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1 Abstract

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The project is currently on track – in the upcoming quarter, we will continue applying the developed algorithms to various data sets and the design/implementation of the multiscale heat kernel coordinates algorithms. No problems are currently anticipated.

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2 Summary

In this quarter, we continued design and implementation of the new multiscale heat kernel coordinates (MHKC) algorithms. The current design variants for MHKC algorithms are documented in this report.

The project is currently on track – in the upcoming quarters, we will continue applying the developed algorithms to various data sets and focus on the design and development of the MHKC algorithms. No problems are currently anticipated.

3 Introduction

The primary project effort over the last quarter focused on completing design/development of the multiscale heat-kernel coordinates algorithms [1]. This provides a power tool for discovering the non-linear geometries in any given dataset. This utilizes the fast randomized Singular Value Decomposition (RSVD) algorithms described in the earlier ONR reports [7][8]. Use of the RSVD effectively reduces the computational complexity from $O(m.n.k)$ to $O((m+n).k^2)$ for an m by n matrix of rank k . In contrast to the multiscale Singular Value Decomposition (MSVD) algorithms that detect linear structures in data at multiple scales, the MHKC uses heat kernels to discover the non-linear manifold structure in which the data resides at various scales. Similar to the MSVD, the MHKC provides an efficient representation using low-dimensional coordinates corresponding to the original data points.

An outline of the MHKC algorithm was presented in the previous quarterly report [10]. While most of the algorithm is automated, the crucial step of selecting the *appropriate* heat-kernel coordinates for any given application required manual intervention on part of the data analyst. In this report, we present two canonical approaches to automating the selection of the MHKC embedding. Further, it also provides a way to visualize the embedded data in lower (2 or 3) dimensions.

4 Methods, Assumptions and Procedures

4.1 Multiscale Heat Kernel Coordinates

The Multiscale Heat Kernel Coordinates (MHKC) algorithms are based on theoretical results presented in [1]. The current algorithm design is described below.

Input: A set of n data points $\{x_1, x_2, \dots, x_n\}$ in R^d . Assume n is large.

Step 1 (Normalization): Normalize the points x_i such that the data cloud is in a ball of unit variance. Define

$$\sigma_x = \sqrt{\frac{1}{n} \sum_{i=1}^n \|x_i - \vec{x}\|^2}$$

where $\vec{x} = \frac{1}{n} \sum_{i=1}^n x_i$. The normalized point y_i corresponding to x_i is given by

$$y_i = \frac{x_i - \vec{x}}{\sigma_x}$$

Note: The translation to mean zero is not necessary for the purposes of building the transition probability matrix in the next step.

Step 2 (Transition Probability Matrix): The second step comprises constructing the data matrix to be provided as input to the RSVD algorithm. Define the heat kernel as

$$k(x, y) = \exp\left(-\frac{\|x - y\|_2^2}{t_0}\right)$$

for any two points x and y . Here, t_0 is a constant (data dependent) representing the kernel window size (set $t_0 = 2^{-s} \sigma_y^2$ for scale $s \geq 0$; select s representing some finer scale of interest). The heat kernel matrix is then defined as

$$K = \{k_{ij}\} \text{ where } k_{ij} = k(x_i, x_j)$$

for $i, j = 1, 2, \dots, n$. The transition probability matrix is $P = D^{-1}K$ where D is the diagonal matrix with the i -th entry as sum of the i -th row of K .

Note: For large n , compute $\beta \approx 25$ elements for each row of K using the randomized approximate nearest neighbor algorithm ([9]). This reduces the computational complexity from $O(n^2 \cdot d)$ to $O(n \cdot \log(n) \cdot d)$ and captures local information.

Note that P is not symmetric. There are various techniques to symmetrize P such that the eigenvalues and eigenfunctions are still easy to compute. One way is to define

$$P' = D^{-1/2} \cdot P \cdot D^{-1/2}$$

P' is symmetric with the same eigenvalues as P . Also, the eigenvectors can easily be easily obtained using a simple transformation of either $D^{-1/2}$ or $D^{1/2}$. The RSVD algorithm may be used to compute the spectrum of P' .

Step 3 (MHKC Embedding): Next, the heat kernel coordinates is defined for each of the original data points. Let the eigenvalues of P be defined as λ_j and the right-eigenvectors as v_j for $j = 1, 2, \dots, \text{rank}(P)$.

