Data analysis and Unsupervised Learning Dimensionality Reduction: Beyond PCA and Non Linear Methods

MAP 573, 2020 - Julien Chiquet

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https://jchiquet.github.io/MAP573





Outline Introduction

Motivations

Part I

Introduction

Packages required for reproducing the slides

```
library(tidyverse) # opinionated collection of packages for data manipulation
library(GGally) # extension to ggplot vizualization system
library(FactoMineR) # PCA and oter linear method for dimension reduction
library(factoextra) # fancy plotting for FactoMineR output
# color and plots themes
library(RColorBrewer)
pal <- brewer.pal(10, "Set3")
theme_set(theme_bw())</pre>
```

Companion data set: 'scRNA'

Subsamples of normalized Single-Cell RNAseq

Description: subsample of a large data set

Gene-level expression of 100 representative genes for a collection of 301 cells spreaded in 11 cell-lines. Original transcription data are measured by counts obtained by *RNAseq* and normalized to be close to Gaussian.



Pollen, Alex A., et al. Low-coverage single-cell mRNA sequencing reveals cellular heterogeneity and activated signaling pathways in developing cerebral cortex.

Nature biotechnology 32.10 (2014): 1053.

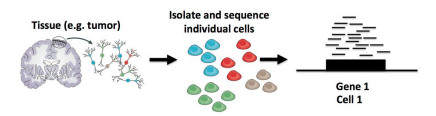


Figure: Single Cell RNA sequnencing data: general principle – source: Stephanie Hicks

Companion data set: 'scRNA'

Brief data summary I

Data manipulation

```
load("../../data/scRNA.RData")
scRNA <- pollen$data %>% t() %>% as_tibble() %>%
   add_column(cell_type = pollen$celltypes)
```

Cell types

```
scRNA %>% dplyr::select(cell_type) %>% summary() %>% knitr::kable()
```

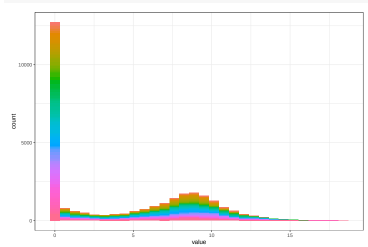
_	11.
	cell_type
	HL60 :54
	K562 :42
	Kera :40
	BJ :37
	GW16 :26
	hiPSC :24
	(Other):78

Companion data set II: 'scRNA'

Brief data summary II

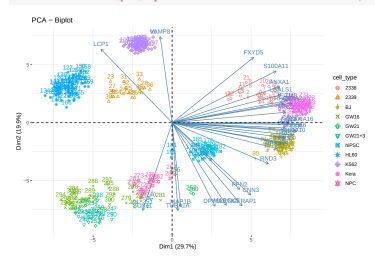
Histogram of normalized expression

```
scRNA %>% dplyr::select(-cell_type) %>% pivot_longer(everything()) %>%
ggplot() + aes(x = value, fill = name) + geom_histogram(show.legend = FALSE)
```



Companion data set: 'scRNA'

```
scRNA %>% FactoMineR::PCA(graph = FALSE, quali.sup = which(colnames(scRNA) == "cell
factoextra::fviz_pca_biplot(select.var = list(contrib = 30), habillage = "cell_t"
```



PCA (and linear methods) limitations

Account for complex pattern

- Linear methods are powerful for planar structures
- May fail at describing manifolds

Preserve local geometry

- High dimensional data are characterized by multiscale properties (local / global structures)
- Non Linear projection helps at preserving local characteristics of distances

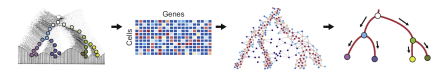


Figure: Intuition of manifolds and geometry underlying sc-data - source: F. Picard

Companion data set II: 'mollusk'

Abundance table (Species counts spread in various sites)

Description: small size count data

Abundance of 32 mollusk species in 163 samples. For each sample, 4 additional covariates are known.



