

# Comp683: Computational Biology

## Lecture 22

April 18, 2024

# Today

- Finish up MOFA
- Modifications with MOFA+
- Intro to convex optimization and ADMM, needed to solve our problem.

# Quick Announcements

- Homework 2 is now available online and due **Friday April 26**. → [https://github.com/natalies-teaching/Comp683\\_CompBio\\_2024/tree/main/Homework2](https://github.com/natalies-teaching/Comp683_CompBio_2024/tree/main/Homework2)
- Please sign up for project presentations, [https://docs.google.com/spreadsheets/d/1LvzE54589TLroS8McWfQ6uAscSf9u0Ls\\_FXnLyScMw4/edit?usp=sharing](https://docs.google.com/spreadsheets/d/1LvzE54589TLroS8McWfQ6uAscSf9u0Ls_FXnLyScMw4/edit?usp=sharing).

# Dealing with Latent Variables

More help with factorized approximations...

- Our situation is the case of having an intractable posterior distribution of unobserved variables  $p(\mathbf{X} \mid \mathbf{Y})$ .
- Here,  $\mathbf{X}$  is denoting our un-observed variables, and  $\mathbf{Y}$  is denoting the observed variables

We can deal with  $p(\mathbf{X} \mid \mathbf{Y})$  by approximating it with a factorized form,

$$q(\mathbf{X}) = \prod_i q(\mathbf{X}_i)$$

.

This can make inference more efficient.

## Reminder About Latent Variables Here

Notably,  $\mathbf{Z}$  and  $(\mathbf{S} \circ \hat{\mathbf{W}})$

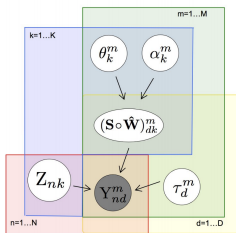


Figure: from Argelaguet *et al.* Molecular Systems Biology. 2018. As always, gray nodes are observed variables and white nodes are the unobserved variables inferred by the model.

## For MOFA : Mean Field Approximation

The assumption under the mean-field approach is that  $q(\mathbf{X})$  factorized over  $M$  disjoint groups of variables as,

$$q(\mathbf{X}) = \prod_{i=1}^M q(\mathbf{x}_i)$$

.  
In the case of MOFA,

$$\begin{aligned} q(\mathbf{Z}, \mathbf{S}, \hat{\mathbf{W}}, \boldsymbol{\alpha}, \boldsymbol{\tau}, \boldsymbol{\theta}) &= q(\mathbf{Z})q(\boldsymbol{\alpha})q(\boldsymbol{\theta})q(\boldsymbol{\tau})q(\mathbf{S}, \hat{\mathbf{W}}) \\ &= \prod_{n=1}^N \prod_{k=1}^K q(z_{n,k}) \prod_{m=1}^M \prod_{k=1}^K q(\alpha_k^m) q(\theta_k^m) \prod_{m=1}^M \prod_{d=1}^{D_m} q(\tau_d^m) \prod_{m=1}^M \prod_{d=1}^{D_m} \prod_{k=1}^K q(\hat{w}_{d,k}^m, s_{d,k}^m) \end{aligned}$$

## Update via EM - $q(\mathbf{Z})$

$$q(\mathbf{Z}) = \prod_{k=1}^K \prod_{n=1}^N q(z_{nk}) = \prod_{k=1}^K \prod_{n=1}^N \mathcal{N}(z_{nk} \mid \mu_{z_{nk}}, \sigma_{z_{nk}})$$

where

$$\sigma_{z_{nk}}^2 = \left( \sum_{m=1}^M \sum_{d=1}^{D_m} \langle \tau_d^m \rangle \langle (s_{dk}^m \hat{w}_{dk}^m)^2 \rangle + 1 \right)^{-1}$$
$$\mu_{z_{nk}} = \sigma_{z_{nk}}^2 \sum_{m=1}^M \sum_{d=1}^{D_m} \langle \tau_d^m \rangle \langle s_{dk}^m \hat{w}_{dk}^m \rangle \left( y_{nd}^m - \sum_{j \neq k} \langle s_{dj}^m \hat{w}_{dj}^m \rangle \langle z_{nj} \rangle \right)$$

# Moving into Results: Summary

After having inferred  $\mathbf{W}^m$ s and  $\mathbf{Z}$ , several things can be done, such as, imputation, annotation of factors, etc.

