Clustering Metagenomic Sequences Using Canopies

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Abstract—Metagenomics is the study of genome sequences from samples which are retrieved directly from environment hosting multiple organisms that co-exist as communities within ecological regions. Metagenomics study has created the necessity and challenges of developing computational tools for the quantification of abundance, diversity, role of different species within communities, phenotype inferences etc. Enormous amount of data is created during metagenome sequencing process. Algorithms have been developed to cluster similar metagenome sequences. We have developed an approach for clustering metagenomic data that uses Canopy Clustering with Locality Sensitive Hashing distance approximation to make clustering process in metagenomic data faster. Canopy Clustering works as a preprocessing step to reduce pairwise distance calculation and enables efficient parallel processing with subsequent expensive cluster methods while LSH provides fast distance approximation and reduces data dimension. We tested our framework with three popular clustering mechanisms in literature on three synthetic and three real world large scale metagenome datasets and observed that our proposed approach can reduce runtime while providing similar in some cases better outcomes.

Keywords-Clustering, Canopy, LSH, Metagenome

I. Introduction

Metagenomics is the study of genetic material recovered directly from samples that comprise organisms as co-existing communities. These samples can be taken from sea, soil and human body [1][2]. Metagenomics has enabled scientists to study all of the genomes in a community as a whole. Analysis of microbial community through metagenome data can reveal interesting relationships between microbial community and the host which in turn can lead to further investigation. For example analyzing metagenomic data from human gut microbiome provides an understanding of the role played by microbes with regards to human health and disease [3].

Instead of producing whole genomic sequences of the members of communities in host samples, latest sequencing technologies produce short contiguous subsequences called *reads* from random positions of actual whole genome. These reads from different organisms are commingled together posing fundamental challenge to further analysis of the data. Combining the reads of different organisms based on overlapping yet discriminating information from organism specific genome sequences is known as sequence assembly

[4]. Many Sequence assemblers require reference genome sequence and the process of assembly is exceedingly complex, challenging and time consuming [5]. For this reason clustering metagenome data for identification of Operational Taxonomic Units (OTU) from 16s rRNA genes has become popular recently. 16S rRNA gene sequencing has been widely used for the analysis of genetic diversity within complex microbial communities. 16S sequences are marker genes, which exists in most microbial species but have variations in their sequences that allow them to be separated into different taxonomic groups [6]. OTU is used to classify groups of closely related individuals from similar or different taxonomic levels [7]. It is the most commonly used microbial diversity unit, especially when analyzing the small subunit 16S or 18S rRNA marker gene microbial datasets [8]. Sequences can be clustered according to their similarity to each other. Microbial OTUs are generally ecologically consistent across the hosts regardless of OTU clustering approaches [9].

Clustering approaches have been developed and used for the rapid analysis of large sets of whole and targeted 16S rRNA metagenomic sequences which are discussed in Section II - Literature Review. Analysis of microbiome datasets typically begins by clustering raw sequence reads and creating potential OTUs. Proper clustering of sequence reads assists in the metagenome assembly problem, allows computation of different species diversity metrics and allows further analysis with phenotype inferences. In this study we have proposed and evaluated a pre-clustering technique based on Canopy cluster method and Locality Sensitive Hashing. Our proposed framework can reduce pairwise comparison between sequences for similarity measure in large scale metagenome datasets by partitioning the dataset with fast LSH based approximation. These initial partitions can be considered independent of each other. More accurate and expensive clustering methods can be deployed for each partition in parallel utilizing multi-core architechture of modern CPUs. Only the sequences inside a canopy will be considered for further sub-clustering not the sequences outside that canopy. This characteristic of Canopy clustering reduces expensive pairwise comparison significantly.

II. LITERATURE REVIEW

Over the years several sequence clustering methods have been developed and used widely for metagenomic sequence data. A comprehensive survey by XXX et. al. [?] benhcmarks these clustering algorithms including CD-HIT [10], DOTUR [11], MOTHUR [12], UCLUST [?] and ESPRIT [13].

CD-HIT [?] is general purpose sequence clustering algorithm that follows an incremental, greedy approach. CD-HIT uses pairwise sequence alignment to find similar sequences. UCLUST [?] is similar to CD-HIT but achieves a signficant speedup over CD-HIT by using seeds (fixed length gapless subsequences) for performing pairwise sequence comparisons. MC-LSH [14] utilizes an efficient locality sensitive based hashing function to approximate the pairwise sequence similarity. Pairwise similarity among sequences is computed based on randomly chosen indices that essentially compresses the input sequences. The MC-MinH [15] algorithm uses the min-wise [16] hashing along with the greedy clustering algorithm to group 16S and whole metagenomic sequences. Mash [17], uses MinHash locality sensitive hashing to reduce large sequences to a representative sketch and rapidly estimate pairwise distances. Other methods for clustering whole metagenome sequence reads include TOSS [18], AbundanceBin [19] and CompostBin [20]. All unique k-mers are first clustered in TOSS and then clusters are merged based on k-mer repetitions. In AbundanceBin reads are modeled as mixture of Poisson distributions. Then Expectation Maximization (EM) algorithm is used to infer model parameters for the final clustering. Principal component analysis is used within CompostBin to project the data into a lower dimensional space and followed with a graph partitioning approach

DOTUR [?] and MOTHUR [?] are use a pairwise distance matrix as input and perform hierarchical clustering. The pairwise distances are computed based on an expensive sequence alignment between all pairs of input sequences.

