Unsupervised Learning on Graphs

Spectral Clustering: Graph = Matrix

	Α	B	С	D	Е	F	G	н	I	J
Α		I		I		I				
В	I		I							
С		I								
D	I					I				
Е						I				
F	I			I	I					
G									I	
н							I		I	I
							I	I		I
J								L	I	



Spectral Clustering: Graph = Matrix Transitively Closed Components = "Blocks"



sometimes called a **block-stochastic** matrix:

- each node has a latent "block"
- fixed probability qi for links between elements of block i
- fixed probability q0 for links between elements of different blocks



Of course we can't see the "blocks" unless the nodes are sorted by cluster...

Spectral Clustering: Graph = Matrix Vector = Node → Weight



Spectral Clustering: Graph = Matrix $M^*v_1 = v_2$ "propogates weights from neighbors"



Spectral Clustering: Graph = Matrix W*v₁ = v₂ "propogates weights from neighbors"

	W	: norn	nalize	ed so	colu	mns s	sum t	οI		W	* ,	V ₁	V ₂		
	Α	В	С	D	Е	F	G	н	I	J					
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J							.5	.5	.3	_	J		J		

Spectral Clustering: Graph = Matrix W*v₁ = v₂ "propogates weights from neighbors"

 $\mathbf{W} \cdot \mathbf{v} = \lambda \mathbf{v} : \mathbf{v}$ is an eigenvector with eigenvalue λ

Q: How do I pick **v** to be an eigenvector for a blockstochastic matrix?



Spectral Clustering: Graph = Matrix W*v₁ = v₂ "propogates weights from neighbors" W · v = λ v : v is an eigenvector with eigenvalue λ

How do I pick **v** to be an eigenvector for a blockstochastic matrix?



Spectral Clustering: Graph = Matrix W*v₁ = v₂ "propogates weights from neighbors"

 $\mathbf{W} \cdot \mathbf{v} = \lambda \mathbf{v} : \mathbf{v}$ is an eigenvector with eigenvalue λ



Spectral Clustering: Graph = Matrix $W^*v_1 = v_2$ "propogates weights from neighbors" $\mathbf{W} \cdot \mathbf{v} = \lambda \mathbf{v}$: v is an eigenvector with eigenvalue λ e₂ $^{\mathrm{a}}$ 0.4 0.2 0.0 D.4 -0.2 0000000 e₃ -0.4 e₁ -D.Z 0,000 -D.4 \mathbf{e}_2 -0.4 -0.2 0.2 0 -0.6 Ш 5 1D 15 20 seg.1 seg.2 seg.3

[Shi & Meila, 2002]

Spectral Clustering: Graph = Matrix W*v₁ = v₂ "propogates weights from neighbors" W · v = λ v : v is an eigenvector with eigenvalue λ

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If W is connected but roughly block diagonal with *k* blocks then

 the top eigenvector is a constant vector

• the next *k* eigenvectors are roughly piecewise constant with "pieces" corresponding to blocks



seg.1 seg.2 seg.3

Spectral Clustering: Graph = Matrix W*v₁ = v₂ "propogates weights from neighbors"

 $\mathbf{W} \cdot \mathbf{v} = \lambda \mathbf{v}$: v is an eigenvector with eigenvalue λ

If **W** is connected but roughly block diagonal with *k* blocks then

 the "top" eigenvector is a constant vector

• the next *k* eigenvectors are roughly piecewise constant with "pieces" corresponding to blocks Spectral clustering: • Find the top k+1eigenvectors $\mathbf{v}_1, \dots, \mathbf{v}_{k+1}$ • Discard the "top" one • Replace every node *a* with *k*-dimensional vector $x_a = \langle \mathbf{v}_2(a), \dots, \mathbf{v}_{k+1}(a) \rangle$ • Cluster with *k*-means

