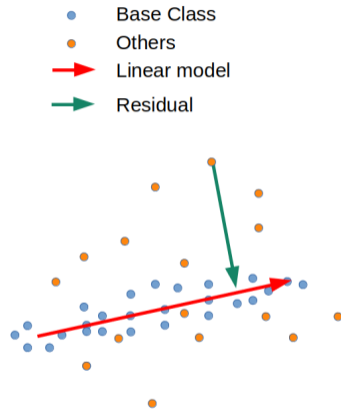


Figure: Creating lens

Lens Correlation Analysis

- For a given lens:
- For each orthogroup:
 - Compute Lens-Orthogroup correlation.
- 6328 correlation values.
- 2.5% most +ve correlations (right tail).
- 2.5% most -ve correlations (left tail).
- 159 Orthogroups in each tail.
- Perform GO Enrichment Analysis for each subset vs all.

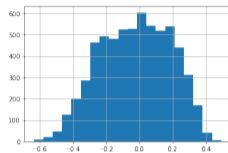


Figure: Leaf lens

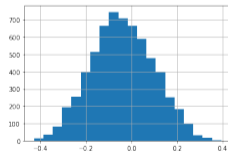


Figure: Stress lens

- Gene Ontology (GO): Standardized vocabulary describing gene function.
 - Cellular component
 - Molecular function
 - Biological Process
- Enrichment: test subset of genes vs all
 - Does the subset contain more representative of certain GO terms compared to chance?
 - Fischer's exact / Hyper-geometric test
- We want to test a subset of orthogroups vs all.
- GO terms only available for genes, not orthogroups!

- Use Arabidopsis genome as reference.
 - Also used to find orthogroups.
 - Why?: Model organism, very well studied genome.
- Use orthogroup - Arabidopsis gene correspondence.
- Create orthogroup - GO term associations.
- GO Analysis tools: <https://pypi.org/project/goatools/>
- Perform enrichment analysis for each tail separately.
- Correct for multiple testing.

Results

Go Enrichment Results

- Tissue lens: Captures photosynthetic vs non-photosynthetic divide.
- GO enrichment of +ve correlated orthogroups:
 - Core metabolic processes, development of non-photosynthetic tissues.
- GO enrichment of -ve correlated orthogroups:
 - Related to photosynthesis, response to light, chloroplast organization.
- Stress lens: healthy vs stressed gene expression
- GO enrichment of +ve correlated orthogroups:
 - Genes involved in stress response.
- GO enrichment of -ve correlated orthogroups:
 - Genes involved in growth and reproduction.

Mapper: Tissue Lens

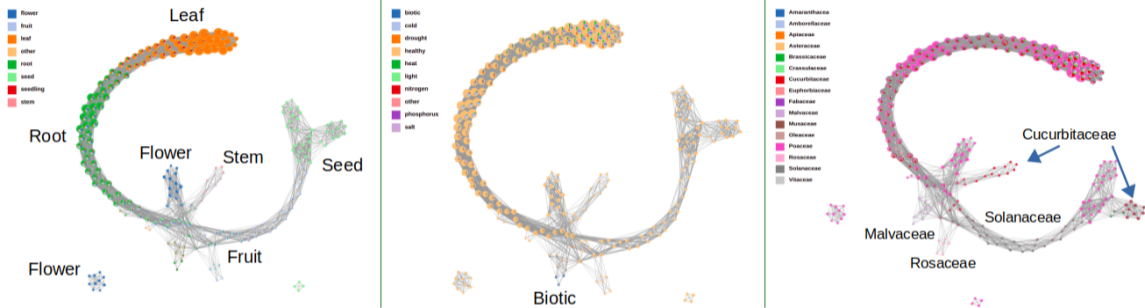


Figure: Tissue (leaf) Mapper Visualization

Mapper: Stress Lens

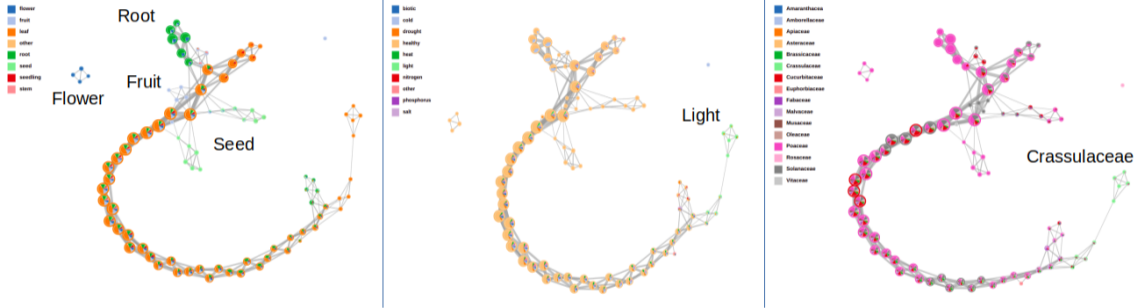


Figure: Stress Mapper Visualization

Discussion

- First cross-species expression study.
- Curated a novel data set.
- Gene expression data observed through phenotype lenses.
- Tissue lens: captures life cycle of plants.
- Stress lens: Deeply conserved stress response signatures.

Future Work

- Proof of concept: Cross-species gene expression can be valuable.
- Expand to include the wealth of public gene expression data sets
- Go beyond mapper. More sophisticated genome alignment and analysis methods.
- Transfer learning across species?

Preprint: <https://www.biorxiv.org/content/10.1101/2022.09.07.506951v1>

Data and Code: <https://github.com/PlantsAndPython/plant-evo-mapper>

email: palandes@msu.edu

Monica Nicolau, Arnold J. Levine, and Gunnar Carlsson. “Topology based data analysis identifies a subgroup of breast cancers with a unique mutational profile and excellent survival”. In: *Proceedings of the National Academy of Sciences* 108.17 (2011), pp. 7265–7270. DOI: [10.1073/pnas.1102826108](https://doi.org/10.1073/pnas.1102826108).

Mapper: Root Lens

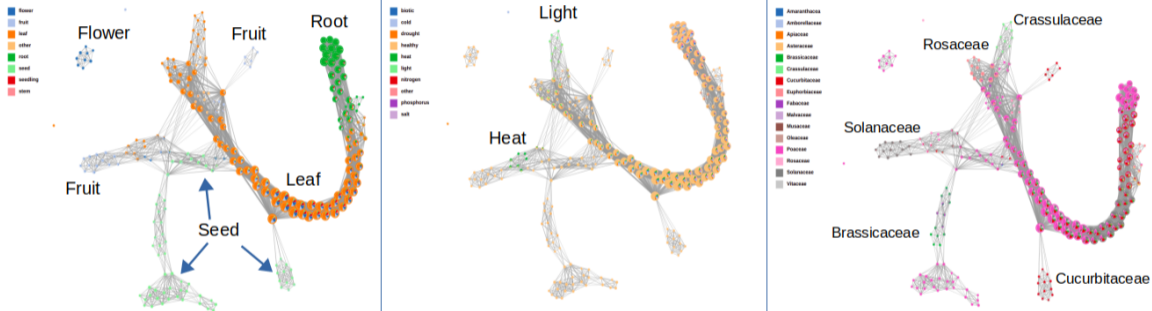


Figure: Root Mapper Visualization