UCLUST [?], SWARM [23][24] and SUMACLUST [25] are considered to be the state-of-the-art metagenomic sequence clustering methods by a recent benchmarking study [26]. SWARM uses an exhaustive single-linkage clustering approach using a fast pairwise sequence alignment. SUMACLUST is similar to UCLUST. Abundance ordered list of input sequences are compared against the representative set of already chosen sequences in SUMACLUST.

In this study introduced a fast initial partitioning of large scale metagenomic datasets using canopies [27] and locality sensitive hashing [28]. This way efficient and scalable parallelism in OTU-clustering can be achieved that is capable of making existing OTU-clustering method multiple times faster by taking advantage of modern multicore CPU architectures. We have used UCLUST, SWARM and SUMACLUST with and without our proposed framework

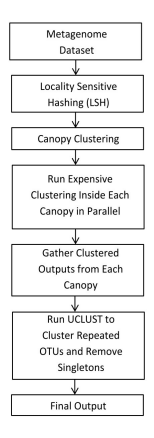


Figure 1. Overview of our proposed Canopy Clustering approach on Metagenome Data

and brief descriptions of these methods are provided in Section III-D.

III. METHODS

A. Overview

In this section we provide an overview of our proposed framework. Figure 1 represents an abstract overview of our proposed approach.

Figure 2 shows a more detailed overview of our proposed Canopy clustering approach for metagenomic data. After computing the kmers and LSH codes our proposed method will start canopy clustering based on the hamming distances between the binary representations of data points created by LSH. Once all reads in datasets are assigned to canopies, our method will create multiple partitions from original dataset based on canopy membership information. Finally each smaller partitions representing a canopy will be provided as input to more accurate sequence clustering methods in parallel. In this study UCLUST, SUMACLUST, SWARM and Minimum Spanning Tree (MST) based clustering will be used as accurate clustering methods for canopies which are briefly discussed in Section III-D.

The final step of our proposed framework is to merge the clustered outputs from each canopy provided by more

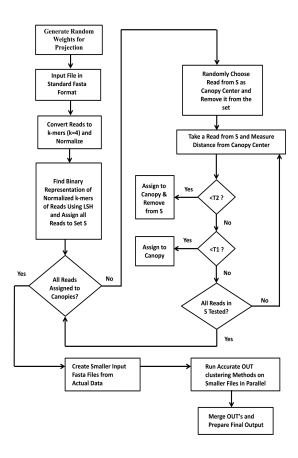


Figure 2. Workflow of Canopy Clustering for Large Scale Metagenome Data

accurate and expensive clustering methods. There is a potential concern in this scenario. According to Canopy cluster algorithm a single data point may belong to multiple canopies as long as the soft threshold is met. As a result same or very similar OTUs will be generated from multiple canopies. This scenario is depicted in Figure 3. The dashed circles titled OTU-A, B, C and D represent *potential OTUs*. Soild outer circles represent canopies titled Canopy-1, 2 and 3. OTU-A and C are shared by Canopy-1 and 2. OTU-B and D are shared by Canopy-2 and 3. The smaller triangles, stars, hexagons and rectangles represent sequence reads in metagenome data. Because of this repetitions another final clustering on resulting OTUs is required which will remove repetitions in OTU. We are proposing two solutions to this scenario.

1) Merging with Greedy Clustering Algorithm: We have chosen UCLUST for merging step since UCLUST is one of the fastest greedy sequence clustering methods. We chose 90% identity threshold for UCLUST in this step. Other sequence clustering algorithm can also be applied in this step. Since number of sequences is already reduced by

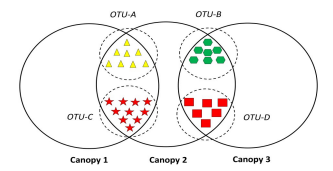


Figure 3. Same OTUs Occurring in Multiple Canopies

Canopy and accurate clustering, this merging step is very fast. Greedy clustering methods like UCLUST always converges to local minima. Thus this approach does not require number of OTUs as prior.

2) Merging with Minimum Spanning Tree: We developed a Minimum Spanning Tree and Locality Sensitive Hashing (LSH) distance based metagenome sequence clustering method to be used in merging step as well as inside Canopies for accurate clustering. A brief description of MST based clustering is given in Section III-D. Unlike merging with greedy clustering methods, this approach requires number of edges to remove as prior which indicates the number of OTUs.

