

# Fundamentals of Computer Graphics

Assignment problems

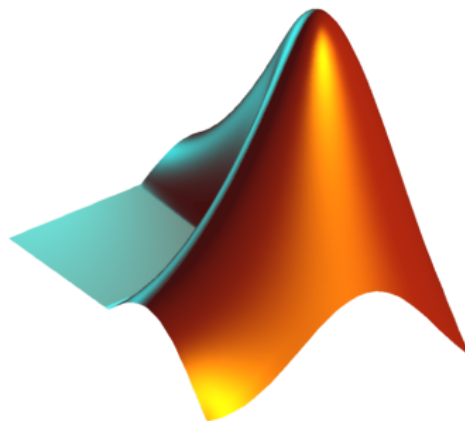
Emanuele Rodolà  
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SAPIENZA  
UNIVERSITÀ DI ROMA

# Exercises

- Voronoi basis



## Gromov-Hausdorff distance

$$d_{\mathcal{GH}}(\mathcal{X}, \mathcal{Y}) = \min_{\mathcal{Z}, f, g} d_{\mathcal{H}}^{\mathcal{Z}}(f(\mathcal{X}), g(\mathcal{Y}))$$

where  $f : \mathcal{X} \rightarrow \mathcal{Z}$  and  $g : \mathcal{Y} \rightarrow \mathcal{Z}$  are isometric embeddings

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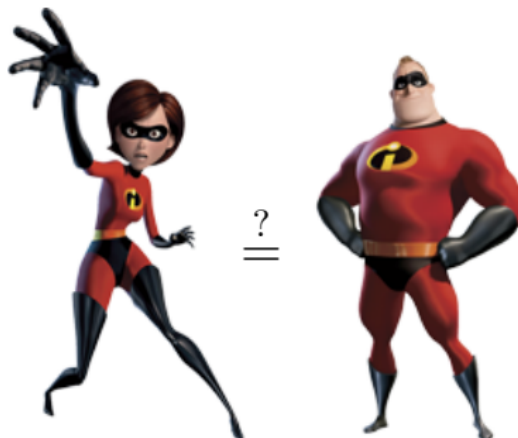
**Why** should we compute this distance?

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Why should we compute this distance? We can compare shapes

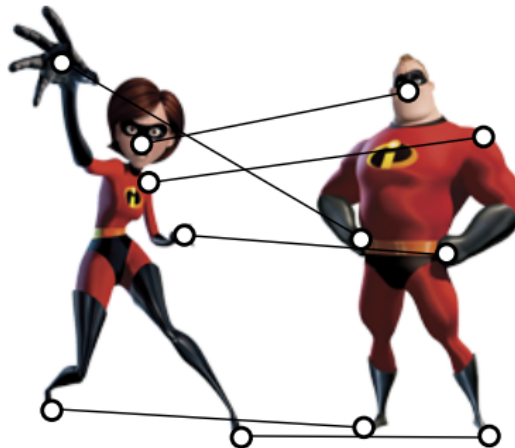


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[Why](#) should we compute this distance? We can find [correspondences](#)



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- It depends on the task (correspondence vs retrieval)

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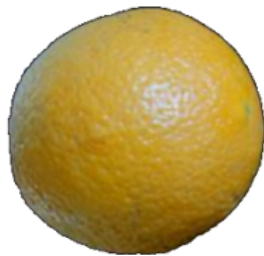
- It depends on the task (correspondence vs retrieval)
- It is convenient to compute approximations of  $d_{\mathcal{GH}}$
- Still an open problem! AKA shape matching

## Example: Texture transfer

With a correspondence we can **transfer** information (e.g. texture coordinates) across shapes



$\mathcal{X}$



$\mathcal{Y}$



texture of  $\mathcal{Y}$   
transferred to  $\mathcal{X}$

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# Correspondence

A **correspondence** between two sets  $\mathcal{X}$  and  $\mathcal{Y}$  is a set  $\mathcal{R} \subset \mathcal{X} \times \mathcal{Y}$  satisfying:

- For every  $x \in \mathcal{X}$ , there exists at least one  $y \in \mathcal{Y}$  such that  $(x, y) \in \mathcal{R}$
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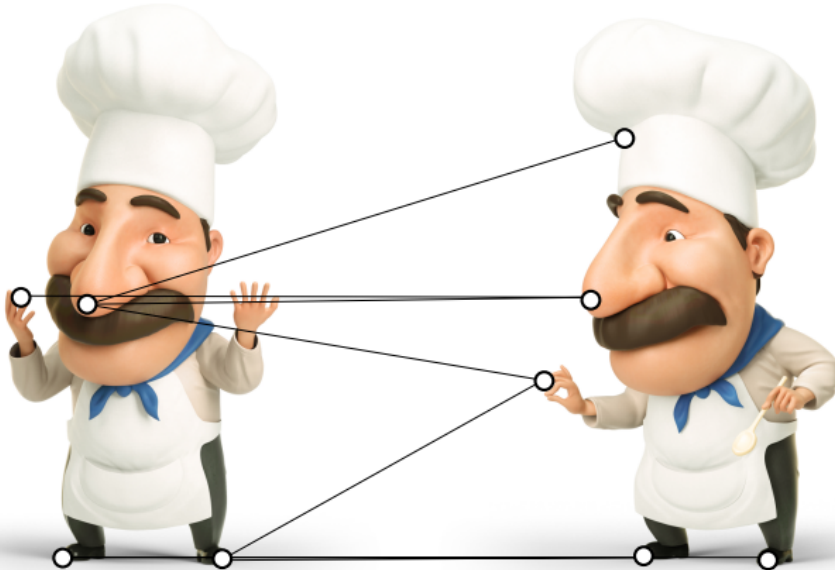
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In other words, this is giving **at least one** match for **all** points

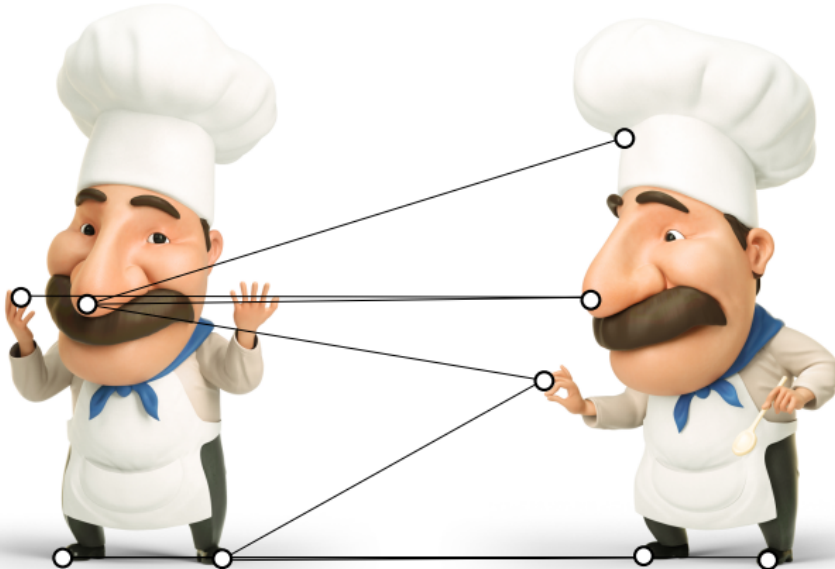
Later on, we will see that **mapping constraints** (e.g. one-to-one correspondence) are often necessary