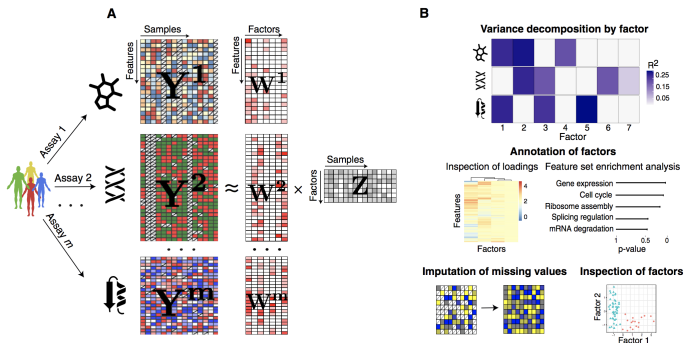


Figure: from Argelaguet *et al.* Molecular Systems Biology. 2018.



# Getting Intuition About Variance Explained by Factors

- **Variance Explained by Factor  $k$  in View  $m$**  : Compute the fraction of variance explained by factor  $k$  in view  $m$  as,

$$R_{m,k}^2 = 1 - \left( \sum_{n,d} y_{nd}^m - z_{nk} w_{kd}^m - \mu_d^m \right)^2 / \left( \sum_{n,d} y_{nd}^m - \mu_d^m \right)^2$$

Here,  $\mu_d$  is the feature-wise mean.

# Variance Explained per Modality

- **Variance Explained In Each Modality Over All Features :**

Compute the fraction of variance explained per view as,

$$R_m^2 = 1 - \left( \sum_{n,d} y_{nd}^m - \sum_k z_{nk} w_{kd}^m - \mu_d^m \right)^2 / \left( \sum_{n,d} y_{nd}^m - \mu_d^m \right)^2$$

Here,  $\mu_d^m$  is the feature-wise mean.

# Results in Dataset 1 : CLL dataset

Find data here

[https://rdrr.io/github/bioFAM/MOFAdata/man/CLL\\_data.html](https://rdrr.io/github/bioFAM/MOFAdata/man/CLL_data.html)

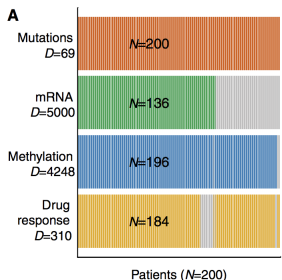


Figure: from Argelaguet *et al.* Molecular Systems Biology. 2018. Modalities and present/missing features for each patient.

# Visualization of Samples

Samples can be projected into two dimensions based on the first two factors inferred in the matrix,  $\mathbf{Z}$

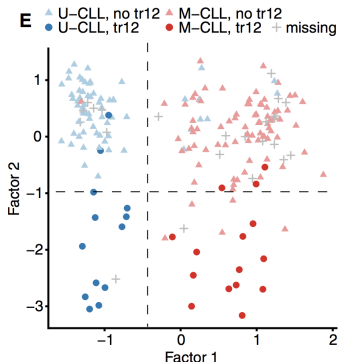


Figure: colors denote something related to the status of the tumor. Shape indicates chromosome 12 trisomy status.

# Jointly Predicting Clinical Outcome in Contrast to a Single Type of Data

Experiment :

- Predict time to treatment for  $N = 174$  patients using multivariate Cox regression trained using the 10 factors from MOFA
- Compare this performance on the prediction accuracy on individual modalities.
  - In this case, reduce each individual modalities to the top 10 PCs.

# Results Predicting Time to Treatment

Note that the Y-axis is simply a statistic reflecting goodness of fit between true and predicted times to treatment (Use top PCs instead of regular feature space).

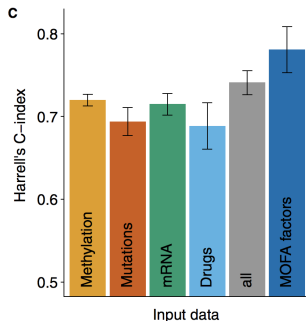


Figure: from Fig. 4 of Argelaguet *et al.* Molecular Systems Biology. 2018.

## Instead of using PC Representations for Each Modality

Instead of using PC representations for each modality, the authors also compared to prediction with all features from each individual modality. Again, predicting time to treatment.