In next sections we provide a brief description of Canopy Clustering, Locality Sensitive Hashing, their potential in metagenome data clustering, sub-clustering methods used in this study and our motivation for using them.

B. Canopy Clustering

Canopy Clustering [27] is an efficient approximate clustering algorithm often used as preprocessing step for other accurate and expensive clustering methods like the K-means algorithm or the Hierarchical clustering algorithm. It is intended to speed up clustering operations on large data sets, where standard clustering algorithms may be impractical in terms of run time and memory consumptions due to the large number of pairwise distance calculations. For a dataset with N instances, worst case calculations without canopy clustering is N^2 . After canopy clustering if the number of canopies is k then worst case calculations with canopy clustering is $\sum_{i=1}^k (c_i)^2$ where c_i is the number of instances in ith canopy. The total difference g in pairwise distance calculations:

$$g = N^2 - \sum_{i=1}^{k} (c_i)^2 \tag{1}$$

Canopy clustering uses two basic distance thresholds namely *soft* threshold T1 and *tight* threshold T2. If data point p_1 is within the soft distance threshold T1 with another

data point p_2 then p_1 will reside in same canopy as p_2 but p_1 may belong to other canopies too assuming that it has only met soft threshold and best match is yet to be found. Thus one data point may belong to multiple canopies in Canopy clustering. On the other hand if data point p_1 is within the tight distance threshold T2 with another data point p_2 then canopy clustering assigns p_1 to the same canopy as p_2 and stops assigning p_1 to any other canopy assuming that tight threshold has been met and best canopy assignment for p_1 has been found. In this case p_1 will not be repeated in other canopies. Canopy centroids are selected randomly. Then all other data points are tested for canopy assignment based on T1 and T2. New canopies are selected until all data points are assigned to canopies. A fast approximation based on Locality Sensitive Hashing (LSH) will be used for Canopy membership identification.

C. Locality Sensitive Hashing

A Locality Sensitive Hashing (LSH) [28][29][30] is a distribution on a family \mathcal{F} is defined for a metric space $\mathcal{M}=(M,d)$ a threshold R>0 and an approximation factor c>1. This family \mathcal{F} is a family of functions $h:\mathcal{M}\to S$ which map elements from the metric space to a bucket $s\in S$. The LSH family either satisfies h(p)=h(q) with probability at least P_1 if $d(p,q)\leq R$ or satisfies h(p)=h(q) with probability at most P_2 if $d(p,q)\geq R$ for any two points $p,q\in \mathcal{M}$, using a function $h\in \mathcal{F}$ which is chosen randomly. A family is interesting when $P_1>P_2$. Such a family \mathcal{F} is called (R,cR,P_1,P_2) -sensitive. LSH has been used for fast comparison between points in very high dimensional space [30]. To preserve approximate cosine distance, data points are often randomly projected to low dimensional bit signatures [34].

In this study we have chosen a random projection based hashing function that projects a d dimensional data point into a n dimensional bit representation where n < d. The intuition behind choosing n < d is to reduce dimension of data. Given a random projection v' of size $1 \times d$ and a data vector a of size $1 \times d$, the dot product between them provides a scalar value. Sign of this scalar value represents which side of the random hyperplane does the data point exist. This information can be represented with a bit. This way a single data point can be represented with limited number of bits. Hamming distance between these binary representations indicates disagreements between two data points.

D. Sub-Clustering Inside Canopies

We have used three recent and popular sequence clustering methods as the expensive clustering measure inside canopies in this study. UCLUST [22], SUMACLUST [25] and SWARM [23] were used for sub-clustering canopies.

1) UCLUST: It is a closed-source software and provides very limited documentations¹. The 32 bit version of

UCLUST executable is available for academic usage. But the 64-bit versions which is necessary to handle large datasets, require expensive license. UCLUST follows a greedy process and creates *seeds* of sequences which generate clusters based on percent identity.

- 2) SUMACLUST: An open-source software. It follows similar approach as UCLUST. Based on greedy strategy SUMACLUST incrementally constructs clusters by comparing an abundance-ordered list of input sequences against the representative set of already-chosen sequences. Initially this list is empty.
- 3) SWARM: SWARM uses exhaustive single-linkage clustering based on optimal sequence alignment. Sequences that are less than a certain distance from any other other sequence in the cluster are clustered together. SWARM attempts to reduce the impact of clustering parameters on the resulting OTUs by avoiding arbitrary global clustering thresholds and input sequence ordering dependence. At first SWARM builds an initial set of OTUs is constructed by iteratively agglomerating similar amplicons. Then amplicon abundance values are used to reveal OTUs internal structures and to break them into sub-OTUs.
- 4) Minimum Spanning Tree (MST): To evaluate Canopy as a standalone clustering method along with UCLUST, SUMACLUST and SWARM, we used Minimum Spanning Tree (MST) based clustering method as an accurate and expensive clustering inside canopies. Minimum Spanning Tree (MST) produces a connected spanning tree of undirected graph. It connects all vertices together such that the total weight of edges remains minimum. This property has been used for clustering [31][32][33]. We represented edges with Locality Sensitive Hashing based distances. Then edges with highest distances were removed from MST to generate OTUs.