## Example: Correspondence



Correspondence: at least one match for each sample

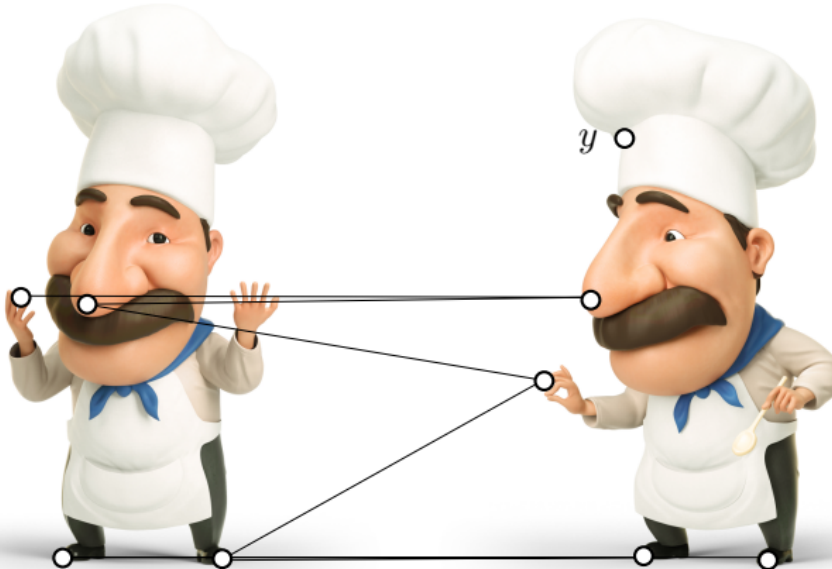
## Example: Correspondence



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**Note:** The mathematical definition of correspondence doesn't have anything to do with **distortion** or **semantics**

## Example: Correspondence



**Not** a correspondence: point  $y \in \mathcal{Y}$  is not mapped

**Note:** The mathematical definition of correspondence doesn't have anything to do with **distortion** or **semantics**

## Correspondence vs. maps

Any **surjective map**  $f : \mathcal{X} \rightarrow \mathcal{Y}$  defines a correspondence:

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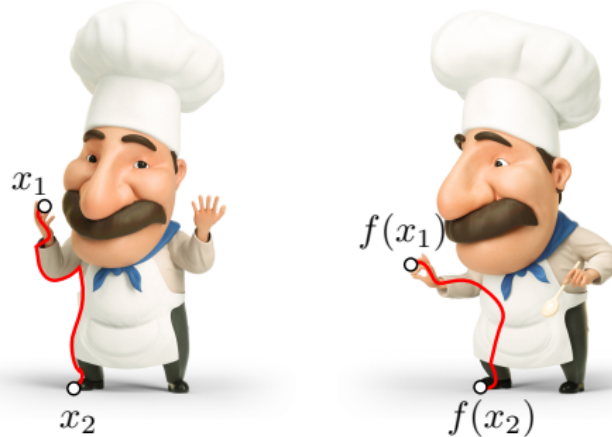
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Intuitively, a correspondence is a “multi-valued” map where a single point may have more than one image

# Metric distortion

For a map  $f : \mathcal{X} \rightarrow \mathcal{Y}$  between metric spaces  $(\mathcal{X}, d_{\mathcal{X}})$  and  $(\mathcal{Y}, d_{\mathcal{Y}})$  we define its (absolute) **distortion** as:

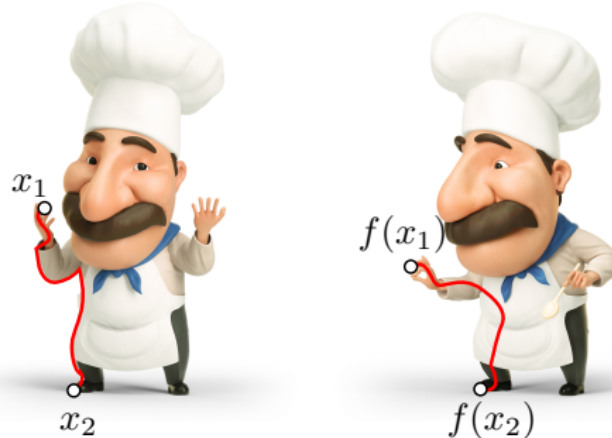
$$\text{dis}(f) = \max_{x_1, x_2 \in \mathcal{X}} |d_{\mathcal{X}}(x_1, x_2) - d_{\mathcal{Y}}(f(x_1), f(x_2))|$$



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Similarly, for a **correspondence**  $\mathcal{R}$  we define:

$$\text{dis}(\mathcal{R}) = \max\{|d_{\mathcal{X}}(x_1, x_2) - d_{\mathcal{Y}}(y_1, y_2)| : (x_1, y_1), (x_2, y_2) \in \mathcal{R}\}$$