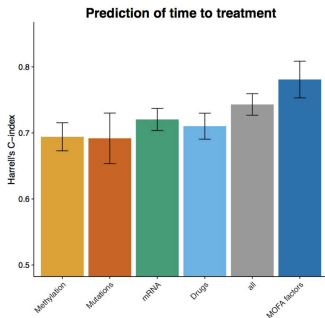


Figure: from Supp. Figure 20. Using all features still does not beat performance based on MOFA factors.

## Certain Combinations of Modalities Can Turn out to be More Effective

Applying MOFA with individual modalities held out reveals sometimes certain combinations of modalities is more effective than just throwing everything in all together!

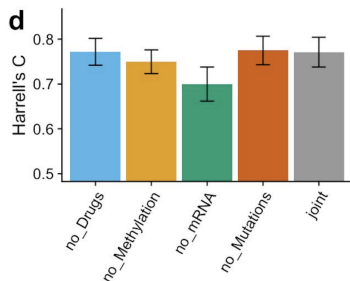


Figure: from Supplementary Figure 8.



# Survival Analysis According to Particular Factors

Survival probability vs time-to-treatment for the individual MOFA factors

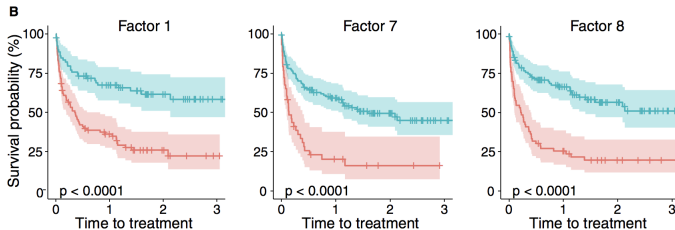


Figure: from Fig. 4. We can see distinct patterns based on survival probability against predicted time to next treatment.

## Dataset 2 : Single-Cell Data for a Differentiation Trajectory

Data from a single-cell multiomics dataset. This is applied to 87 mouse embryonic stem cells, with 16 that were cultured in '2i1' media, giving them a naive pluripotent state. The remaining cells were serum grown and hence committed to a differentiation trajectory.

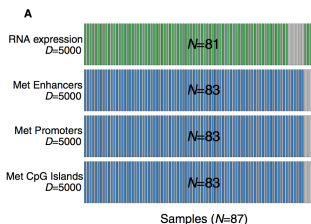


Figure: from Fig. 5. Distribution of samples profiled in each modality.

# Visualizing Cells Based on $\mathbf{Z}$

Using  $\mathbf{Z}$  to plot cells based on the first two factors, the cells separate according to how they were cultured (which also corresponds to differentiation status!)

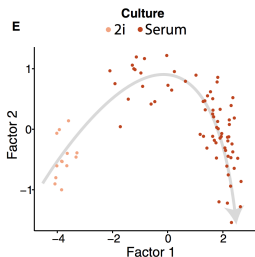


Figure: from Fig. 5.

# Inferred Factors on Genes and How They Relate to Pluripotency and Differentiation

Using the  $\mathbf{W}^m$ s for the mRNA data, the authors compared the ranks of genes involved in pluripotency and differentiation, respectively to the loadings.

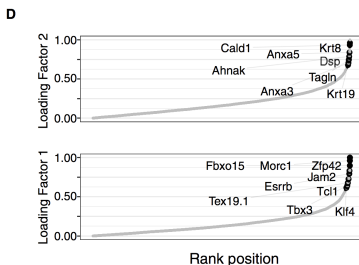


Figure: from Fig. 5. Pluripotency genes (top) vs differentiation related genes (bottom)

# An Extension, MOFA+

A trivial extension....almost the same, except now we learn a  $\mathbf{Z}$  per group.