Spectral Clustering: Pros and Cons

- Elegant, and well-founded mathematically
- Tends to avoid local minima
 - Optimal solution to relaxed version of mincut problem (Normalized cut, aka NCut)
- Works quite well when relations are approximately transitive (like similarity, social connections)
- Expensive for very large datasets
 - Computing eigenvectors is the bottleneck
 - Approximate eigenvector computation not always useful
- Noisy datasets sometimes cause problems
 - Picking number of eigenvectors and k is tricky
 - "Informative" eigenvectors need not be in top few
 - Performance can drop suddenly from good to terrible

Experimental results: best-case assignment of class labels to clusters

			N	Cut	N	IW	
	Dataset	k	Accuracy	Macro-F1	Accuracy	Macro-F1	
	Iris	3	0.673	0.570	0.807	0.806	
	PenDigits01	2	1.000	1.000	1.000	1.000	
	PenDigits17	2	0.755	0.753	0.755	0.754	
	UBMCBlog	2	0.953	0.953	0.953	0.953	
L	AGBlog	2	0.520	0.342	0.520	0.342	
	20ngA	2	0.955	0.955	0.955	0.955	
	20ngB	2	0.505	0.344	0.550	0.436	
	20ngC	3	0.613	0.621		s. ⁵ , ,	. <u>^ / * /</u>
	20ngD	4	0.469	0.432	~ * *	t all t	i A A A A A A A A A A A A A A A A A A A
	Average	-	0.716	0.663	Start and a start of the start		
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Another way to think about spectral clustering....

Spectral Clustering: Graph = Matrix W*v₁ = v₂ "propogates weights from neighbors"

 $\mathbf{W} \cdot \mathbf{v} = \lambda \mathbf{v} : \mathbf{v}$ is an eigenvector with eigenvalue λ

- smallest eigenvecs of D-A are largest eigenvecs of A
- smallest eigenvecs of I-W are largest eigenvecs of W Suppose each *y*(*i*)=+1 or -1:
- Then *y* is a cluster indicator that splits the nodes into two
- what is $\mathbf{y}^{\mathsf{T}}(\mathsf{D}-\mathsf{A})\mathbf{y}$?

$$\mathbf{y}^{T}(D-A)\mathbf{y} = \mathbf{y}^{T}D\mathbf{y} - \mathbf{y}^{T}A\mathbf{y} = \sum_{i} d_{i}y_{i}^{2} - \sum_{i,j} a_{i,j}y_{i}y_{j}$$

$$= \frac{1}{2} \left[2\sum_{i} d_{i}y_{i}^{2} - 2\sum_{i,j} a_{i,j}y_{i}y_{j} \right]$$

$$= \frac{1}{2} \left[\sum_{i} \left(\sum_{j} a_{ij} \right) y_{i}^{2} + \sum_{j} \left(\sum_{i} a_{ij} \right) y_{j}^{2} - 2\sum_{i,j} a_{i,j}y_{i}y_{j} \right]$$

$$= \frac{1}{2} \left[\sum_{i,j} a_{ij}y_{i}^{2} + \sum_{i,j} a_{ij}y_{j}^{2} - 2\sum_{i,j} a_{i,j}y_{i}y_{j} \right]$$

$$= \frac{1}{2} \left[\sum_{i,j} a_{ij}(y_{i} - y_{j})^{2} \right] = \text{size of CUT}(\mathbf{y})$$

 $\mathbf{y}^{T}(I - W)\mathbf{y} = \text{size of NCUT}(\mathbf{y})$

NCUT: roughly minimize ratio of transitions between classes vs transitions within classes

Spectral Clustering: Graph = Matrix W*v₁ = v₂ "propogates weights from neighbors"

 $\mathbf{W} \cdot \mathbf{v} = \lambda \mathbf{v} : \mathbf{v}$ is an eigenvector with eigenvalue λ