In the following Sections UCLUST, SUMACLUST, SWARM and MST with Canopy clustering approach are represented as CC_{UCLUST} , $CC_{SUMACLUST}$, CC_{SWARM} and CC_{MST} respectively where the term CC stands for Canopy Clustering.

IV. EXPERIMENTAL EVALUATION

A. Dataset Description

To evaluate the performance of our developed approach we have used previously published synthetic and real world sequence benchmarks. Specifically, we used three synthetic 16S rRNA gene mock community datasets (Bokulich₂, Bokulich₃, and Bokulich₆) from Bokulich et. al. [35] and three real data sets: a 16S rRNA gene soil data set (Canadian Soil) [36], a 16S rRNA gene human data set (Body Sites) [37] and 18S rRNA gene soil data set (Global Soil) [38]. Key statistics and relevant information regarding these datasets are presented in Table I.

Synthetic Datasets:

¹http://www.drive5.com/uclust/uclust_userguide_1_1_579.pdf

Table I DATASET STATISTICS

Datasets	Type	Reference	# of Reads	# of Samples	Read Length	Platform
Bokulich ₂	Mock	[35]	6,938,836	4	189-251	HiSeq
Bokulich ₃	Mock	[35]	3,594,237	4	114-151	HiSeq
Bokulich ₆	Mock	[35]	250,903	1	114-150	HiSeq
Canadian Soil	Genuine	[36]	2,966,053	13	76–10	HiSeq
Body Sites	Genuine	[37]	886,630	602	117-351	GS FLX
Global Soil	Genuine	[38]	9,252,764	57	119-151	HiSeq

- 1) Bokulich₂: This dataset was prepared using the Illumina TruSeq v2 paired—end library preparation kit. It is a simulated 16S rRNA gene microbial community data set. This dataset contains 19 taxonomic Families, 19 Genera, 22 Species and 22 Strains in total. This dataset can be found in the QIIME database (identifier 1685).
- 2) Bokulich₃: Similar to Bokulich₂ except that it was prepared with the TruSeq v1 paired-end library kit at Illumina Cambridge and is also available in the QIIME database (identifier 1686).
- 3) Bokulich₆: This 16S rRNA dataset was sequenced at Washington University School of Medicine and contains evenly distributed microbial communities. This dataset contains 13 taxonomic Families, 23 Genera, 44 Species and 48 Strains in total. All these datasets from Bokulich et al.[35] are available at QIIME database² under their respective ID's. Since, these are simulated datasets the taxonomic profile of microbial organisms within them are known.

Real World Datasets:

- *4) Canadian Soil:* The Canadian Soil dataset³ contains genomic data of soil spanning from Arctic Tundra to Agricultural soil suitable for different agricultural products.
- 5) Body Sites: This dataset contains composition of bacterial communities from up to 27 different body sites in healthy adults. A collection of 602 samples acquired from different body sites of human subjects are provided with meta-data.
- 6) Global Soil: The global soil data was taken from Ramirez et al. [38] which is a study of the below-ground diversity in New York City's Central Park.

All of these datasets have been used in previous studies [35],[39]. Any analysis on these datasets requires appropriate preprocessing which can significantly change the results of clustering and taxonomic classification based on them [35]. The performance of different open source sequence clustering methods were assessed and compared in a study by Kopylova et. al. [39] using these specific datasets. As such, in this paper we use the same benchmarks as done in the prior study and publicly available ⁴.

B. Evaluation Metrics

We assess the performance of our proposed approach using the following types of commonly used metrics: (i) that are used for the assessment of biodiversity within metagenomic samples, (ii) output of clustering algorithms and (iii) computational run time.

- 1) Faiths phylogenetic diversity metric (PD): Faiths phylogenetic diversity [40] is based on the Phylogenetic tree. It combines all the branch lengths of the tree as a measure of diversity. So, if a new OTU (cluster) is found and it is closely related to another OTU in the sample, it will contribute to a small increase to the PD score. However, if a new OTU is found and is from a different lineage than anything else in the sample, it will contribute to a large increase in the PD score.
- 2) Shannon Entropy: Shannon-Wiener diversity index is defined as:

$$H = -\sum_{i=1}^{s} (p_i \log_2 p_i)$$
 (2)

where s is the number of OTUs and p_i is the proportion of the community represented by OTU i. Since by definition p_i will be between zero and one, the natural log makes all of the terms of the summation negative, which is why we take the inverse of the sum. The Shannon index increases as both the richness and evenness of the community increase. The fact that the index incorporates both components of biodiversity can be seen as both a strength and a weakness. It is a strength because it provides a simple summary, but it is a weakness because it makes it difficult to compare communities that differ greatly in richness.