Richardot-Coulet, M., Chessel D. and Bournaud M. Typological value of the benthos of old beds of a large river. Methodological approach. Archiv fur Hydrobiologie, 107.

```
library(PLNmodels); data(mollusk)
mollusk <-
    prepare_data(mollusk$Abundance, mollusk$Covariate[c("season", "site")]) %>%
    as_tibble() %>%
    distinct() # remove duplicates
```

External Covariates

mollusk %>% dplyr::select(site, season) %>% summary() %>% t() %>% knitr::kable()

site	Negria1 :24	Negria2 :24	Pecheurs1:24	Pecheurs2:24	GGravier3:22	GGravie
season	automn:41	spring:44	summer:44	winter:34	NA	NA

Companion data set: 'mollusk'

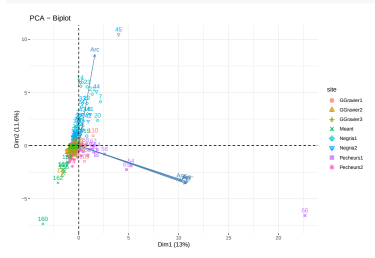
Brief data summary II

Histogram of raw counts

```
mollusk %>% dplyr::select(-site, -season) %>%
  pivot_longer(everything()) %>%
  ggplot() + aes(x = value, fill = name) + geom_histogram(show.legend = FALSE)
## Error: Aesthetics must be either length 1 or the same as the data (316): x
```

Companion data set: 'mollusk'

```
mollusk %>% PCA(graph = FALSE, quali.sup = which(map_lgl(mollusk, is.factor))) %>%
fviz_pca_biplot(select.var = list(contrib = 5), habillage = "site")
```



PCA (and linear methods) limitations

Account for complex data distribution

- Linear methods /PCA are tied to an hidden Gaussian assumption
- Fail with Count data
- Fail with Skew data

Possible solutions

- Probabilistic (non Gaussian) models
- Need transformed (non-linear) input space

Dimension reduction: revisiting the problem setup

Settings

- Training data : $\mathcal{D} = \{\mathbf{x}_1, \dots, \mathbf{x}_n\} \in \mathbb{R}^p$, (i.i.d.)
- Space \mathbb{R}^p of possibly high dimension $(n \ll p)$

Dimension Reduction Map

Construct a map Φ from the space \mathbb{R}^p into a space \mathbb{R}^q of smaller dimension:

$$\Phi: \quad \mathbb{R}^p \to \mathbb{R}^q, q \ll p$$
$$\mathbf{x} \mapsto \Phi(\mathbf{x})$$

How should we design/construct Φ ?

Criterion

- Geometrical approach (see slides on PCA)
- Reconstruction error
- Relationship preservation

Form of the map Φ

- Linear or non-linear ?
- tradeoff between interpretability and versatility ?
- tradeoff between high or low computational resource

Part II

Non-linear methods

Non-linear methods

Motivated by reconstruction error

General goal

Kernel-PCA

Non-negative matrix factorization

Auto-Encoder

2 Relation preservation

Non-linear methods

 Motivated by reconstruction error General goal

> Kernel-PCA Non-negative matrix factorization Auto-Encoder

2 Relation preservation

Reconstruction error approach

1 Construct a map Φ from the space \mathbb{R}^p into a space \mathbb{R}^q of smaller dimension:

$$\Phi: \quad \mathbb{R}^p \to \mathbb{R}^q, q \ll p$$
$$\mathbf{x} \mapsto \Phi(\mathbf{x})$$

- 2 Construct $\widetilde{\Phi}$ from \mathbb{R}^q to \mathbb{R}^p (reconstruction formula)
- 3 Control an error ϵ between ${\bf x}$ and its reconstruction $\hat{{\bf x}}=\tilde{\Phi}(\Phi({\bf x}))$

For instance, the error measured with th Frobenius between the original data matrix \mathbf{X} and its approximation:

$$\epsilon(\mathbf{X}, \hat{\mathbf{X}}) = \left\| \mathbf{X} - \hat{\mathbf{X}} \right\|_F^2 = \sum_{i=1}^n \left\| \mathbf{x}_i - \tilde{\Phi}(\Phi(\mathbf{x}_i)) \right\|^2$$

Reconstruction error approach

1 Construct a map Φ from the space \mathbb{R}^p into a space \mathbb{R}^q of smaller dimension:

$$\Phi: \quad \mathbb{R}^p \to \mathbb{R}^q, q \ll p$$
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Reinterpretation of PCA

PCA model

Let ${f V}$ be a $p \times q$ matrix whose columns are of q orthonormal vectors.

