Improving Clustering in Metagenomic Data Using Canopy and Locality Sensitive Hashing

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Abstract-Recent sequencing technologies like Shotgun sequencing, Next generation sequencing etc. allow sequencing of genomes from samples taken directly from environments hosting multiple organisms that co-exist as communities within the ecological regions. This collective genomic process known as metagenomics has created the necessity 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 3 popular clustering mechanisms in literature on 3 synthetic and 3 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

With earlier sequencing technology nearly 50 million bases of nucleotide sequence were available in public archives 1. Now a single sequencing instrument can produce over 1 trillion nucleotide bases in just a single run [1]. Latest genome sequencing technologies like Shotgun and High Throughput sequencing have paved the way to Metagenomics - 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 [2][3]. 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 can provide an understanding of the role played by microbes with regards to human health and disease [4].

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 [5]. Many Sequence assemblers require reference genome sequence and the process of assembly is exceedingly complex, challenging and time consuming [6]. 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 [7]. OTU is used to classify groups of closely related individuals from similar or different taxonomic levels [8]. It is the most commonly used microbial diversity unit, especially when analyzing the small subunit 16S or 18S rRNA marker gene microbial datasets [9]. 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 [10].

Many Unsupervised and Supervised 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

¹http://www.ncbi.nlm.nih.gov/genbank/statistics/

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

Analyzing metagenome sequences has poses great computational challenges since current DNA sequencing technologies generate hundreds of gigabytes of data in a single run and many computational methods have been developed over the years [11], [12]. Both availability of data and recent computational methods have enabled new detailed research into the human microbiome [13] and characterization of the Earth ecosystems microbiome, such as the Earth Microbiome Project (EMP)². Metagenome dataset analysis usually begins with clustering raw biological sequence reads into operational taxonomic units (OTUs) based on sequence similarity. This process frequently referred to as OTU clustering also known as delineating. Focus of study is OTU clustering methods and ways to improve them in terms of OTU quality and runtime.

Some of the popular and early OTU clustering methods proposed were CD-HIT [14], DOTUR [15], MOTHUR [16] and ESPRIT [17]. Algorithm used in CD-HIT is a greedy incremental clustering algorithm and performs ordering on sequences. The ordering used in CD-HIT is determined by sequence length. CD-HIT sorts all sequences by length in decreasing order. The first sequence is considered as the representative of the first cluster. Rest of the sequences are compared with first sequence. If any of the sequences fall above the threshold for similarity, then it is grouped together with the first sequence. Subsequent comparisons have to fall above the threshold for similarity with all sequences in this cluster group. After this first round of cluster, the next ungrouped sequence is set as the representative of the next cluster. This process is repeated untill all sequences are part of a cluster. DOTUR, MOTHUR and ESPRIT exclusively use pairwise distance matrix as input and then perform hierarchical clustering clustering on 16S sequences. MOTHUR and DOTUR utilizes global alignment score between all pairs of sequences for pairwise distances computation. ESPRIT computes k-mer distance for each pair of input sequences, avoiding the expensive global alignment distance calculation. Number of sequence comparisons is also reduced by ESPRIT because of usage of heuristics. Hashing techniques have been utilized in earlier works for clustering metagenomic datasets. MC-LSH [18] utilizes an efficient locality sensitive based hashing function to approximate the pairwise sequence similarity. Similarity among sequences is computed based on randomly chosen indices that essentially compresses the input sequence. MC-MinH [19] algorithm uses the min-wise [20] hashing approach, along with a greedy clustering algorithm to group 16S and whole metagenomic sequences. Unequal length sequences are represented using contiguous subsequences or k-mers, and then pairwise similarity are approximated using independent min-wise hashing. Mash [21], uses MinHash locality sensitive hashing to reduce large sequences to a representative sketch and rapidly estimate pairwise distances between genomes or metagenomes. More recent methods for clustering whole metagenome sequence reads include TOSS [22], AbundanceBin [23], CompostBin [24], LikelyBin [25] and MetaCluster [26]. All unique k-mers are first clustered in TOSS then clusters are merged based on k-mer repetition.In AbundanceBin sequencing reads are modeled as mixture of Poisson distributions. Then Maximization (EM) algorithm is used to infer model parameters and final clustering. Principal component analysis is used in CompostBin to project the data into lower dimensional space and then uses the normalized cut clustering [27] algorithm to group sequences into taxa specific clusters. CompostBin also uses phylogenetic markers to assign clusters. LikelyBin uses an unsupervised, maximum-likelihood approach for clustering based on k-mer distributions of sequences. MetaCluster creates a top-down separation followed by a bottom-up merging for clustering metagenomic fragments based on k-mer frequencies and Spear-man distance computation. UCLUST [28] follows a greedy process and creates seeds of sequences which generate clusters based on percent identity. It is very fast but closed source software. Only the 32 version is made available for free with very limited documentations for academic and non-profit usage. 64-bit version of the software require expensive license. In last 2 years, 2 relatively new clustering methods have been introduced Swarm [29][30] and SUMACLUST [31]. These methods are some of the currently popular, relatively fast [32] and open source softwares. 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. SUMACLUST follows a similar approach as UCLUST. Abundance-ordered list of input sequences are compared against the representative set of already-chosen sequences in SUMACLUST.