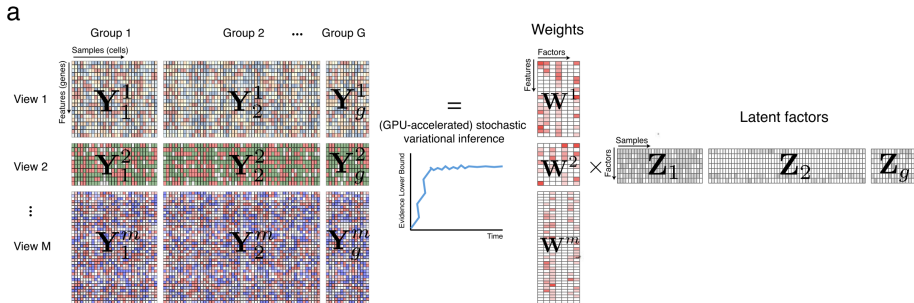


Figure: from Argalaguet *et al.* Genome Biology. 2020. Views are modalities. Groups are some partitioning of the samples.

# Downstream Analysis

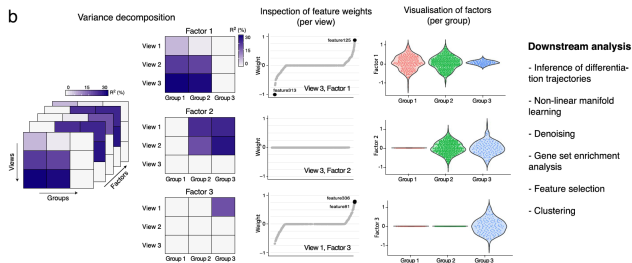


Figure: Variance decomposition by factor, view, and group

So what do we think? Does anyone have any ideas for when this would be useful for your application?

## Defining $Y_{gm}$

Similar to what we have seen with the regular MOFA,

$$Y_{gm} = Z_g W_m^T + \epsilon_{gm}$$

- $Y_{gm}$  is the matrix of observations for the  $m$ th modality and  $g$ th group
- $W_m$  is the weight matrix for the  $m$ th modality
- $Z_g$  is the factor matrix for the  $g$ th group
- $\epsilon_{gm}$  is the residual noise for the  $m$ th modality in the  $g$ th group



# Example

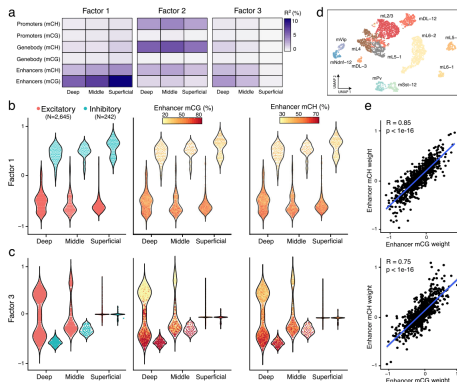


Figure: Context dependent DNA methylation signature associated with cellular diversity in the mammalian cortex. Views are genomic content (DNA methylation signatures), groups are cortical layer.

## Switching Gears for an optimization tool we will need for one more omics study: ADMM

- ADMM stands for alternating direction method of multipliers.
- The basic idea is to break up a big problem into sub-problems.
- The complicated objective here will be solved using multi-block ADMM, an extension of regular ADMM by putting the problem in the following form,

$$\begin{array}{ll} \min_{x_i} & f_1(x_1) + f_2(x_2) + \cdots + f_K(x_K) \\ \text{subject to} & \mathbf{E}_1 x_1 + \mathbf{E}_2 x_2 + \cdots + \mathbf{E}_K x_K = c \end{array}$$

# Quick Intro to Convex Optimization

First regarding **optimization** in general.

- All of us are learning parameters all of the time (even when we do something simple, like regression.....)

The magic here is that if you can formulate your problem as a **convex problem**,

- It can often be solved efficiently and exactly (similar to how we deal with simple problems, like least squares)
- Often a challenge is knowing whether or not an objective function is convex
- Sometimes problems can be re-stated as convex problems to use all of the nice theory.

# Quick Plug for CVX

- CVX is software for convex optimization.
- It is nice because you can easily specify the objective function and constraints
- <https://www.cvxpy.org/index.html>

# An Example for a Classic Least Squares Problem

All of us have seen the following many times before,

$$\min ||Ax - b||_2^2$$

Here  $A$  is an  $M \times N$  matrix, with a corresponding  $N$ -length vector,  $b$  of the response variables.

