





Université de Caen Basse-Normandie

Graph Classification

Septempber 3, 2019 Salerno

Pros and cons of graphs Graph Metrics Graph Neural Networks Conclusion Bibliography

Pros and cons of graphs

Graph Metrics

- Graph alignment metric: Orbifold Space
- Graph Edit Distance: Krein Spaces
- Graph Kernels: Hilbert Spaces

③ Graph Neural Networks

Agregation

Conclusion





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- Interest: provide a compact encoding of both :
 - a decomposition of objects into meaningful sub-parts,
 - the relationships between these sub-parts.
- Applications:
 - Image processing: segmentation, boundary detection,
 - Pattern Recognition: printed characters, documents, objects (buildings, brain structures), faces, gestures, molecules,...,
 - Image registration,
 - Understanding of structured scenes.
 - . . .





- Even simple questions are difficult:
 - Are this two graphs the same ?
 - \mathcal{NP} -intermediate
 - Is this graph a sub-part of this graph ?
 - $\bullet \ \mathcal{NP}\text{-complete}$
 - What is the distance between these two graphs ?
 - \mathcal{NP} -hard (for the usual distance)
 - What is the mean/median of a set of graphs ?
 - \mathcal{NP} -hard (for the usual distance)
- Pattern Recognition implies:
 - Metrics,
 - In the security of the secu

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Graph space as an Orbifold

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[Jain, 2016, Jain and Wysotzki, 2004, Jain, 2014].

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Orbifolds theory: basics

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- Let \mathcal{T} be the manifold of $n \times n$ matrices.
- One graph may have multiple matrix representations



• Let \mathcal{P} denote the set of $n \times n$ permutation matrices:

$$\forall (x,y) \in \mathcal{T} \ x \sim y \Leftrightarrow \exists P \in \mathcal{P} | x = P^t y P$$

- ${\cal T}/\sim$ is called an orbifold
- A graph G is encoded in \mathcal{T}/\sim by $[X_G] = \{x, y, z, \dots\}$.



Orbifolds theory: basics

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• Let X and Y denote two elements of \mathcal{T}/\sim :

$$\begin{cases} < X, Y > = \max\{< x, y > x \in X, y \in Y\} \\ \delta(X, Y) = \sqrt{\|X\|^2 + \|Y\|^2 - 2 < X, Y >} \\ = \min_{x \in X, y \in Y} \|x - y\| \end{cases}$$

This metric is called the graph alignment metric.

- Using real attributes, the space (\mathcal{G}, δ) is:
 - A complete metric space,
 - a geodesic space,
 - locally compact,
 - every closed bounded subset of (\mathcal{G}, δ) is compact.

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Orbifold theory: Applications

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- Computation of the sample mean [Jain, 2016]
 - The sample mean of a set of graphs always exists.
 - Under some conditions, the set of sample means reduces to a singleton.
- Central clustering algorithms [Jain and Wysotzki, 2004],
- Generalized linear classifiers [Jain, 2014]



- The enumeration of [X] requires n! computations.
- Graph metric is induced by graph representation: Both concepts can not be distinguished.



$$\delta(G_1, G_2) = \sqrt{a^2 + \alpha^2}$$

We should search for a more flexible metric.



Graph Edit distance

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Pros and cons of graphs Graph Metrics Graph Neural Networks Conclusion Bibliography Edit Paths

Definition (Edit path)

Given two graphs G_1 and G_2 an **edit path** between G_1 and G_2 is a sequence of node or edge removal, insertion or substitution which transforms G_1 into G_2 .



A substitution is denoted $u \rightarrow v$, an insertion $\epsilon \rightarrow v$ and a removal $u \rightarrow \epsilon$.

Alternative edit operations such as merge/split have been also proposed[Ambauen et al., 2003].

Costs Graph Metrics Graph MetricsGraph Metrics

- All cost are positive: $c() \ge 0$,
- A node or edge substitution which does not modify a label has a 0 cost: c(l→ l) = 0.



If all costs are equal to 1, the cost of this edit path is equal to 5.

Conversely to graph alignment metric, costs allow to distinguish graph representation and graph metrics.

Pros and cons of graphs



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Definition (Graph edit distance)

The graph edit distance between G_1 and G_2 is defined as the cost of the less costly path within $\Gamma(G_1, G_2)$. Where $\Gamma(G_1, G_2)$ denotes the set of edit paths between G_1 and G_2 .

$$d(G_1, G_2) = \min_{\gamma \in \Gamma(G_1, G_2)} cost(\gamma) = \min_{\gamma \in \Gamma(G_1, G_2)} \sum_{e \in \gamma} c(e)$$

An $\mathcal{NP}\text{-hard}$ problem.