## Metric distortion

For a surjective map  $f : \mathcal{X} \rightarrow \mathcal{Y}$  and the associated correspondence  $\mathcal{R} = \{(x, f(x)) : x \in \mathcal{X}\}$ , we have:

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Finally, if

$$\text{dis}(f) \leq \epsilon$$

we call  $f$  an  **$\epsilon$ -isometry** (which is what we see in the real world)

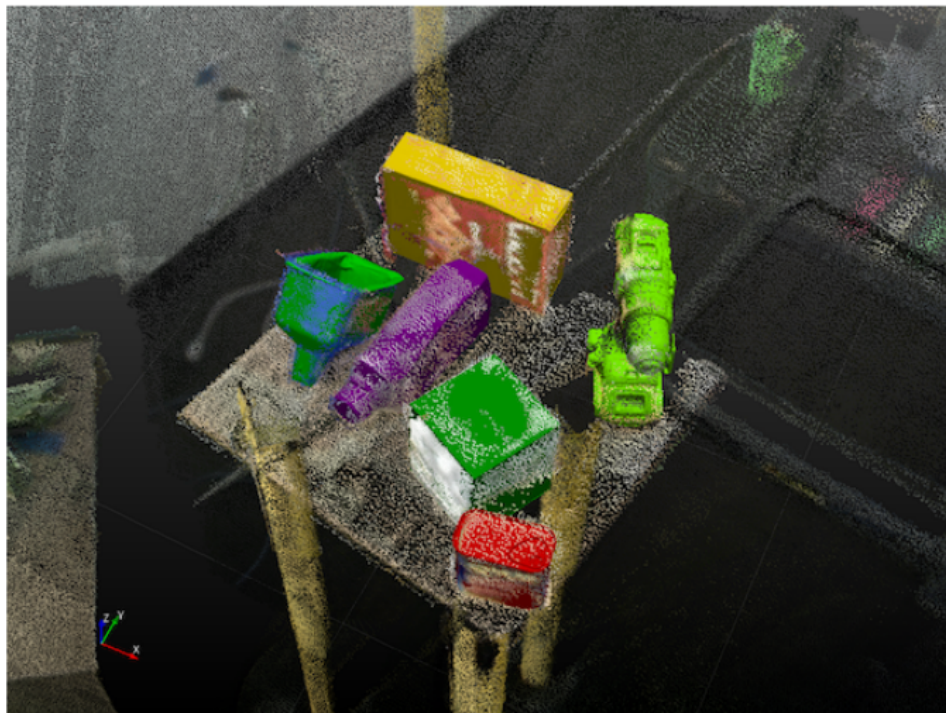
## Rigid correspondence

We first consider the simple case where  $d_{\mathcal{X}} = d_{\mathcal{Y}} = \|\cdot\|_2$

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If we want zero distortion, we then expect our shapes to be **rigid**



## Rigid alignment

We consider the Hausdorff distance:

$$d_{\mathcal{H}}(\mathcal{X}, \mathcal{Y}) = \max\{\max_x \text{dist}_{\mathbb{R}^3}(x, \mathcal{Y}), \max_y \text{dist}_{\mathbb{R}^3}(y, \mathcal{X})\}$$

In particular, its **non-symmetric** version:

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**Rigid alignment** methods minimize  $d_{\mathcal{H}}$  over all rigid isometries:

$$\min_{i \in \text{iso}(\mathbb{R}^3)} d_{\mathcal{H}}(i(\mathcal{X}), \mathcal{Y})$$

Shape  $\mathcal{X}$  is moved in  $\mathbb{R}^3$  until the Hausdorff distance to  $\mathcal{Y}$  is minimum

# Iterative closest point (ICP)

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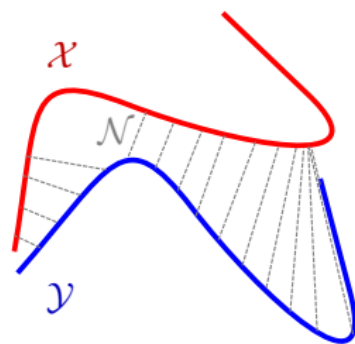
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- Repeat