# Corresponding CVX Code

```
# Import packages.
import cvxpy as cp
import numpy as np

# Generate data.
m = 20
n = 15
np.random.seed(1)
A = np.random.randn(m, n)
b = np.random.randn(m)

# Define and solve the CVXPY problem.
x = cp.Variable(n)
cost = cp.sum_squares(A @ x - b)
prob = cp.Problem(cp.Minimize(cost))
prob.solve()

# Print result.
print("\nThe optimal value is", prob.value)
print("The optimal x is")
print(x.value)
print("The norm of the residual is ", cp.norm(A @ x - b, p=2).value)
```

# Who Cares about Least Squares

You can also solve harder problems, like general linear programs.

$$\begin{array}{ll}\text{minimize} & c^T x \\ \text{subject to} & Ax \leq b\end{array}$$

# Also easy with CVX

```
# Import packages.
import cvxpy as cp
import numpy as np

# Generate a random non-trivial linear program.
m = 15
n = 10
np.random.seed(1)
s0 = np.random.randn(m)
lamb0 = np.maximum(-s0, 0)
s0 = np.maximum(s0, 0)
x0 = np.random.randn(n)
A = np.random.randn(m, n)
b = A @ x0 + s0
c = -A.T @ lamb0

# Define and solve the CVXPY problem.
x = cp.Variable(n)
prob = cp.Problem(cp.Minimize(c.T @ x),
                  [A @ x <= b])
prob.solve()

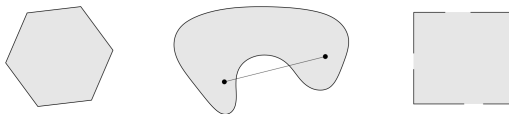
# Print result.
print("\nThe optimal value is", prob.value)
print("A solution x is")
print(x.value)
print("A dual solution is")
print(prob.constraints[0].dual_value)
```



# Convex Sets

A set  $C$  is **convex** if the line segment between any two points in  $C$  lies in  $C$ . That is, for any  $x_1, x_2 \in C$  and for any  $\theta$  with  $0 \leq \theta \leq 1$ , we have,

$$\theta x_1 + (1 - \theta)x_2 \in C$$



**Figure 2.2** Some simple convex and nonconvex sets. *Left.* The hexagon, which includes its boundary (shown darker), is convex. *Middle.* The kidney shaped set is not convex, since the line segment between the two points in the set shown as dots is not contained in the set. *Right.* The square contains some boundary points but not others, and is not convex.

Figure: from the CVX book by Boyd and Vandenberghe

# Convex Function

A function  $f : \mathbb{R}^n \rightarrow \mathbb{R}$  is convex, if the domain of  $f$  is a convex set and if for all  $x$  and  $y$  in the domain of  $f$  with  $0 \leq \theta \leq 1$ , we have

$$f(\theta x + (1 - \theta)y) \leq \theta f(x) + (1 - \theta)f(y)$$



**Figure 3.1** Graph of a convex function. The chord (*i.e.*, line segment) between any two points on the graph lies above the graph.

Figure: from the CVX book by Boyd and Vandenberghe. Geometrically, the inequality means that the line segment between  $(x, f(x))$  and  $(y, f(y))$  lies above  $f$

# Examples of Convex Functions

- $e^{ax}$
- $|x|^p$
- $x \log x$
- $x^a$
- $ax + b$

# Dual Problem

Consider a convex equality constrained optimization problem,

$$\begin{array}{ll}\text{minimize} & f(x) \\ \text{subject to} & Ax = b\end{array}$$

We can write the Lagrangian as,

$$L(x, y) = f(x) + y^T (Ax - b)$$

The idea of Lagrangian duality is to take the constraints into account by augmenting the objective function with a weighted sum of the constraints.

## Dual Problem Continued

Given the Lagrangian,

$$L(x, y) = f(x) + y^T (Ax - b)$$

we denote the dual function as,

$$g(y) = \inf_x L(x, y)$$

.

We can also consider the dual problem,

$$\max g(y)$$

Finally recover an optimal value,  $x^*$  from a dual optimal,  $y^*$ ,

$$x^* = \arg \min_x L(x, y^*)$$

# Dual Ascent

Use a gradient method for the dual problem,

$$y^{k+1} = y^k + \alpha^k \nabla g(y^k)$$

In this case, the gradient is,

$$\nabla g(y^k) = A\tilde{x} - b,$$

where

$$\tilde{x} = \operatorname{argmin}_x L(x, y^k)$$

So, the dual ascent method is,

$$x^{k+1} := \operatorname{argmin}_x L(x, y^k) \quad // x\text{-minimization}$$

$$y^{k+1} := y^k + \alpha^k (Ax^{k+1} - b) \quad // \text{dual update}$$

# Dual Decomposition

Suppose  $f$  is separable :