Graph edit distance: Main meth Gas Metrics Graph Retrict Graph Netrics

Conclusion Bibliography

- A* like algorithms [Abu-Aisheh, 2016],
- Formulation as a quadratic problem [Bougleux et al., 2017] solved by Franck-Wolfe [Frank and Wolfe, 1956] like algorithms (see also:
 [Liu and Qiao, 2014, Boria et al., 2018, Daller et al., 2018].
- Integer Programming [Lerouge et al., 2017, Darwiche, 2018]
- Fast (and often rough) approximations [Riesen and Bunke, 2009, Gaüzère et al., 2014a, Carletti et al., 2015, Blumenthal et al., 2018]

Pros and cons of graphs Graph edit distance: Comparison Graph Metrics Graph Neural Networks

Conclusion Bibliography

• Relative comparison:





Graph Edit distance: Conclusion Graph Metrics Graph Metrics

- Allows to dissociate graph representation and graph metric.
- \bigcirc Constitutes a fine and intuitive metric between graphs.
- $\stackrel{\textbf{R}}{\mathrel{>}}$ It is \mathcal{NP} -hard to compute,
- 🙁 It is not conditionally definite negative (Krein space).
 - Most of machine learning machinery should be adapted [Loosli et al., 2016].
 - Weak properties: E.g. the median is usually not unique.

Library: https://github.com/Ryurin/Python_GedLib



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Input Space

Feature Space



• A kernel k is a symmetric similarity measure on a set χ

$$\forall (x,y) \in \chi^2, \ k(x,y) = k(y,x)$$

• k is said to be **definite positive** (d.p.) iff k is symmetric and iff:

$$\begin{array}{rcl} \forall (x_1,\ldots,x_n) & \in & \chi^n \\ \forall (c_1,\ldots,c_n) & \in & \mathbb{R}^n \end{array} \right\} \sum_{i=1}^n \sum_{j=1}^n c_i c_j k(x_i,x_j) \ge 0$$

• $\mathcal{K} = (k(x_i, x_j))_{(i,j) \in \{1,...,n\}}$ is the Gramm matrix of k. k is d.p. iff:

$$\forall c \in \mathbb{R}^n - \{0\}, \ c^t K c \ge 0$$



Kernels and scalar products

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[Aronszahn, 1950] :

A kernel k is d.p. on a space χ if and only if it exists

• one Hilbert space $\mathcal H$ and • a function $\varphi: \chi \to \mathcal H$ such that:

$$k(x,y) = < \varphi(x), \varphi(y) >$$

Open the way for rich interactions between graphs and usual machine learning methods: SVM, kPCA, MKL, \ldots



Graph Kernel: methods

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$$K(G, G') = \sum_{x \in \mathcal{B}(G)} \sum_{y \in \mathcal{B}(G')} k(x, y)$$

where $\mathcal{B}(G)$ is a bag of patterns deduced from G [Haussler, 1999].

- Linear Patterns: Random Walk Kernel *n* order path kernel: Shortest Path
- [Kashima et al., 2003, Gärtner et al., 2003 [Ralaivola et al., 2005, Dupé and Brun, 20 [Hermansson et al., 2015]
- Non linear patterns:

Tree Pattern kernel

Graphlet Kernel Treelet kernels [Mahé and Vert, 2009, Shervashidze and Borgwardt, 2009] [Shervashidze et al., 2009] [Gaüzère et al., 2014b]



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Method	RMSE(°C)	Time(s)	
		Learning	Prediction
Gaussian edit distance	10.27	1.35	0.05
Random Walks	18.72	19.10	0.57
Path Kernel	12.24	7.83	0.18
Tree Pattern Kernel	11.02	4.98	0.03
Treelet Kernel (TK)	8.10	0.07	0.01
TK + MKL	5.24	70	0.01

Boiling point prediction on acyclic molecule dataset using 90% of the dataset as train set and remaining 10% as test set.



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- Pros:
 - Graph kernels provide an implicit embedding of graphs,
 - Its open the way to the application of many statistical tools to graphs,
- Cons:
 - Graph kernels are usually based on a notion of bag which only provides a rough similarity measure.
 - The graph feature extraction process has been moved to the design of a similarity measure (the kernel). Such a measure remains largely "handcrafted".
- Libraries:
 - https://github.com/jajupmochi/py-graph
 - http://chemcpp.sourceforge.net/html/index.html



Graph Neural Network

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Graph Neural Networks: Three normal Networks: Three

Conclusion Bibliography

- Agregation,
- Oecimation,
- Pooling





$$\begin{cases} h_{v} = f_{w}(I_{v}, I_{CON(v)}, h_{\mathcal{N}(v)}, I_{\mathcal{N}(v)}) \\ o_{v} = g_{w}(h_{v}, I_{v}) \end{cases}$$

with
$$CON(v) = \{(v, v') | v' \in \mathcal{N}(v)\}$$



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$$\begin{cases} h_v^t = f_w(l_v, l_{CON(v)}, h_{\mathcal{N}(v)}^{t-1}, l_{\mathcal{N}(v)}) \\ o_v = g_w(h_v^T, l_v) \end{cases}$$



Pros and cons of graphs Graph Metrics Graph Neural Networks Conclusion Bibliograph he problem

• Using images we learn $w_0 \ldots, w_8$:

<i>W</i> ₅	W_1	W ₆
W3	w ₀	W4
W7	<i>W</i> ₂	W ₈

 w_1 denotes the weigh of the pixel above the central pixel.