Each point x_i is then represented as $HKC(x_i) = (\exp(-\lambda_1 t) \cdot v_{1i}, \exp(-\lambda_2 t) \cdot v_{2i}, \dots, \exp(-\lambda_r t) \cdot v_{ri})$ where v_{ji} is the i -th coordinate of the eigenvector v_j . Here t is the time/scale parameter that is to be varied to look at the geometries of the data set at various scales.

Note: The first eigenvalue/eigenvector of P is trivial and should not be used.

Next, we provide two approaches to automating the selection of heat-kernel coordinates in Step 2 of the algorithm described above.

4.1.1 Representation Using Canonical Clustering

Choose a small integer l representing the number of non-trivial eigenvectors v_j to consider (obtained via the algorithm in Section 4.1). Construct a binary tree using the first l non-trivial eigenvectors to divide the n points into $N_c = 2^l$ clusters as follows. First, divide the points into two sets defined by

$$\left\{ x_i \mid y_i^1 \geq \frac{1}{n} \sum_{j=1}^n y_j^1 \right\} \text{ and } \left\{ x_i \mid y_i^1 < \frac{1}{n} \sum_{j=1}^n y_j^1 \right\}$$

where $y_i^k = v_k(x_i)$ using the first diffusion vector. Apply the 2^{nd} diffusion vector to both these sets; repeat recursively to obtain N_c disjoint clusters. For each cluster $c = 1, 2, \dots, N_c$, find the closest point $v(x_{\mu(c)})$ to the cluster mean.

Select a suitable time t (same for all the clusters) and use the heat kernel $k(x, y)$ to get a vector of length N_c given by

$$z_i = \left(k(x_i, x_{\mu(1)}), k(x_i, x_{\mu(2)}), \dots, k(x_i, x_{\mu(N_c)}) \right)$$

for each x_i . Now, perform standard PCA on the dataset $\{z_i\}$. To visualize the dataset, simply use projections on the first 2 or 3 dimensions.

4.1.2 Representation Using Multiscale SVD

For each k , normalize $(y_1^k, y_2^k, \dots, y_n^k)$ to have unit length. Define $z_i^k = y_i^k / \sum_{j=1}^n y_j^k$. Pick a small integer, say 2. Out of the top ten non-trivial diffusion eigenvectors, choose the two values k_1, k_2 such that the set (in 2D) consisting of all the points

$$\{(z_i^{k_1}, z_i^{k_2}) \mid i = 1, 2, \dots, n\}$$

has the smallest average value of multiscale SVD at scales 2^{-s} for $s = 0, 1, 2, 3$ (at each location and scale, compute the average squared distance to the best fitting line). Amongst the various “best” choices, pick the one with the smallest value of $k_1 + k_2$.

4.2 Deliverables / Milestones

| Date | Deliverables / Milestones | Status |
|----------|--|--------|
| Oct 2010 | Progress report for period 1, 1 st quarter | ✓ |
| Jan 2011 | Progress report for period 1, 2 nd quarter / complete randomized matrix decompositions task | ✓ |
| Apr 2011 | Progress report for period 1, 3 rd quarter / complete approximate nearest neighbors task | ✓ |
| Jul 2011 | Progress report for period 1, 4 th quarter / complete experiments – part 1 | ✓ |
| Oct 2011 | Progress report for period 2, 1 st quarter | ✓ |
| Jan 2012 | Progress report for period 2, 2 nd quarter / complete multiscale SVD task | ✓ |
| Apr 2012 | Progress report for period 2, 3 rd quarter | ✓ |
| Jul 2012 | Progress report for period 2, 4 th quarter / complete experiments – part 2 | ✓ |
| Oct 2012 | Progress report for period 3, 1 st quarter | ✓ |
| Jan 2013 | Progress report for period 3, 2 nd quarter / complete multiscale Heat Kernel task | ✓ |
| Apr 2013 | Progress report for period 3, 3 rd quarter | |
| Jul 2013 | Final project report + software + documentation on CDROM / complete experiments – part 3 | |

5 Results and Discussion

We described two approaches to automate the process of selecting the “appropriate” diffusion vectors for a given dataset. The first approach in itself provides an agnostic and canonical way of “clustering”. In terms of computational cost, the first approach is much better as it avoids the potential combinatorial explosion in the second approach. However, the second approach directly evaluates the information content for each diffusion vector in a multiscale sense and picks the “best” combination. We will experimentally evaluate both these techniques against real-world datasets.

6 Conclusions

The project is on track with design/implementation of the new multiscale heat kernel coordinates algorithms. We will continue with algorithmic improvements and experimentation using the developed algorithms in the next quarter.

No problems are currently anticipated.

7 References

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