$$f(x) = f_1(x_1) + \cdots + f_N(x_N), \quad x = (x_1, \dots, x_N)$$

Then  $L$  is separable in  $x$  :

$$L(x, y) = L_1(x_1, y) + \cdots + L_N(x_N, y) - y^T b$$

with

$$L_i(x_i, y) = f_i(x_i) + y^T A_i x_i$$

## Dual Decomposition, Continued

$x$ -minimization in dual ascent splits into  $N$  separate minimizations can be carried out in parallel.

$$x_i^{k+1} := \operatorname{argmin}_{x_i} L_i(x_i, y^k)$$

Updates are,

$$\begin{aligned} x_i^{k+1} &:= \operatorname{argmin}_{x_i} L_i(x_i, y^k), \quad i = 1, \dots, N \\ y^{k+1} &:= y^k + \alpha^k \left( \sum_{i=1}^N A_i x_i^{k+1} - b \right) \end{aligned}$$



# Augmented Lagrangian

For  $\rho > 0$ , form the augmented Lagrangian,

$$L_\rho(x, y) = f(x) + y^T(Ax - b) + (\rho/2)\|Ax - b\|_2^2$$

- The augmented Lagrangian was developed to in part bring robustness to the dual ascent method, and to encourage convergence without strict assumptions about convexity of  $f$ .

Applying dual ascent (also called here method of multipliers),

$$\begin{aligned}x^{k+1} &:= \operatorname{argmin}_x L_\rho(x, y^k) \\ y^{k+1} &:= y^k + \rho(Ax^{k+1} - b)\end{aligned}$$

# Alternating Direction Method of Multipliers

The ADMM problem form (where  $f$  and  $g$  are convex) is,

$$\begin{array}{ll}\text{minimize} & f(x) + g(z) \\ \text{subject to} & Ax + Bz = c\end{array}$$

You can further write down the associated augmented Lagrangian as,

$$L_\rho(x, z, y) = f(x) + g(z) + y^T(Ax + Bz - c) + (\rho/2)\|Ax + Bz - c\|_2^2$$

ADMM will therefore do updates as follows,

$$x^{k+1} := \operatorname{argmin}_x L_\rho(x, z^k, y^k)$$

$$z^{k+1} := \operatorname{argmin}_z L_\rho(x^{k+1}, z, y^k)$$

$$y^{k+1} := y^k + \rho(Ax^{k+1} + Bz^{k+1} - c)$$

$\rho$  is the particular update step size.

# ADNI Data (Multimodal Brain Imaging + Biomarkers + Genetic + Clinical Data)



ABOUT

STUDY DESIGN

DATA & SAMPLES

METHODS & TOOLS

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## DATA TYPES

ADNI researchers collect several types of data from study volunteers throughout their participation in the study, using a standard set of protocols and procedures to eliminate inconsistencies. This information is available for free to authorized investigators through the [LONI Image and Data Archive \(IDA\)](#).

Click each data type below for details.

CLINICAL

GENETIC

MRI IMAGE

PET IMAGE

BIOSPECIMEN

Figure: Access ADNI data

<http://adni.loni.usc.edu/data-samples/data-types/>

# A Joint Model of Cognitive Scores and Diagnosis

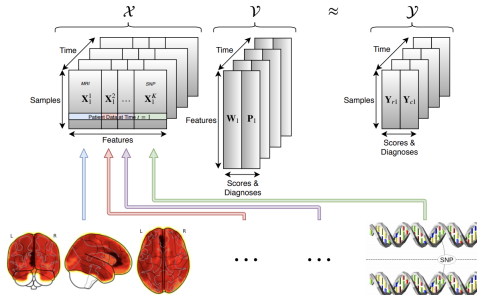


Figure: from Brand, Wang, *et al.* PacSym Biocomputing. 2020.