Without embedding nothing distinguishes the cyan, red and green neighbors.

How to become permutation invariants Graphicer Conclusion



Bibliography

$$h_{v}^{t} = f_{w}(I_{v}, I_{CON(v)}, h_{\mathcal{N}(v)}^{t-1}, I_{\mathcal{N}(v)})$$
$$h_{v}^{t} = \sum_{v' \in \mathcal{N}(v)} f(I_{v}, I_{v,v'}, I_{v'}, h_{v'}^{(t-1)})$$

where f may be:

• An affine function [Scarselli et al., 2009],

$$f(l_{v}, l_{v,v'}, l_{v'}, h_{v'}^{(t-1)}) = A^{(l_{v}, l_{v,v'}, l_{v'})} h_{v'}^{(t-1)} + b^{(l_{v}, l_{v,v'}, l_{v'})}$$

• A MLP [Massa et al., 2006]



More complex agregation functions of graphs Conclusion Graph Neural Networks Conclusion Bibliography

- A long Short-term Memory [Hochreiter and Schmidhuber, 1997, Peng et al., 2017, Zayats and Ostendorf, 2018]
- A Gated Reccurent Unit [Li et al., 2016]

$$\begin{aligned} h_{v}^{(1)} &= [x_{v}^{T}, 0] & (1) \\ a_{v}^{(t)} &= A_{v}^{T} [h_{1}^{(t-1)} , \dots, h_{|V|}^{(t-1)}]^{T} + b & (2) \\ z_{v}^{t} &= \sigma (W^{z} a_{v}^{(t)} + U^{z} h_{v}^{(t-1)}) & (3) \\ r_{v}^{t} &= \sigma (W^{r} a_{v}^{(t)} + U^{r} h_{v}^{(t-1)}) & (4) \\ \tilde{h}_{v}^{(t)} &= \tanh \left(W a_{v}^{(t)} + U \left(r_{v}^{t} \odot h_{v}^{(t-1)} \right) \right) & (5) \\ h_{v}^{t} &= (1 - z_{v}^{t}) \odot h_{v}^{(t-1)} + z_{v}^{t} \odot \tilde{h}_{v}^{t} & (6) \end{aligned}$$

$$\begin{split} & z_{v}^{t}: \text{ update gate, } r_{v}^{t}: \text{ reset gate, } A_{v}: \text{ weight by edges types.} \\ \bullet \text{ Learned weight by edge type:} \\ & a_{v}^{(t)} = \sum_{w \in \mathcal{N}(v)} A_{l_{v,w}} h_{w}^{(t-1)} \text{ [Gilmer et al., 2017]} \end{split}$$



Graph attention Networks

Pros and cons of graphs Graph Metrics Graph Neural Networks Conclusion [Velickövic^pet al., 2018]

• Not all neighbors have a same importance for update:

$$\alpha_{v,v'} = softmax_{v'}(e_{v,v'}) = \frac{exp(e_{v,v'})}{\sum_{v'' \in \mathcal{N}_i} exp(e_{v,v''})}$$



- With : e_{v,v'} = LeakyReLU(a^T[Wh_v||Wh_{v'}])
 a, W : weight vector and matrix.
- Update rule:

$$\mathbf{h}'_{\mathbf{v}} = \sigma(\sum_{\mathbf{v}' \in \mathcal{N}_{\mathbf{v}}} \alpha_{\mathbf{v},\mathbf{v}'} W \mathbf{h}_{\mathbf{v}'})$$

• With K features:

$$h'_{\mathbf{v}} = ||_{k=1}^{\kappa} \sigma(\sum_{\mathbf{v}' \in \mathcal{N}_{\mathbf{v}}} \alpha_{\mathbf{v},\mathbf{v}'}^{k} W^{k} h_{\mathbf{v}'})$$



Graph Convolution

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Input image



Convolution Kernel

$$\begin{bmatrix} -1 & -1 & -1 \\ -1 & 8 & -1 \\ -1 & -1 & -1 \end{bmatrix}$$

Feature map



• Graph Laplacian:

$$L = D - A$$
 with $D_{ii} = \sum_{j=1}^{n} A_{ij}$

A adjacency matrix of a graph G.