$$\begin{split} \Phi(\mathbf{x}) &= \mathbf{V}^{\top}(\mathbf{x} - \boldsymbol{\mu}) = \tilde{\mathbf{x}} \\ \mathbf{x} &\simeq \tilde{\Phi}(\tilde{\mathbf{x}}) = \boldsymbol{\mu} + \mathbf{V}\tilde{\mathbf{x}} \end{split}$$

→ Model with Linear assumption + ortho-normality constraints

PCA reconstruction error

$$\min_{oldsymbol{\mu} \in \mathbb{R}^p, \mathbf{V} \in \mathcal{O}_{p,q}} \sum_{i=1}^n \left\| (\mathbf{x}_i - oldsymbol{\mu}) + \mathbf{V}^ op \mathbf{V} (\mathbf{x}_i - oldsymbol{\mu})
ight\|^2$$

Solution (explicit)

- ullet $\mu=ar{\mathbf{x}}$ the empirical mean
- V an orthonormal basis of the space spanned by the q first eigenvectors of the empirical covariance matrix

Reinterpretation of PCA

PCA model

Let V be a $p \times q$ matrix whose columns are of q orthonormal vectors.

$$\begin{split} \Phi(\mathbf{x}) &= \mathbf{V}^{\top}(\mathbf{x} - \boldsymbol{\mu}) = \tilde{\mathbf{x}} \\ \mathbf{x} &\simeq \tilde{\Phi}(\tilde{\mathbf{x}}) = \boldsymbol{\mu} + \mathbf{V}\tilde{\mathbf{x}} \end{split}$$

→ Model with Linear assumption + ortho-normality constraints

PCA reconstruction error

$$\underset{\boldsymbol{\mu} \in \mathbb{R}^p, \mathbf{V} \in \mathcal{O}_{p,q}}{\operatorname{minimize}} \sum_{i=1}^n \left\| (\mathbf{x}_i - \boldsymbol{\mu}) + \mathbf{V}^\top \mathbf{V} (\mathbf{x}_i - \boldsymbol{\mu}) \right\|^2$$

Solution (explicit)

- ullet $\mu=ar{\mathbf{x}}$ the empirical mean
- V an orthonormal basis of the space spanned by the q first eigenvectors of the empirical covariance matrix

Important digression: SVD

Singular Value Decomposition (SVD)

The SVD of ${\bf M}$ a $n \times p$ matrix is the factorization given by

$$\mathbf{M} = \mathbf{U}\mathbf{D}\mathbf{V}^{\mathsf{T}},$$

where $r = \min(n, p)$ and

- $\mathbf{D}_{r \times r} = \operatorname{diag}(\delta_1, ... \delta_r)$ is the diagonal matrix of singular values.
- ullet U is orthonormal, whose columns are eigen vectors of $(\mathbf{M}\mathbf{M}^T)$
- V is orthonormal whose columns are eigen vectors of $(\mathbf{M}^T\mathbf{M})$
- ightharpoonup Time complexity in $\mathcal{O}(npqr)$ (less when $k \ll r$ components are required)

Connection with eigen decomposition of the covariance matrix

$$\begin{split} \mathbf{M}^{\top}\mathbf{M} &= \mathbf{V}\mathbf{D}\mathbf{U}^{\top}\mathbf{U}\mathbf{D}\mathbf{V}^{\top} \\ &= \mathbf{V}\mathbf{D}^{2}\mathbf{V}^{\top} = \mathbf{V}\boldsymbol{\Lambda}\mathbf{V}^{\top} \end{split}$$

Important digression: SVD

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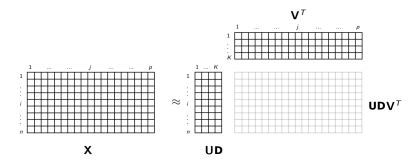
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- U is orthonormal, whose columns are eigen vectors of $(\mathbf{M}\mathbf{M}^T)$
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- \leadsto Time complexity in $\mathcal{O}(npqr)$ (less when $k \ll r$ components are required)