Besl and McKay, "A Method for Registration of 3-D Shapes". TPAMI 14(2), 1992

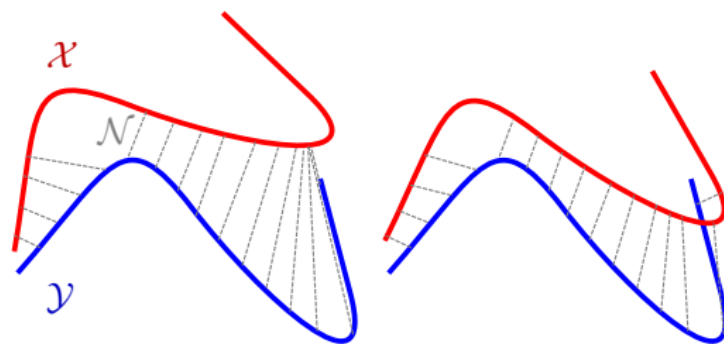
## Example: ICP

At each step, we have both a correspondence and a rigid transformation



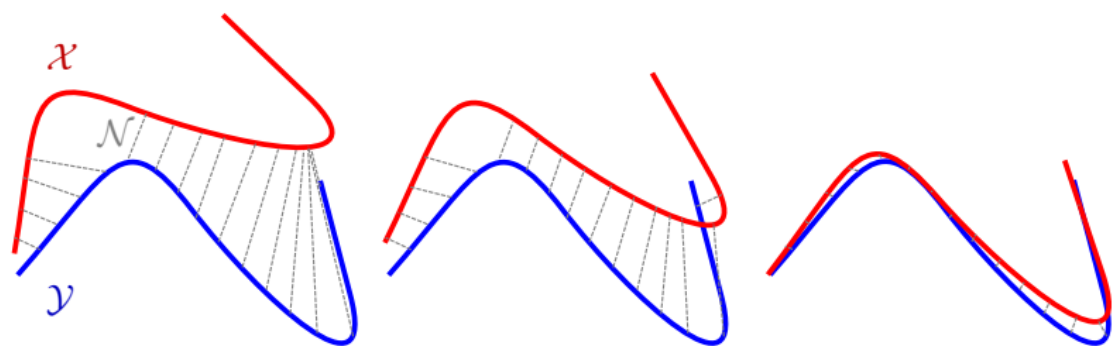
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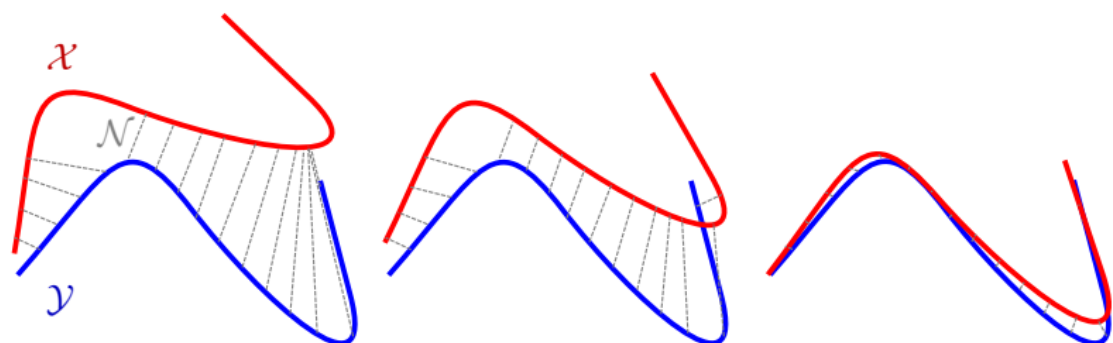
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- In practice converges to local optima
- Requires a good initialization
- Only works well in the rigid setting

## Correspondence and Gromov-Hausdorff

One can prove the following relationship<sup>1</sup>:

$$d_{\mathcal{GH}}(\mathcal{X}, \mathcal{Y}) < r \iff |d_{\mathcal{X}}(x_1, x_2) - d_{\mathcal{Y}}(y_1, y_2)| < 2r$$

for all  $(x_1, y_1), (x_2, y_2) \in \mathcal{R}$ . More formally, the above holds when such a correspondence  $\mathcal{R} \subset \mathcal{X} \times \mathcal{Y}$  exists.