• Matrix *L* is real symmetric semi definite positive:

 $L = U\Lambda U^T$

U orthogonal, Λ real(positive) diagonal matrix.

• A classical result from signal processing:

$$x * y = \mathcal{F}^{-1}(\hat{x}.\hat{y})$$

*: convolution operation, \mathcal{F}^{-1} inverse Fourrier transform, \hat{x} fourrier transform of x, \ddot{x} term by term multiplication.

(GREYC)



Graph Convolution

Pros and cons of graphs Graph Metrics Graph Neural Networks Conclusion The spectral approach

• If x is a signal on G, $\hat{x} = U^T x$ can be considered as its "Fourrier" transform. We have:

$$U\hat{x} = UU^T x = x$$

U is thus the inverse Fourrier transform.

• By analogy:

$$z * x = U(\hat{z} \odot \hat{x}) = U\left(U^{\mathsf{T}} z \odot U^{\mathsf{T}} x\right) = U\left(diag(U^{\mathsf{T}} z)U^{\mathsf{T}} x\right)$$

 \odot : Hadamard product.

• Let $g_{\theta}(\Lambda)$ be a diagonal matrix. The filtering of x by g_{θ} is:

$$y = U\left(g_{\theta}(\Lambda)U^{\mathsf{T}}x\right) = \left(Ug_{\theta}(\Lambda)U^{\mathsf{T}}\right)x$$



• If:

$$g_{ heta}(\Lambda) = \sum_{i=0}^{K-1} heta_i \Lambda^i$$

Then:

$$y = \left(Ug_{\theta}(\Lambda) U^{T} \right) x = U\left(\sum_{i=0}^{K-1} \theta_{i} \Lambda^{i} \right) U^{T} x = \left(\sum_{i=0}^{K-1} \theta_{i} L^{i} \right) x$$

- One parameter per ring:
 - Lx : one step (direct) neighborhood,
 - L^2x : two step neighborhood (idem for $L^3, L^4, ...$)
- Problem: Computing Lⁱ for i ∈ {0,..., K − 1} is problematic for large matrices (SVD computation)



• Let us consider Chebyshev polynomial
$$T_k(x) = 2xT_{k-1}(x) - T_{k-2}(x)$$
, with $T_0 = 1$ and $T_1(x) = x$.

$$g_{ heta}(\Lambda) = \sum_{i=0}^{K-1} heta_i \Lambda^i o g_{ heta}(\Lambda) = \sum_{i=0}^{K-1} heta_i T_i(ilde{\Lambda})$$

 $\tilde{\Lambda}$ normalized version of $\Lambda.$

• we have:

$$ilde{x}_k = 2 ilde{L} ilde{x}_{k-1} - ilde{x}_{k-2}$$
 with $ilde{x}_0 = x$ and $ilde{x}_1 = ilde{L}x$

 $\mathcal{O}(\mathcal{K}|\mathcal{E}|)$ operations to get \tilde{x}_k .

• If K = 2 it simplifies to [Kipf and Welling, 2017]: $y = \theta L' x$ where L' is a regularized version of the normalized Laplacian.

(GREYC)



• [Simonovsky and Komodakis, 2017]

$$y_i = rac{1}{|\mathcal{N}(i)|} \sum_{j \in \mathcal{N}(i)} F_{ heta}(L(j,i)) x_j + b$$

F: Parametric function of θ which associates one weigh to each edge label L(j, i).

• [Verma et al., 2017]:

$$y_i = rac{1}{|\mathcal{N}(i)|} \sum_{m=1}^M \sum_{j \in \mathcal{N}(i)} q_{\theta_m}(x_j, x_i) W_m x_j + b$$

 $q_{\theta_m}(.,.)$ m^{th} learned soft-assignment function. W_m weight matrix.

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Graph Propagation

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Recurent networks

[Hochreiter and Schmidhuber, 1997] [Massa et al., 2006] [Scarselli et al., 2009] [Li et al., 2016] [Gilmer et al., 2017] [Peng et al., 2017] [Zayats and Osten

Convolution

 al., 2009]
 [Bruna et al., 2014]

 , 2016]
 [Defferrard et al., 2016]

 ner et al., 2017]
 [Kipf and Welling, 2017]

 Peng et al., 2017]
 [Simonovsky and Komodakis, 2017]

 [Zayats and Ostendorf, 2018]
 [Verma et al., 2017]





• Graph Downsampling, Graph pooling, Graph final decision: Some solutions but still the jungle.





- Graph neural network:
 - Still in their infancy,
 - A great potential.

Pros and cons of graphs Graph Metrics Graph Neural Networks Conclusion **Bibliography**

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