Connection with eigen decomposition of the covariance matrix

$$\begin{split} \mathbf{M}^{\top}\mathbf{M} &= \mathbf{V}\mathbf{D}\mathbf{U}^{\top}\mathbf{U}\mathbf{D}\mathbf{V}^{\top} \\ &= \mathbf{V}\mathbf{D}^{2}\mathbf{V}^{\top} = \mathbf{V}\boldsymbol{\Lambda}\mathbf{V}^{\top} \end{split}$$

PCA solution is given by SVD of the centered data matrix



Since $\tilde{\mathbf{X}} = \mathbf{X}^c \mathbf{V} = \mathbf{U} \mathbf{D} \mathbf{V}^{\top} \mathbf{V} = \mathbf{U} \mathbf{D}$, PCA can be rephrased as

$$\hat{\mathbf{X}}^c = \mathbf{F} \mathbf{V}^\top = \operatorname*{arg\ min}_{\mathbf{F} \in \mathcal{M}_{n,q}, \mathbf{V} \in \mathcal{O}_{p,q}} \left\| \mathbf{X}^c - \mathbf{F} \mathbf{V}^\top \right\|_F^2 \text{ with } \|\mathbf{A}\|_F^2 = \sum_{ij} a_{ij}^2,$$

 $ilde{\mathbf{X}} \in \mathbb{R}^{n imes q}, \mathbf{V} \in \mathbb{R}^{p imes q} \Big\}$ Best linear low-rank representation of \mathbf{X}

Non-linear methods

Motivated by reconstruction error

General goal

Kernel-PCA

Non-negative matrix factorization Auto-Encoder

2 Relation preservation

Non-linear methods

Motivated by reconstruction error

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NMF Model

Example on 'mollusk' I

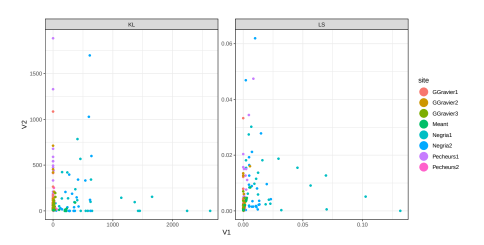
Run the fit

```
nmf_KL <- mollusk %>% select(-site, -season) %>%
nmf(rank = 2, method = 'brunet') %>% basis() %>%
as.data.frame() %>% add_column(algo = "KL") %>% add_column(site = mollusk$site)
nmf_LS <- mollusk %>% select(-site, -season) %>%
nmf(rank = 2, method = 'lee') %>% basis() %>%
as.data.frame() %>% add_column(algo = "LS") %>% add_column(site = mollusk$site)
```

Compare algorithms

```
rbind(nmf_KL, nmf_LS) %>%
  ggplot(aes(x = V1, y = V2, color = site)) +
    geom_point(size=1.25) +
    guides(colour = guide_legend(override.aes = list(size=6))) +
  facet_wrap(.~algo, scales = 'free')
```

Example on 'mollusk' II



Non-linear methods

1 Motivated by reconstruction error General goal Kernel-PCA

Non-negative matrix factorization

Auto-Encoder

2 Relation preservation

Non-linear methods

- Motivated by reconstruction error
- 2 Relation preservation

General goal

MDS

t-SNE

UMAP

Non-linear methods

- Motivated by reconstruction error
- Relation preservation General goal MDS t-SNE UMAP

Pairwise Relation

Focus on pairwise relation $\mathcal{R}(\mathbf{x}_i, \mathbf{x}_{i'})$.

Distance Preservation

• Construct a map Φ from the space \mathbb{R}^d into a space $\mathbb{R}^{d'}$ of smaller dimension:

$$\begin{split} \Phi: \quad \mathbb{R}^d &\to \mathbb{R}^{d'}, d' \ll d \\ \mathbf{x} &\mapsto \Phi(\mathbf{x}) \end{split}$$
 such that
$$\quad \mathcal{R}(\mathbf{x}_i, \mathbf{x}_{i'}) \sim \mathcal{R}'(\mathbf{x}_i', \mathbf{x}_{i'}')$$

Multidimensional scaling

Try to preserve inner product related to the distance (e.g. Euclidean)

t-SNE – Stochastic Neighborhood Embedding

Try to preserve relations with close neighbors with Gaussian kernel

Non-linear methods

- Motivated by reconstruction error
- 2 Relation preservation

General goal

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t-SNE

Non-linear methods

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MDS

 $\mathsf{t}\text{-}\mathsf{SNE}$

UMAP

Stochastic Neighbor Embedding [van der Maaten and Hinton, 2008]