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This allows us to speak about  $d_{\mathcal{GH}}$  just by using [correspondences](#)  $\mathcal{R}$ :

$$d_{\mathcal{GH}}(\mathcal{X}, \mathcal{Y}) = \frac{1}{2} \min_{\mathcal{R}} \text{dis}(\mathcal{R})$$

Still huge!

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## Correspondence computation

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Also, restrict the feasible set to the set of **one-to-one** correspondences;  
this is done by optimizing over **permutations**:

$$d_{\mathcal{P}}(\tilde{\mathcal{X}}, \tilde{\mathcal{Y}}) = \frac{1}{2} \min_{\pi \in \mathcal{P}_n} \max_{1 \leq i, j \leq n} |d_{\mathcal{X}}(x_1, x_2) - d_{\mathcal{Y}}(y_{\pi i}, y_{\pi j})|$$

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Q: how big is the set of permutations?

## Correspondence computation

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this is done by optimizing over **permutations**:

$$d_{\mathcal{P}}(\tilde{\mathcal{X}}, \tilde{\mathcal{Y}}) = \frac{1}{2} \min_{\pi \in \mathcal{P}_n} \max_{1 \leq i, j \leq n} |d_{\mathcal{X}}(x_1, x_2) - d_{\mathcal{Y}}(y_{\pi i}, y_{\pi j})|$$

Q: how big is the set of permutations?  $n!$

In fact, one can prove that  $d_{\mathcal{GH}}(\mathcal{X}, \mathcal{Y}) \leq d_{\mathcal{P}}(\tilde{\mathcal{X}}, \tilde{\mathcal{Y}})$

## Example: Sort by distortion

Assume we know the correspondence  $\pi$ , then we can directly compute:

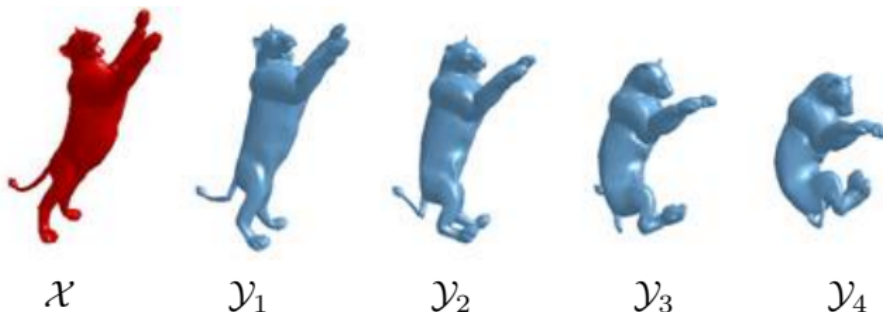
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And we can use it to sort shapes by distortion:



where

$$d_{\mathcal{P}}(\mathcal{X}, \mathcal{Y}_1) \leq d_{\mathcal{P}}(\mathcal{X}, \mathcal{Y}_2) \leq d_{\mathcal{P}}(\mathcal{X}, \mathcal{Y}_3) \leq d_{\mathcal{P}}(\mathcal{X}, \mathcal{Y}_4)$$

## Correspondence matrix

A common representation is the **correspondence matrix**  $\mathbf{T} \in \{0, 1\}^{m \times n}$

$$\mathbf{T} = \begin{pmatrix} 0 & 1 & \cdots & 0 \\ 1 & 0 & \cdots & 1 \\ \vdots & & & \vdots \\ 0 & 0 & \cdots & 1 \end{pmatrix}$$

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In other words, it maps functions to functions

## Permutation matrix

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In turn, one-to-one correspondences are associated to **bijective maps**  
 $f : \mathcal{X} \rightarrow \mathcal{Y}$

## Correspondence problem

$$\pi^* = \arg \min_{\pi \in \mathcal{P}_n} \max_{1 \leq i, j \leq n} |d_{\mathcal{X}}(x_1, x_2) - d_{\mathcal{Y}}(y_{\pi i}, y_{\pi j})|$$