- smallest eigenvecs of D-A are largest eigenvecs of A
- smallest eigenvecs of I-W are largest eigenvecs of W
 Suppose each y(i)=+1 or -1:
- Then y is a cluster indicator that cuts the nodes into two
- what is $\mathbf{y}^{\mathsf{T}}(\mathsf{D}-\mathsf{A})\mathbf{y}$? The cost of the graph cut defined by \mathbf{y}
- what is y^T(I-W)y ? <u>Also a cost of a graph cut defined by y</u>
 How to minimize it?
 - Turns out: to minimize $\mathbf{y}^T \times \mathbf{y} / (\mathbf{y}^T \mathbf{y})$ find *smallest* eigenvector of X
 - But: this will not be +1/-1, so it's a "relaxed" solution

Spectral Clustering: Graph = Matrix W*v₁ = v₂ "propogates weights from neighbors"

 $\mathbf{W} \cdot \mathbf{v} = \lambda \mathbf{v} : \mathbf{v}$ is an eigenvector with eigenvalue λ



Another way to think about spectral clustering....

- Most normal people think about spectral clustering like that - as relaxed optimization
- ...me, not so much
- I like the connection to "averaging"... because....

Spectral Clustering: Graph = Matrix W*v₁ = v₂ "propogates weights from neighbors"

	W	: norn	nalize	ed so	colu	mns s	sum t	οI		W	* ,	V ₁	V ₂		
	Α	В	С	D	Е	F	G	н	I	J					
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С	.3	.5	_								С	3	B	3*.3+3*.5	B
D				_	.5	.3					D		C	3*.33+2*.5	
Е				.5	_	.3					F		D		
F	.3			.5	.5	_					-		E		F
									2	2			F		F
G							-				G		G		
н								-	.3	.3	н		н		
I							.5	.5	-	.3	L		I		
J							.5	.5	.3	_	J		J		

Repeated averaging with neighbors as a clustering method

- Pick a vector v⁰ (maybe at random)
- Compute $v^1 = Wv^0$
 - i.e., replace v⁰[x] with weighted average of v⁰[y] for the neighbors y of x
- Plot v¹[x] for each x
- Repeat for v^2 , v^3 , ...
- Variants widely used for *semi-supervised* learning
 - HF/CoEM/wvRN average + clamping of labels for nodes with known labels
- Without clamping, will converge to constant v⁺
- What are the *dynamics* of this process?

Spectral Clustering: Graph = Matrix W*v₁ = v₂ "propogates weights from neighbors"

 $\mathbf{W} \cdot \mathbf{v} = \lambda \mathbf{v} : \mathbf{v}$ is an eigenvector with eigenvalue λ





- Create a graph, connecting all points in the 2-D initial space to all other points
 Weighted by distance
- Run power iteration (averaging) for 10 steps
- Plot node id x vs v¹⁰(x)
 - nodes are ordered and colored by actual cluster number









PIC: Power Iteration Clustering run power iteration (repeated averaging w/ neighbors) with early stopping

- 1. Pick an initial vector \mathbf{v}^0 .
- 2. Set $\mathbf{v^{t+1}} \leftarrow \frac{W\mathbf{v^t}}{\|W\mathbf{v^t}\|_1}$ and $\delta^{t+1} \leftarrow |\mathbf{v^{t+1}} \mathbf{v^t}|$.
- 3. Increment t and repeat above step until $|\delta^t \delta^{t-1}| \simeq 0$.
- 4. Use k-means to cluster points on v^t and return clusters $C_1, C_2, ..., C_k$.
- V⁰: random start, or "degree matrix" D, or ...
- Easy to implement and efficient
- Very easily parallelized
- Experimentally, often better than traditional spectral methods
- Surprising since the embedded space is 1-dimensional!