3) Simpson's Index: Simpsons index is defined as 1-dominance or

$$1 - \sum p_i^2 \tag{3}$$

where where p_i is the proportion of the community represented by OTU i. Simpsons index is based on the probability of any two individuals drawn at random from an infinitely large community belonging to the same species. It measures *evenness* of the community from 0 to 1. Higher value of this index refers higher similarity and relatively lower diversity of microorganisms.

²http://giime.org/home_static/dataFiles.html

³http://www.cm2bl.org/

⁴https://github.com/ekopylova/otu-clustering

4) F-measure: In case of synthetic datasets, false-positive (FP; taxonomy/OTU string exists in observed but not expected), false-negative (FN; taxonomy/OTU string exists in expected but not observed), and true-positive (TP; taxonomy/OTU string exists in both observed and expected) measures were computed from cluster output and the ground truth which is the expected taxonomic composition. The following definitions were used:

$$precision = \frac{TP}{(TP + FP)} \tag{4}$$

$$recall = \frac{TP}{(TP + FN)} \tag{5}$$

$$FScore = \frac{2 \times precision \times recall}{(precision + recall)} \tag{6}$$

5) Pearson Coefficient Correlation (ρ -value): After getting Operational Taxonomic Units (OTU) from a clustering method we created the taxonomic profile at Genus level. Pearsons correlation coefficient was computed to measure the relatedness of taxonomic assignment between a pair of tools. Values can range between -1 and 1, with -1 indicating a negative correlation, 0 indicating no correlation, and 1 indicating a positive correlation or strong relationship.

C. Experimental Details

For kmers, the value of parameter K was set to 4. There are 4 possible nucleotide representatives (A,C,G and T) and the total number of features became $4^4 = 256$. This way we were able to convert the string representation into numeric. For Locality Sensitive Hashing the number of Hyperplanes (parameter d) was set to 150. We wanted to reduce the number of features while maintaining a relatively accurate distance approximation. So value of d was chosen to be less than 256. For performance analysis and comparison we have chosen T1 = 0.45 and T2 = 0.34.

We performed all the experiments on computers with Intel 5th generation Core i7 2.70GHz 64bit processor with 8 core CPUs and 12GB memory. For implementation we used Python 2.7.12 and QIIME [41] version 1.9.0 - a popular open source Bioinformatics pipeline that combines many metagenome clustering methods including the ones which has been used in this work. One important aspect our proposed approach is parallelism. Each canopy can be clustered in parallel. To take full advantage of today's multicore CPU based computing systems we utilized Python's multiprocessing module instead of threading module. According to Python's documentation⁵ multiprocessing module allows the programmer to fully leverage multiple processors on a given machine by spawning subprocesses instead of threads. After receiving output from all canopy clustering we merged the outputs to make a single clustering results. Some performance metric used in this study like Faiths Phylogenetic Diversity (PD) metric require sequence alignment. We used PyNast⁶ [42] open source sequence aligner for aligning clustered output.

V. RESULTS DISCUSSION

A. Clustering Performance Comparison

Table II shows the performance of UCLUST [22], SUMACLUST [25] and SWARM [23] and their corresponding versions with our proposed approach. Table II compares F-scores and Pearson Correlation Coefficient (ρ). F-scores are only available for synthetic benchmarks since taxanomic profile for them is known as ground truth. But F-scores are not available for real benchmarks as no ground truth is available for them. Correlation values were generated based on taxonomic profiles at Genus level generated from outputs from clustering methods. From Table II we can see that F-scores obtained from a clustering method and its corresponding Canopy clustering version are very closer, in some cases better. We got higher F-score for Bokulich₂ benchmark from CC_{SUMACLUST}. CC_{UCLUST} and CCSWARM provided same F-scores as their respective naive versions for Bokulich₂. For Bokulich₃ benchmark we observed higher F-scores for all LSH based Canopy clustering methods. Finally for Bokulich₆ benchmark Fscores of UCLUST and SWARM were improved comparing to respective naive versions.

For all benchmarks we got very strong correlations between taxonomic profiles at genus level from UCLUST, SUMACLUST, SWARM and their respective LSH based Canopy counterparts. The highest correlation was observed for Bokulich₆ benchmark between UCLUST and CC_{UCLUST} with 0.9831 and the lowest was observed for Canadian Soil metagenome data between SUMACLUST and $CC_{SUMACLUST}$ with 0.7581. From observed F-scores and correlation values we can say that running a sequence clustering method with our proposed LSH based Canopy cluster framework can bring similar or better results. Next we will discuss about Runtime Comparisons. CC_{MST} provided relatively similar F-scores and high ρ values indicating strong correlations of taxonomic outcomes with other methods.