# Notation and Problem Formulation

- **Input Features:**  $\mathcal{X} = \{\mathbf{X}_1, \mathbf{X}_2, \dots, \mathbf{X}_T\} \in \mathbb{R}^{n \times d \times T}$  This represents patients  $\times$  features  $\times$  timepoints.
- Note that each  $\mathbf{X}_t$  can be broken down across the  $K$  modalities as,  $\{\mathbf{X}_t\}_{j=1}^K$
- The output diagnoses and cognitive scores are represented by another tensor,  $\mathcal{Y} = \{\mathbf{Y}_1, \mathbf{Y}_2, \dots, \mathbf{Y}_T\} \in \mathbb{R}^{n \times c \times T}$
- Each  $\mathbf{Y}_t = [\mathbf{Y}_{rt}, \mathbf{Y}_{ct}]$  represents concatenated response variables for regression ( $r$ ) and classification ( $c$ ).
- The goal is to learn a tensor,  $\mathcal{V} = \{[\mathbf{W}_1, \mathbf{P}_1], [\mathbf{W}_2, \mathbf{P}_2], [\mathbf{W}_T, \mathbf{P}_T]\}$  which represents the coefficients for each feature for regression ( $\mathbf{W}$ s) and classification ( $\mathbf{P}$ s) across the  $T$  timepoints.

# A Regularized Joint Learning Model

$$\min_{\mathcal{W}, \mathcal{P}} \mathcal{L}_r(\mathcal{W}) + \mathcal{L}_c(\mathcal{P}) + \mathcal{R}(\mathcal{V})$$

- Here,  $\mathcal{L}_r(\mathcal{W})$  and  $\mathcal{L}_c(\mathcal{P})$  are the loss functions for the regression and classification tasks, respectively.
- Regression and classification coefficient matrices are  $\mathbf{W}_t \in \mathbb{R}^{d \times c_r}$  and  $\mathbf{P}_t \in \mathbb{R}^{d \times c_c}$ . This yields  $c$  total coefficients.
- $\mathcal{R}(\mathcal{V})$  is a regularization function applied to the matrix unfolded from the tensor,  $\mathcal{V} \rightarrow \mathbf{V}^{d \times cT}$ . Here,  $\mathbf{V}^{d \times cT}$  is constructed by taking the  $(\mathbf{W}_t, \mathbf{P}_t)$  matrix pairs and joining by the columns.

## Regularization, $\mathcal{R}(\mathcal{V})$

To associate image and genomic features to cognitive scores and diagnoses over time, apply an  $\ell_{2,1}$  norm to unfolded coefficient matrix as,

$$\mathbf{V} : \|\mathbf{V}\|_{2,1} = \sum_{d=1}^d \|\mathbf{v}^i\|_2$$

Next, to capture the impact of each modality (e.g. MRI, SNP, etc), the authors use the group  $\ell_1$ -norm ( $G_1$  norm) on the rows of  $\mathbf{V}$  corresponding to modality  $j$  as,

$$\|\mathbf{V}\|_{G_1} = \sum_{j=1}^K \|\mathbf{v}^j\|_2$$

## Regularization, $\mathcal{R}(\mathcal{V})$ , Continued

Finally, to account for inter-modal relationships (or relatedness of features within a modality to cognitive outcome), they use trace norm regularization of  $\mathbf{V}$  as,

$$\mathbf{V} : \|\mathbf{V}\|_* = \sum \sigma_i(\mathbf{V})$$

. Here,  $\sigma_i(\mathbf{V})$  are the singular values of  $\mathbf{V}$



# Objective

Incorporating the three regularizations, the objective can be specified as follows,

$$\min_{\mathbf{V}} J = \sum_{t=1}^T \left[ \|\mathbf{X}_t \mathbf{W}_t - \mathbf{Y}_{rt}\|_F^2 \right] + \sum_{t=1}^T \sum_{i=1}^n \sum_{k=1}^{c_c} \left[ \left( 1 - (\mathbf{x}^{it} \mathbf{p}_{kt} + b_{kt}) y_{ikt} \right)_+ \right] \\ + \gamma_1 \|\mathbf{V}\|_{2,1} + \gamma_2 \|\mathbf{V}\|_{G_1} + \gamma_3 \|\mathbf{V}\|_* ,$$

- The second term is the loss of  $c_c \times T$  one-vs-all multi-class SVM
- $y_{ikt} \in \{-1, 1\}$  is the class label associated with the  $i$ -th patient at time  $t$
- $b_{kt}$  is the bias associated with the  $(k \times t)$ -th SVM
- $(\cdot)_+$  is defined as  $(a)_+ = \max(0, a)$