Experiments

- "Network" problems: natural graph structure
 - PolBooks: 105 political books, 3 classes, linked by copurchaser
 - UMBCBlog: 404 political blogs, 2 classes, blogroll links
 - AGBlog: 1222 political blogs, 2 classes, blogroll links
- "Manifold" problems: cosine distance between classification instances
 - Iris: 150 flowers, 3 classes
 - PenDigits01,17: 200 handwritten digits, 2 classes (0-1 or 1-7)
 - 20ngA: 200 docs, misc.forsale vs soc.religion.christian
 - 20ngB: 400 docs, misc.forsale vs soc.religion.christian
 - 20ngC: 20ngB + 200 docs from talk.politics.guns
 - 20ngD: 20ngC + 200 docs from rec.sport.baseball

Experimental results: best-case assignment of class labels to clusters

			NO	Cut	N.	IW	PIC		
	Dataset	k	Accuracy	Macro-F1	Accuracy	Macro-F1	Accuracy	Macro-F1	
	Iris	3	0.673	0.570	0.807	0.806	0.980	0.980	
	PenDigits01	2	1.000	1.000	1.000	1.000	1.000	1.000	
	PenDigits17	2	0.755	0.753	0.755	0.754	0.755	0.753	
	UBMCBlog	2	0.953	0.953	0.953	0.953	0.948	0.948	
	AGBlog	2	0.520	0.342	0.520	0.342	0.957	0.957	
	20ngA	2	0.955	0.955	0.955	0.955	0.960	0.960	
	20ngB	2	0.505	0.344	0.550	0.436	0.905	0.904	
	20ngC	3	0.613	0.621	0.635	0.639	0.737	0.730	
_	20ngD	4	0.469	0.432	0.535	0.534	0.580	0.570	
	Average	-	0.716	0.663	0.746	0.713	0.869	0.867	

Table 1: Clustering performance of PIC and spectral clustering algorithms on several real datasets.





Experiments: run time and scalability

		NCut	NJW	P	IC	
Dataset	Size	Runtime	Runtime	Runtime	Iterations	
Iris	150	589	242	59	6	
PenDigits01	200	965	326	56	6	
PenDigits17	200	1197	528	62	6	
UBMCBlog	404	4205	1589	85	21	
AGBlog	1222	114821	58145	211	34	
20ngA	200	1113	355	72	15	
20ngB	400	4085	1864	139	13	
20ngC	600	13070	6383	190	13	
20ngD	800	33191	16295	278	11	

Time in millisec

More experiments



Varying the number of clusters for PIC and PIC4 (starts with random 4-d point rather than a random 1-d point).

More experiments



Varying the amount of noise for PIC and PIC4 (starts with random 4-d point rather than a random 1-d point).

More experiments

Table 1: Dataset Statistics (N/E/C indicates Nodes / Edges / Clusters)(a) Social network(b) Author disambiguation

Dataset	N/E/C	Dataset	N/E/C	Dataset	N/E/C	Dataset	N/E/C
karate	34 / 156 / 2	umbc	404 / 4764 / 2	jsmith	4120 / 21452 / 30	jrobinson	686 / 2846 / 12
polbooks	105 / 882 / 3	mgemail	280 / 1344 / 55	akumar	801 / 2476 / 14	ktanaka	827 / 2758 / 10
dolphin	62/318/2	citeseer	2114 / 7396 / 6	cchen	424 / 1558 / 16	mbrown	579 / 2112 / 13
football	115 / 1226 / 10	cora	2485 / 10138 / 7	djohnson	1381 / 5344 / 15	mmiller	2106/9918/12
msp	4324 / 37254 / 2			jmartin	424 / 1558 / 16	jlee	5820/23110/100
ag	1222 / 33428 / 2			agupta	2485 /10208 / 26	ychen	5472/25584/71
senate	98 / 9506 / 2			mjones	961 / 3450 / 13	slee	5963 / 23086 / 86