B. Runtime Comparison

Table III shows the runtime in minutes of UCLUST, SUMACLUST, SWARM and their respective versions with our proposed Canopy clustering pipeline. Time information in Table III was recorded while populating performance metric values for Table II for the parameter setting mentioned in Experimental Details section. Improving runtime while maintaining similar or better clustering results was one of the major motivations of our proposed approach. We can see from Table III that CC_{UCLUST} outperforms

⁵https://docs.python.org/2/library/multiprocessing.html

⁶http://biocore.github.io/pynast/

UCLUST for all datasets. The highest speed up for UCLUST was observed for Bokulich₂ dataset where UCLUST was 2.10 times faster with LSH based Canopy than the naive version. For SUMACLUST we can see from Table III that $CC_{SUMACLUST}$ outperforms naive SUMACLUST in most cases. Major improvement in runtime was observed in large scale real world metagenome datasets specially Canadian Soil and Global Soil datasets. The highest speed-up for SUMACLUST with Canopy clustering was observed to be 7.70x for Global Soil dataset.

We can see a similar footprints for CC_{SWARM} from Table III. The highest reduction in runtime for SWARM with Canopy clustering was observed to be 4.79x for Global Soil dataset. From these observations we can conclude that LSH based Canopy clustering not only reduces the runtime of an expensive clustering method but also scales well and performs better for larger benchmarks comparing to the smaller ones.

C. Biodiversity Comparison

Clustering sequences in metagenome data will provide OTUs that represent biodiversity contained in the samples from which data is collected. Hence we also need to compare the biodiversity represented by OTUs from naive methods and their respective Canopy clustering variations used in this study. Table IV shows Faiths phylogenetic diversity metric (PD), Shannon and Simpson index after clustering with different methods. These metrics are some of the popular Alpha Diversity metrics meaning that they measure species diversity in sites or habitats at a local scale. These metric values are generated per sample basis. Table IV shows ranges of metric values in [minimum - maximum] format provided by methods with and without Canopy clustering. Any sample of a dataset will take value from this range. These ranges may not be same since OTUs vary over clustering methods. But they should be similar. We can see from Table IV that Canopy clustering based methods produce similar ranges of values as their naive counterparts. No significant changes in diversity metric values were observed which indicates that Canopy based approaches reproduce similar biodiversity while reducing runtime.

D. Effect of Varying Number of Multi-processes

Figure 4a-4f shows how the number of multiprocess can affect runtime of CC_{UCLUST} , $CC_{SUMACLUST}$ and CC_{SWARM} on six benchmarks used in this study. We observed that increasing the number of multiprocess can reduce total runtime. The steepest curve showing reduction in runtime can be found in Figure 4e for $CC_{SUMACLUST}$ on Canadian Soil dataset. For single process version $CC_{SUMACLUST}$ is slower in most cases except Bokulich₂ dataset where it's runtime is better comparing to CC_{SWARM} . For Body Site dataset (Figure 4d) the

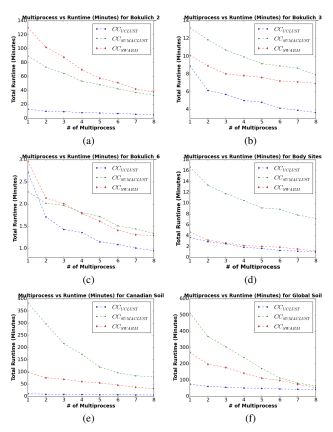


Figure 4. Runtime comparison of Canopy clustering framework with UCLUST, SUMACLUST and SWARM on Bokulich₂, Bokulich₃, Bokulich₆, Body Sites, Canadian Soil and GLobal Soil datasets.

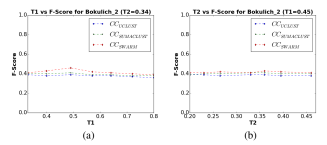


Figure 5. 5a and 5b show effect of varying T1 and T2 on F-scores for the largest synthetic dataset Bokulich₂

 CC_{UCLUST} and CC_{SWARM} have closer runtime with respect to increasing number of multiprocess. Figure 4f shows reduction in runtime for the largest dataset used in this study namely Global Soil with more than nine million reads.

E. Effect of Varying T1 and T2 Parameter of Canopy Clustering

Figure 5a-5b shows effect of varying T1 (T2 fixed at 0.34) and T2 (T1 fixed at 0.45) on F-scores for the largest synthetic dataset Bokulich₂. Reducing T1 means comparatively *tighter* soft-threshold which will reduce repetitions of instances in multiple canopies. For a fixed T2 this

Table II

PERFORMANCE COMPARISON [F-SCORE AND PEARSON CORRELATION COEFFICIENT (ρ)]