In this study we have introduced a very fast initial partitioning of large scale metagenomic datasets with Canopy Clustering [33] and Locality Sensitive Hashing [34][35][36]. This way efficient and scalable parallelism in OTU-clustering can be achieved that is capable of making

²http://www.earthmicrobiome.org/

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 on 6 standard large scale metagenomic datasets to validate and compare our proposed approach. Our proposed framework has shown similar biodiversity metric values, higher F-measure values for datasets with known taxonomic profiles as ground truth and significantly less amount of runtime specially for large datasets.

III. METHODS

A. Overview

In this section we will give a brief description of Canopy Clustering and Locality Sensitive Hashing, their potential in metagenome data clustering, our motivation for using them in this work and a framework that enables them to be used with other expensive clustering harnessing the power of parallel processing in multi-core CPUs.

B. Canopy Clustering

Canopy Clustering was introduced in [33]. It is often used as preprocessing step for other accurate and expensive clustering methods like K-means algorithm or the Hierarchical clustering algorithm. It is intended to speed up clustering operations on large data sets, where using another algorithm directly may be impractical in terms of run time and memory consumption for large number of pairwise distance calculations due to the size of the data set.

Canopy algorithm can greatly reduce the number of distance computations required for clustering by roughly partitioning the data into overlapping subsets. Then more accurate clustering can be performed inside the formed canopies by only measuring distances between pairs of data points that belong to same canopies. Points outside a canopy will not participate in clustering for that canopy. If a dataset contains total N instances then worst case calculations without canopy clustering is N^2 . After canopy clustering if the number of canopies is N^2 where N^2 where N^2 is the number of instances in N^2 total difference N^2 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. The process of canopy clustering begins with all the data points to be clustered. A point is randomly removed from the set, beginning a new canopy. All other data points are tested and assigned to the new canopy if the distance less than the soft threshold T_1 . Moreover if the distance of the point is less than the tight threshold T_2 then it is removed from the original set. This process is repeated until there are no more data points in the set to cluster. These relatively cheaply clustered canopies can be clustered using a more expensive but accurate algorithm. A fast approximation which is discussed in next section will be used for Canopy membership identification. Once the dataset is partitioned other popular metagenome clustering can be used to subcluster canopies in parallel achieving both advantages of reduced calculations due to canopies and lower runtime due to proper utilization of parallel processing.

C. Locality Sensitive Hashing

A Locality Sensitive Hashing (LSH) [34][35][36] scheme 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 satisfies the following conditions for any two points $p,q\in \mathcal M$, using a function $h\in \mathcal F$ which is chosen uniformly at random:

- (i) If $d(p,q) \leq R$, then h(p) = h(q) with probability at least P_1 .
- (ii) If $d(p,q) \ge R$, then h(p) = h(q) with probability at most P_2 .

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 [36]. In [37] data points were randomly projected to low dimensional bit signatures such that cosine distance is approximately preserved. LSH provides a convenient way of projecting high dimensional data points into low dimensional space and compute fast approximate distance between pairs of data points in this transformed low dimensional space. These characteristics of LSH make it appropriate to be used as a fast distance calculation mechanism between 2 reads in large scale metagenomic datasets.

LSH function based on random projection and bit sampling for hamming distance was introduced in [36]. This approach works for the Hamming distance over d-dimensional vectors $\{0,1\}^d\{0,1\}^d$. Here, the family $\mathcal F$ of hash functions is the family of all the projections of points on one of the d coordinates, for example $\mathcal F = \{h: \{0,1\}^d \to \{0,1\} \mid h(x) = x_i \text{ for some } i \in \{1,...,d\}\},$

where x_i is the ith coordinate of x. A random function h from \mathcal{F} selects a random bit from the input point. This family has the following parameters: $P_1 = 1 - R/d$, $P_2 = 1 - cR/d$. In this study we have chosen a similar 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 the actual data dimension for faster distance calculations. Afterwards we used Hamming distance between 2 projected data points with reduced dimensions to get approximate distance between them. Given a random projection matrix v' of size $1 \times d$:

$$\begin{bmatrix} w_{1,1} & w_{1,2} & \dots & w_{1,d} \end{bmatrix}$$

The dot product of these vector

And a vector a of size $1 \times d$ representing a single data point in dataset :

$$\begin{bmatrix} a_{1,1} & a_{1,2} & \dots & a_{1,d} \end{bmatrix}$$

The dot product of these 2 vectors provides a scalar value:

$$v \cdot a = \sum_{i=1}^{d} v_i a_i \tag{2}$$

$$= v_1 a_1 + v_2 a_2 + \ldots + v_d a_d \tag{3}$$

$$=s$$
 (4)