- (x_1,\ldots,x_n) are the points in the high dimensional space \mathbb{R}^p ,
- Consider a similarity between points:

$$p_{i|j} = \frac{\exp(-\|x_i - x_j\|^2 / 2\sigma_i^2)}{\sum_{k \neq i} \exp(-\|x_k - x_j\|^2 / 2\sigma_k^2)}, \ p_{ij} = (p_{i|j} + p_{j|i}) / 2N$$

- ullet σ smooths the data (linked to the regularity of the target manifold)
- ullet σ is chosen such that the entropy of p is fixed to a given value of the so-called perplexity

$$\exp\left(-\sum_{ij}p_{ij}\log(p_{ij})\right)$$

The perplexity parameter

- σ_i Should adjust to local densities (neighborhood of point i)
- Define the Shannon entropy of $p_i = (p_{1|i}, \dots, p_{n|i})$

$$H(p_i) = -\sum_{j=1}^{n} p_{j|i} \log_2 p_{j|i}$$

• The perplexity is defined by:

$$Perp(p_i) = 2^{H(p_i)}$$

- Interpreted as the smoothed effective number of neighbors.
- ullet SNE performs a binary search for the value of si that produces a p_i with a fixed perplexity that is specified by the user.

tSNE and Student / Cauchy kernels

- Consider (y_1, \ldots, y_n) are points in the low dimensional space \mathbb{R}^2
- Consider a similarity between points in the new representation:

$$q_{i|j} = \frac{\exp(-\|y_i - y_j\|^2)}{\sum_{k \neq i} \exp(-\|y_k - y_j\|^2)}$$

Robustify this kernel by using Student(1) kernels (ie Cauchy)

$$q_{i|j} = \frac{(1 + ||y_i - y_j||^2)^{-1}}{\sum_{k \neq i} (1 + ||y_i - y_k||^2)^{-1}}$$

Optimizing tSNE

 Minimize the KL between p and q so that the data representation minimizes:

$$C(y) = \sum_{ij} KL(p_{ij}, q_{ij})$$

The cost function is not convex

$$\left[\frac{\partial C(y)}{\partial y}\right]_i = \sum_j (p_{ij} - q_{ij})(y_i - y_j)$$

- Interpreted as the resultant force created by a set of springs between the map point y_i and all other map points $(y_j)_j$. All springs exert a force along the direction $(y_i y_j)$.
- $(p_{ij}-q_{ij})$ is viewed as a stiffness of the force exerted by the spring between y_i and y_j .

Customed Gradient descent

- Gradient descent initialized by sampling map points randomly from an isotropic Gaussian with small variance centered around the origin
- Gradient update using

$$y^{(t)} = y^{(t-1)} + \eta \frac{\partial C(y)}{\partial y} + \alpha(t)(y^{(t-1)} - y^{(t-2)})$$

- η learning rate, $\alpha(t)$ momentum at iteration t.
- Gaussian noise is added to the map points to perform simulated annealing.

Properties of t-SNE

- good at preserving local distances (intra-cluster variance)
- not so good for global representation (inter-cluster variance)
- hence good at creating clusters of points that are close, but bad at positionning clusters wrt each other
- preprocessing very important : initialize with PCA and feature selection plus log transform (non linear transform)
- ullet percent of explained variance ? interpretation of the q distribution ?

Example on scRNA I

Run the fit

```
scRNA_expr <- scRNA %>% select(-cell_type) %>% as.matrix()
tSNE_perp2 <- Rtsne(scRNA_expr, perplexity = 2)$Y %>%
  as.data.frame() %>% add_column(perplexity = 2) %>% add_column(cell_type = scRNA$
## Error in Rtsne(scRNA_expr, perplexity = 2): could not find function "Rtsne"
tSNE_perp10 <- Rtsne(scRNA_expr, perplexity = 10)$Y %>%
  as.data.frame() %% add_column(perplexity = 10) %>% add_column(cell_type = scRNAs
## Error in Rtsne(scRNA_expr, perplexity = 10): could not find function
"Rtsne"
tSNE_perp100 <- Rtsne(scRNA_expr, perplexity = 100)$Y %>%
  as.data.frame() %% add_column(perplexity = 100) %>% add_column(cell_type = scRN)
## Error in Rtsne(scRNA_expr, perplexity = 100): could not find function
"Rtsne"
```