Methods	Comparison Metric	Datasets							
		Synthetic			Real World				
		Bokulich ₂	Bokulich ₃	Bokulich ₆	Body Sites	Canadian Soil	Global Soil		
UCLUST	F-Measure	0.39	0.40	0.51	N/A	N/A	N/A		
CC	F-Measure	$ \overline{0.39}$	0.41	0.52	\bar{N}/A	- $ N/A$ $ -$	$\overline{N/A}$		
CC_{UCLUST}	Correlation (ρ) with Respect to UCLUST	0.9831	0.9753	0.9831	0.9682	0.8419	0.9824		
SUMACLUST	F-Measure	0.40	0.41	0.51	N/A	N/A	N/A		
$CC_{SUMACLUST}$	F-Measure	0 .41	0.42	0.51	\bar{N}/A	$ N/\bar{A}$ $ -$	$\overline{N/A}$		
	Correlation (ρ) with Respect to SUMACLUST	0.9709	0.9813	0.9538	0.9518	0.7643	0.8714		
SWARM	F-Measure	0.46	0.48	0.55	N/A	N/A	N/A		
CC_{SWARM}	F-Measure	$ \bar{0.46}$	0.49	0.56	$-\bar{N}/A$	- $ N/A$ $ -$	- $ N/A$ $-$		
	Correlation (ρ) with Respect to SWARM	0.9817	0.97861	0.9251	0.9648	0.7581	0.9143		
CC_{MST}	F-Measure	0.41	0.42	0.52	N/A	N/A	N/A		
	Correlation (ρ) with respect to UCLUST	0.7461	0.8143	0.8272	0.7417	0.6935	0.7974		
	Correlation (ρ) with respect to SUMACLUST	0.7182	0.6903	0.8613	0.8293	0.6791	0.8213		
	Correlation (ρ) with respect to SWARM	0.7914	0.7352	0.8904	0.7619	0.7372	0.8502		

Table shows values of F-measures and Pearson Correlation Coefficient (ρ -value) of UCLUST, SUMACLUST, SWARM and their respective versions with Canopy clustering. F-measures is only available for synthetic datasets but not for real world datasets since no ground truths like known taxonomic profiles are available for them. ρ -value was calculated based on the taxonomy profiles at Genus level generated from clustered OTUs provided by a method and it's corresponding LSH based Canopy counterpart. Higher F-measures reflect better clustering by adhering to ground truth. Higher ρ -values reflect stronger correlation between taxonomic profiles.

Table III
RUNTIME COMPARISON (IN MINUTES)

	Datasets		Methods											
			UCLUST			SUMACLUST				SWARM				
Type	Title	# of Reads	UCLUST	Canopy	with	Speed Up	SUMACLUST	Canopy	with	Speed Up	SWARM	Canopy	with	Speed Up
					Canopy				Canopy				Canopy	
	Bokulich ₂	6,938,836	12.71	4.81	6.03	2.10x	87.53	5.73	33.03	2.65x	128.12	5.29	37.87	3.38x
Synthetic	Bokulich ₃	3,594,237	8.91	2.91	4.89	1.82x	11.73	3.05	7.89	1.48x	9.10	2.98	6.93	1.31x
	Bokulich ₆	250,903	1.21	0.49	1.08	1.12x	1.28	0.52	1.34	0.96	1.97	0.51	1.29	1.53x
	Body Sites	886,630	2.12	1.03	1.51	1.40x	15.42	1.09	7.88	1.95x	3.64	0.98	1.71	2.13x
Real World	Canadian Soil	2,966,053	9.55	2.18	5.91	1.62x	363.96	2.91	81.18	4.48x	97.50	3.11	32.16	3.03x
	Global Soil	9,252,764	72.47	21.38	43.53	1.67x	510.92	20.82	66.27	7.70x	269.07	21.17	56.14	4.79x

means lower T1 will yield better canopy assignment. From Figure 5a we can see that when T1's range is in 0.4 to 0.6 our proposed approach provides better F-Scores. On the other hand, increasing T2 means comparatively *softer* tight-threshold for canopies. As a result instances will be prematurely assigned to canopies without waiting for best match. From Figure 5b we can see that when T2's range is in 0.2 to 0.35 our proposed approach provides better F-Scores. Lower T2 may cause higher runtime since instances will continue to reappear until T2 is met.

VI. CONCLUSION AND FUTURE WORK

We propose a framework that can be used as preclustering for any accurate and relatively expensive clustering on large scale metagenomic datasets. Our proposed approach scales well with large datasets and provide significant reduction in computation time. We demonstrate that our approach provides similar outcome in terms of biodiversity metrics used in this study, scores based on ground truth and taxonomic correlation with corresponding relatively expensive cluster. Our approach takes advantage of the multi-core CPU systems by partitioning the large dataset roughly with fast and cheaper pairwise distance measure and then deploying comparatively expensive clustering in parallel which considers only data points that are inside the partition. We plan to develop standalone cluster mechanism for the canopies in future.