More "real" network datasets from various domains

(c) Best alignment: Social networks

Dataset	PSK	PIC _D	PIC _R	PIC _{R4}	NCut	NJW
Karate	1.00	0.91	0.93	0.95	0.95	0.95
Dolphin	0.90	0.98	0.98	0.98	0.98	0.98
UMBC	0.95	0.93	0.95	0.95	0.95	0.96
AG	0.95	0.91	0.94	0.94	0.52	0.51
MSP	0.88	0.63	0.63	0.63	0.63	0.64
Senate	0.98	0.99	0.99	0.99	0.99	0.99
PolBook	0.78	0.80	0.81	0.83	0.82	0.80
Football	0.76	0.47	0.51	0.66	0.72	0.67
MGEmail	0.28	0.39	0.40	0.64	0.59	0.56
CiteSeer	0.33	0.51	0.48	0.55	0.48	0.52
Cora	0.47	0.46	0.40	0.45	0.29	0.42
Average	0.75	0.73	0.73	0.78	0.72	0.73

(d) Best alignment: Author disambiguation

Dataset	PSK	PIC _D	PIC _R	PIC _{R4}	NCut	NJW
AGupta	0.13	0.26	0.24	0.37	0.26	0.34
AKumar	0.20	0.29	0.31	0.37	0.35	0.40
CChen	0.30	0.43	0.44	0.53	0.24	0.50
DJohnson	0.15	0.24	0.33	0.46	0.47	0.35
JLee	0.11	0.20	0.23	0.41	0.17	0.39
JMartin	0.28	0.42	0.43	0.53	0.25	0.49
JRobinson	0.26	0.37	0.42	0.49	0.26	0.48
JSmith	0.11	0.22	0.21	0.41	0.31	0.42
KTanaka	0.19	0.36	0.41	0.45	0.45	0.43
MBrown	0.21	0.35	0.41	0.52	0.47	0.50
MJones	0.19	0.29	0.34	0.38	0.38	0.35
MMiller	0.14	0.30	0.41	0.52	0.52	0.53
SLee	0.08	0.19	0.23	0.41	0.23	0.39
YChen	0.10	0.23	0.28	0.47	0.23	0.46
Average	0.18	0.30	0.34	0.45	0.33	0.43

LEARNING ON GRAPHS FOR NON-GRAPH DATASETS

Why I'm talking about graphs

- Lots of large data *is* graphs
 - Facebook, Twitter, citation data, and other *social* networks
 - The web, the blogosphere, the semantic web, Freebase, Wikipedia, Twitter, and other *information* networks
 - Text corpora (like RCV1), large datasets with discrete feature values, and other *bipartite* networks
 - nodes = documents or words
 - links connect document \rightarrow word or word \rightarrow document
 - Computer networks, biological networks (proteins, ecosystems, brains, ...), ...
 - Heterogeneous networks with multiple types of nodes
 - people, groups, documents

Simplest Case: Bi-partite Graphs



Outline

- Background on spectral clustering
- "Power Iteration Clustering"
 - Motivation
 - Experimental results
- Analysis: PIC vs spectral methods
- PIC for sparse bipartite graphs
 - "Lazy" Distance Computation
 - "Lazy" Normalization
 - Experimental Results

Motivation: Experimental Datasets are...

- "Network" problems: natural graph structure
 - PolBooks: 105 political books, 3 classes, linked by copurchaser
 - UMBCBlog: 404 political blogs, 2 classes, blogroll links
 - AGBlog: 1222 political blogs, 2 classes, blogroll links
 - Also: Zachary's karate club, citation networks, ...
- "Manifold" problems: cosine distance between <u>all pairs</u> of classification instances
 Gets expensive fast
 - Iris: 150 flowers, 3 classes
 - PenDigits01,17: 200 handwritten digits, 2 classes (0-1 or 1-7)
 - 20ngA: 200 docs, misc.forsale vs soc.religion.christian
 - 20ngB: 400 docs, misc.forsale vs soc.religion.christian

— ...