Compare perplexity

Example on scRNA II

```
rbind(tSNE_perp2,tSNE_perp10,tSNE_perp100) %>%
    ggplot(aes(x = V1, y = V2, color = cell_type)) +
        geom_point(size=1.25) +
        guides(colour = guide_legend(override.aes = list(size=6))) +
    facet_wrap(.~perplexity, scales = 'free')

## Error in rbind(tSNE_perp2, tSNE_perp10, tSNE_perp100): object 'tSNE_perp2'
not found
```

Example on 'mollusk' I

Run the fit

```
mollusk_ab <- mollusk %>% select(-site, -season) %>% as.matrix()
tSNE_perp2 <- Rtsne(mollusk_ab, perplexity = 2)$Y %>%
  as.data.frame() %% add_column(perplexity = 2) %>% add_column(site = mollusk_ab$;
## Error in Rtsne(mollusk_ab, perplexity = 2): could not find function "Rtsne"
tSNE_perp10 <- Rtsne(log(1 + mollusk_ab), perplexity = 10)$Y %>%
  as.data.frame() %% add_column(perplexity = 10) %>% add_column(site = mollusk_abs
## Error in Rtsne(log(1 + mollusk_ab), perplexity = 10): could not find
function "Rtsne"
tSNE_perp50 <- Rtsne(log(1 + mollusk_ab), perplexity = 50)$Y %>%
  as.data.frame() %% add_column(perplexity = 50) %>% add_column(site = mollusk_abs
## Error in Rtsne(log(1 + mollusk_ab), perplexity = 50): could not find
function "Rtsne"
```

Compare perplexity

Example on 'mollusk' II

```
rbind(tSNE_perp2,tSNE_perp10,tSNE_perp50) %>%
   ggplot(aes(x = V1, y = V2, color = site)) +
        geom_point(size=1.25) +
        guides(colour = guide_legend(override.aes = list(size=6))) +
   facet_wrap(.~perplexity, scales = 'free')

## Error in rbind(tSNE_perp2, tSNE_perp10, tSNE_perp50): object 'tSNE_perp2'
not found
```

Outline

Non-linear methods

- Motivated by reconstruction error
- 2 Relation preservation

 General goal

MDS

t-SNE

UMAP

Error in library(umap): there is no package called 'umap'

Uniform Manifold Approximation and Projection [McInnes et al., 2018]

Properties of UMAP

Example on scRNA I

Run the fit

```
scRNA_expr <- scRNA %>% select(-cell_type) %>% as.matrix()
umap_fit <- umap(scRNA_expr)$layout %>%
  as.data.frame() %>% add_column(cell_type = scRNA$cell_type)
## Error in umap(scRNA_expr): could not find function "umap"
```

Visualization

```
umap_fit %>%
ggplot(aes(x = V1, y = V2, color = cell_type)) +
    geom_point(size=1.25) +
    guides(colour = guide_legend(override.aes = list(size=6)))
## Error in eval(lhs, parent, parent): object 'umap_fit' not found
```

Example on 'mollusk' I

Run the fit

```
duplicated <- duplicated(mollusk %>% select(-site, -season))
mollusk_ab <- mollusk %>% select(-site, -season) %>% filter(!duplicated) %>% as.ma
umap_fit <- umap(mollusk_ab)$layout %>%
   as.data.frame() %>% add_column(site = mollusk$site[!duplicated])
## Error in umap(mollusk_ab): could not find function "umap"
```

Visualization

```
umap_fit %>%
  ggplot(aes(x = V1, y = V2, color = site)) +
    geom_point(size=1.25) +
    guides(colour = guide_legend(override.aes = list(size=6)))
## Error in eval(lhs, parent, parent): object 'umap_fit' not found
```

References I



McInnes, L., Healy, J., and Melville, J. (2018).

Umap: Uniform manifold approximation and projection for dimension reduction.

arXiv preprint arXiv:1802.03426.



van der Maaten, L. and Hinton, G. (2008).

Visualizing Data using t-SNE.

Journal of Machine Learning Research, 9(Nov):2579–2605.