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Table IV
BIODIVERSITY COMPARISON [FAITHS PHYLOGENETIC DIVERSITY METRIC (PD), SHANNON AND SIMPSON]

Methods	Comparison Metric	Datasets							
	-			Real World					
		Bokulich ₂	Bokulich ₃	Bokulich ₆	Body Sites	Canadian Soil	Global Soil		
	PD Range	[171.95 - 221.85]	[186.90 - 212.84]	[104.51 - 104.51]	[1.46 - 46.79]	[0.30 - 1352.73]	[2.98 - 3.29]		
UCLUST	Shannon Range	[2.52 - 3.51]	[2.43 - 3.54]	[5.87 - 5.87]	[0.29 - 7.67]	[2.32 - 10.85]	[1.84 - 8.30]		
	Simpson Range	[0.55 - 0.75]	[0.55 - 0.76]	[0.96 - 0.96]	[0.049 - 0.98]	[0.80 - 0.99]	[0.0 - 0.98]		
	PD Range	[164.79 - 217.72]	[169.41 - 198.36]	[109.33 - 109.33]	[2.37 - 47.13]	[0.52 - 1419.31]	[3.02 - 3.81]		
CC_{UCLUST}	Shannon Range	[2.61 - 3.83]	[2.92 - 3.91]	[6.41 - 6.41]	[0.93 - 7.13]	[3.26 - 7.61]	[3.72 - 7.38]		
	Simpson Range	[0.64 - 0.87]	[0.56 - 0.93]	[0.96 - 0.96]	[0.081 - 0.99]	[0.84 - 0.99]	[0.21 - 0.99]		
	PD Range	[106.00 - 162.78]	[142.85 - 174.19]	[89.22 - 89.22]	[0.93 - 39.47]	[0.59 - 1279.29]	[2.98 - 3.29]		
SUMACLUST	Shannon Range	[2.00 - 3.01]	[2.19 - 3.28]	[5.48 - 5.48]	[0.16 - 7.43]	[2.32 - 7.32]	[1.00 - 7.89]		
	Simpson Range	[0.52 - 0.73]	[0.54 - 0.75]	[0.95 - 0.95]	[0.027 - 0.98]	[0.80 - 0.99]	[0.40 - 0.98]		
	PD Range	[114.96 - 171.57]	[147.85 - 187.91]	[93.81 - 93.81]	[0.86 - 41.63]	[0.81 - 1292.34]	[1.37 - 4.89]		
$CC_{SUMACLUST}$	Shannon Range	[2.51 - 3.94]	[2.96 - 4.11]	[5.94 - 5.94]	[1.21 - 7.13]	[3.12 - 7.79]	[2.17 - 7.25]		
	Simpson Range	[0.68 - 0.79]	[0.51 - 0.74]	[0.96 - 0.96]	[0.06 - 0.99]	[0.88 - 0.99]	[0.23 - 0.98]		
	PD Range	[18.37 - 24.73]	[17.36 - 19.81]	[30.84 - 30.84]	[1.44 - 28.66]	[0.54 - 706.57]	[5.79 - 6.18]		
SWARM	Shannon Range	[2.98 - 3.91]	[2.01 - 3.04]	[5.03 - 5.03]	[0.28 - 7.63]	[1.0 - 7.79]	[1.66 - 7.81]		
	Simpson Range	[0.70 - 0.82]	[0.53 - 0.74]	[0.95 - 0.95]	[0.05 - 0.98]	[0.50 - 0.99]	[0.00 - 0.99]		
	PD Range	[19.18 - 26.87]	[18.43 - 22.61]	[31.48 - 31.48]	[2.34 - 29.97]	[1.37 - 748.71]	[2.34 - 8.46]		
CC_{SWARM}	Shannon Range	[1.66 - 4.87]	[1.19 - 4.13]	[6.03 - 6.03]	[0.89 - 7.13]	[2.81 - 8.06]	[2.81 - 7.87]		
	Simpson Range	[0.66 - 0.88]	[0.41 - 0.81]	[0.91 - 0.91]	[0.11 - 0.99]	[0.74 - 0.99]	[0.14 - 0.99]		
	PD Range	[10.43 - 89.61]	[12.83 - 92.41]	[63.21 - 63.21]	[1.46 - 34.15]	[2.73 - 824.62]	[1.81 - 5.37]		
CC_{MST}	Shannon Range	[0.96 - 3.91]	[1.21 - 3.16]	[4.91 - 4.91]	[2.02 - 6.51]	[2.72 - 7.83]	[2.17 - 6.29]		
	Simpson Range	[0.14 - 0.68]	[0.32 - 0.73]	[0.92 - 0.92]	[0.07 - 0.94]	[0.58 - 0.94]	[0.14 - 0.98]		

Table shows ranges of values for Faiths Phylogeny Diversity (PD), Shannon and Simpson coefficient over all samples in a dataset in the format [minimum - maximum]. Most of these datasets contain multiple samples and Alpha diversity metrics like PD, Shannon and Simpson values are generated for each of these samples separately. Biodiversity metric values changes significantly over samples e.g. diversity from hair samples and teeth cavity are supposedly different. So instead of mean values this Table represents [minimum - maximum] ranges of values a sample can take. Similar ranges reflect similar diversity.

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