Spectral Clustering: Graph = Matrix $A^*v_1 = v_2$ "propogates weights from neighbors"



Spectral Clustering: Graph = Matrix W*v₁ = v₂ "propogates weights from neighbors"



 $\mathbf{W} = \mathbf{D}^{-1} \mathbf{A}$

D[i,i]=1/degree(i)

Lazy computation of distances and normalizers

- Recall PIC's update is
 - $v^{t} = W * v^{t-1} = D^{-1}A * v^{t-1}$

1 is a column vector of 1's

- ...where D is the [diagonal] degree matrix: $D = A^*1$

 My favorite distance metric for text is lengthnormalized TFIDF:
 Vertication =
 Vertication =

- Def'n:
$$A(i,j) = \langle v_i, v_j \rangle / ||v_i||^* ||v_j||$$

- Let $N(i,i) = ||v_i||$... and N(i,j) = 0 for i!=j
- Let F(i,k)=TFIDF weight of word w_k in document v_i
- -Then: $A = N^{-1}F^{T}FN^{-1}$

Lazy computation of distances and normalizers

• Recall PIC's update is

 $- v^{t} = W * v^{t-1} = D^{-1}A * v^{t-1}$

Equivalent to using TFIDF/ cosine on all pairs of examples but requires only *sparse* matrices

- ...where D is the [diagonal] degree matrix: D=A*1
- Let F(i,k)=TFIDF weight of word w_k in document v_i
- Compute $N(i,i) = ||v_i|| \dots$ and N(i,j) = 0 for i!=j
- **Don't** compute $A = N^{-1}F^{T}FN^{-1}$
- Let $D(i,i) = N^{-1}F^{T}FN^{-1*1}$ where 1 is an all-1's vector
 - Computed as D=N⁻¹(F^T(F(N⁻¹*1))) for efficiency
- -New update:
 - $v^t = D^{-1}A * v^{t-1} = D^{-1}N^{-1}F^TFN^{-1}*v^{t-1}$

Experimental results

- RCV1 text classification dataset
 - 800k + newswire stories
 - Category labels from *industry* vocabulary
 - Took single-label documents and categories with at least 500 instances
 - Result: 193,844 documents, 103 categories
- Generated 100 random category pairs
 - Each is all documents from two categories
 - Range in size and difficulty
 - Pick category 1, with m_1 examples
 - Pick category 2 such that $0.5m_1 < m_2 < 2m_1$

Results

	ACC-Avg	NMI-Avg
baseline	57.59	-
k-means	69.43	0.2629
NCUTevd	77.55	0.3962
NCUTiram	61.63	0.0943
PIC	76.67	0.3818

NCUTevd: Ncut with exact eigenvectors
NCUTiram: Implicit restarted Arnoldi method
No stat. signif. diffs between NCUTevd and PIC







Runtime (sec)

- Linear run-time implies *constant* number of iterations
- Number of iterations to "accelerationconvergence" is hard to analyze:
 - Faster than a single complete run of power iteration to convergence
 - -On our datasets
 - 10-20 iterations is typical
 - 30-35 is exceptional



Size vs PIC Accuracy



From SemiSupervised to Unsupervised Learning ... and back again

- Implicit manifolds work for unsupervised learning (PIC)
- But PIC is *so close* to SSL methods

PIC: Power Iteration Clustering run power iteration (repeated averaging w/ neighbors) with early stopping

- 1. Pick an initial vector \mathbf{v}^0 .
- 2. Set $\mathbf{v^{t+1}} \leftarrow \frac{W\mathbf{v^t}}{\|W\mathbf{v^t}\|_1}$ and $\delta^{t+1} \leftarrow |\mathbf{v^{t+1}} \mathbf{v^t}|$.
- 3. Increment t and repeat above step until $|\delta^t \delta^{t-1}| \simeq 0$.
- 4. Use k-means to cluster points on \mathbf{v}^t and return clusters $C_1, C_2, ..., C_k$.