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The metric distortion terms can be encoded into a matrix  $\mathbf{C} \in \mathbb{R}^{n^2 \times n^2}$ :

$$C_{(i\ell)(jm)} = |d_{\mathcal{X}}(x_i, x_j) - d_{\mathcal{Y}}(y_\ell, y_m)|$$

where  $(i\ell)$  and  $(jm)$  identify the matches  $(x_i, y_\ell), (x_j, y_m) \in \mathcal{R}$

$$\mathbf{C} = \begin{pmatrix} (x_1, y_1) & (x_3, y_5) & \cdots & (x_i, y_\ell) & \cdots \\ & & & \vdots & \\ & & & \vdots & \\ & \cdots & \cdots & \cdots & |d_{\mathcal{X}}(x_i, x_j) - d_{\mathcal{Y}}(y_\ell, y_m)| & \cdots \\ & & & & \vdots & \\ \end{pmatrix} \begin{pmatrix} (x_1, y_1) \\ (x_3, y_5) \\ \vdots \\ (x_j, y_m) \\ \vdots \end{pmatrix}$$

## $L_p$ Gromov-Hausdorff

$$\frac{1}{2} \min_{\mathbf{P} \in \mathcal{P}} \max_{i,j,\ell,m} C_{(i\ell)(jm)} P_{ij} P_{\ell m}$$

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$$\frac{1}{2} \min_{\mathbf{P} \in \mathcal{P}} \sum_{i,j,\ell,m} C_{(i\ell)(jm)} P_{ij} P_{\ell m}$$

Or, more in general, we can use the  $L_p$  distortion

$$C_{(i\ell)(jm)}^{(p)} = |d_{\mathcal{X}}(x_i, x_j) - d_{\mathcal{Y}}(y_\ell, y_m)|^p$$

yielding the minimization problem

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# Quadratic assignment problem

Rewriting in matrix notation, we get:

$$\begin{aligned} \min_{\mathbf{P} \in \{0,1\}^{n \times n}} & \text{vec}(\mathbf{P})^\top \mathbf{C} \text{vec}(\mathbf{P}) \\ \text{s.t. } & \mathbf{P}\mathbf{1} = \mathbf{1}, \mathbf{P}^\top \mathbf{1} = \mathbf{1} \end{aligned}$$

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also known as **quadratic assignment problem** (QAP)

- The problem is **quadratic** in  $\mathbf{P}$
- It is **NP-hard**
- Still an open problem (a survey might be a final project)

## Continuous relaxation

Replace  $\{0, 1\}$  with continuous values in  $[0, 1]$

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We are replacing permutations by **doubly-stochastic** matrices:

$$\mathbf{S} = \begin{pmatrix} 0.2 & 0.3 & \cdots & 0.5 \\ 0.1 & 0 & \cdots & 0 \\ \vdots & & & \vdots \\ 0.7 & 0 & \cdots & 0.1 \end{pmatrix}$$

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Doubly-stochastic matrices are **convex combinations** of permutations:

$$\mathbf{S} = t\mathbf{P}_1 + (1 - t)\mathbf{P}_2, \quad t \in [0, 1]$$

## Example: Soft maps

Each row / column of a soft map can be interpreted as a probability distribution on the shape:



## Exercise: Sort by distortion

Load the shapes `tr_reg_000`, `tr_reg_001`, `tr_reg_002`, and `tr_reg_003` (download from course website).

- The **ground-truth** correspondence  $T^{\mathcal{X}, \mathcal{Y}}$  between each pair of shapes  $(\mathcal{X}, \mathcal{Y})$  is the **identity map**

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- Sort the shapes by increasing distortion

## Exercise: ICP

Load the shapes bun000 and bun045 (download from course website).

- Compute a **FPS** of  $n =$  a few hundred points for each scan

Visualize the alignment with the entire shapes (not just the farthest point samplings) **without** running ICP on the entire shapes

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- Print the alignment error across the iterations

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## Suggested reading

- An extensive treatment of Gromov-Hausdorff distances and correspondences is in Chapter 7 of:

Burago, Burago, Ivanov, “A course in metric geometry”. AMS, 2001

- A very accessible survey of ICP and its variants:

Rusinkiewicz and Levoy, “Efficient variants of the ICP algorithm”.  
Proc. 3DIM, 2011