Harmonic Functions/CoEM/wvRN

- 1. Pick an initial vector \mathbf{v}^0 .
- 2. Set $\mathbf{v^{t+1}} \leftarrow \frac{W\mathbf{v^t}}{\|W\mathbf{v^t}\|_1}$ then replace $\mathbf{v^{t+1}}(i)$ with seed values +1/-1 for labeled data
- 3. Increment t and repeat above step for 5-10 iterations
- 4. Classify data using final values from **v**

Implicit Manifolds on the NELL datasets



Name	20NG	RCV1	City	44Cat
Instances	19K	194K	88K	9,846K
Features	61K	47K	99K	8,622K
NZF	2M	11M	21M	121M
Cats	20	103	1	44
Туре	doc	doc	NP	NP
Manifold	cosine	cosine	bipart	bipart
Input Size	39MB	198MB	330MB	2GB
IM Size	40MB	207MB	335MB	2.4GB
EM Size	5.6GB	*540GB	*80GB	*4TB

Table 1: Dataset comparison. *NZF* is the total number of nonzero feature values and *Cats* is the number of categories. *Type* is the dataset type, where *doc* and *NP* correspond to document collection and noun phrase-context data, respectively. *Manifold* is the choice of manifold for the dataset, where *cosine* and *bipart* refers to cosine similarity and bipartite graph walk, respectively. *Input Size* is the MATLAB memory requirement for the original sparse feature matrix; *IM Size* is the total memory requirement for using the implicit manifold, including the feature matrix; *EM Size* is the memory requirement for constructing a explicit manifold. * indicates that the memory requirement is estimated using random sampling and extrapolation.



Figure 1: F1 scores on the 20NG and RCV1 datasets. The xaxis indicates the number of labeled instances and the y-axis indicates the macro-averaged F1 score. Vertical lines indicate standard deviation (over 20 trials for 20NG and 10 for RCV1) using randomly selected seed labels.



Figure 2: F1 scores on the 20NG and RCV1 datasets using preferred (high feature weight sum) seeds. Subscript HFS indicates result using high feature-sum seeds and R indicates result using random seeds—included for comparison.

Method	SVM	HF	MRW	HF	MRW
Manifold	-	inner	inner	bipart	bipart
NDCG	0.0263	0.0402	0.0405	0.0406	0.0408
AP	0.0208	0.6728	0.7067	0.7130	0.7389
P@10%	0.0123	0.8732	0.8926	0.8796	0.9094
P@20%	0.0143	0.8698	0.8991	0.8941	0.9162
P@30%	0.0168	0.8773	0.9093	0.9036	0.9116
P@40%	0.0199	0.8574	0.8957	0.9118	0.9179
P@50%	0.0210	0.8227	0.8647	0.8832	0.9038
P@60%	0.0236	0.7591	0.7990	0.8093	0.8307
P@70%	0.0265	0.6337	0.6743	0.6805	0.7189
P@80%	0.0267	0.4131	0.4533	0.5087	0.5297
P@90%	0.0272	0.1927	0.2155	0.2521	0.2926
P@100%	0.0274	0.0275	0.0279	0.0280	0.0289

Table 2: City dataset result. Boldfaced font indicates the highest number in a row. *inner* refers to the inner product manifold and *bipart* refers to the bipartite graph walk manifold. Note that HF with bipart is equivalent to co-EM as used in [11]



Figure 4: Sampled per-category accuracies of the top 1000 retrieved NPs on the 44Cat dataset. The categories are ordered from left to right according to the difference between the MRW accuracy and HF accuracy, from the high to low.

A smoothing trick:

$$V^{t+1} \leftarrow (1 - \alpha - \beta)SD^{-1}V^t + \alpha R + \beta(\mathbf{1}/n)$$



Figure 3: Parameter sensitivity. The x-axis correspond to parameter values and the y-axis shows average precisions. α ranges from 0.05 to 0.65, β ranges from 0.0001 to 0.01; the number of iterations T are indicated below x-axes.