The value s in eq. 4 is a scalar quantity. If vector v is considered as hyperplane in d-dimensional space then the sign of the scalar output of the dot product v.a refers to which side of the hyperplane v does the data point a exists. We interpret this information with a bit value. Considering matrix V of size $n \times d$ as a collection all random weights:

$$\begin{bmatrix} w_{1,1} & w_{1,2} & \cdots & w_{1,d} \\ w_{2,1} & w_{2,2} & \cdots & w_{2,d} \\ \vdots & \vdots & \ddots & \vdots \\ w_{n,1} & w_{n,2} & \cdots & w_{n,d} \end{bmatrix}$$

the dot products between each of the rows of matrix V and a data point a will result in a collection of n bits for that data point. In other words the d-dimensional data point will take the form of $\{0,1\}^n$. Hamming distance between 2 such binary number indicates the number of randomly generated hyperplanes for which 2 corresponding data points reside in opposing sides. This is the approximate distance measure that we will use for next step of our proposed approach - Canopy Clustering.

D. Sub-Clustering Inside Canopies

We have used 3 recent and popular sequence clustering methods as an expensive clustering measure inside canopies in this study. UCLUST [28], SUMACLUST [31] and SWARM [29] were used for sub-clustering canopies.

UCLUST is closed-source software and provides very limited documentations³. 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. SUMACLUST is 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. Finally 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.

E. Proposed Framework

Figure 1 shows an overview of our proposed LSH-Canopy clustering approach for metagenomic data. After computing the kmers and LSH code 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 smaller input files from original dataset based on canopy membership information. This will give one smaller input file per canopy. This way larger dataset will be partitioned and new smaller datasets will be created for these partition. Finally each smaller input files representing canopies will be provided as input to more accurate nucleotide sequence clustering methods in parallel. In this study UCLUST, SUMACLUST and SWARM will be used as accurate clustering methods for canopies.

IV. EXPERIMENTAL EVALUATIONS

A. Dataset Description

Different types of previously published mocked and real world genuine datasets were used for extensive evaluation of our approach and comparison with other methods. We have used three different previously published mock community data sets- three of them are 16S rRNA gene mock community data sets (Bokulich_2, Bokulich_3, and Bokulich_6) from [38]. We have also used three previously published natural data sets: a 16S rRNA gene soil data set (canadian_soil)from [39], a 16S rRNA gene human data set (body_sites) from [40], and an 18S rRNA gene soil data set

³http://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	[38]	6,938,836	4	189-251	HiSeq
Bokulich_3	Mock	[38]	3,594,237	4	114-151	HiSeq
Bokulich_6	Mock	[38]	250,903	1	114-150	HiSeq
Canadian Soil	Genuine	[39]	2,966,053	13	76–10	HiSeq
Body Sites	Genuine	[40]	886,630	602	117-351	GS FLX
Global Soil	Genuine	[41]	9,252,764	57	119-151	HiSeq

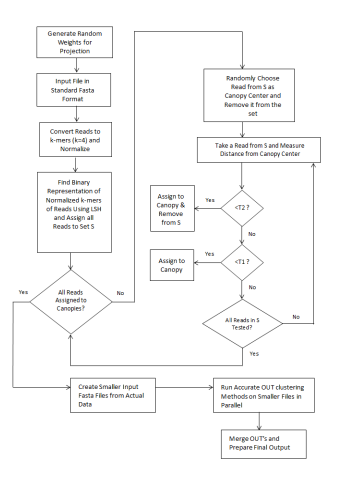


Figure 1. Workflow of LSH-Canopy for Large Scale Metagenome Data

(global_soil) from [41]. Statistics and relevant information regarding the datasets are provided in Table I. Description of the various dataset used in this paper is as follows:

Mock Datasets

Bokulich_2 - This dataset was prepared using the Illumina TruSeq v2 paired—end library preparation kit. It is a 16S rRNA gene mock microbial community data set. It has total 4 samples. For 2 of these samples abundance of mock communities were distributed evenly. For the other 2 samples abundance of mock communities were inserted randomly. This dataset contains 19 taxonomic Families, 19

Genera, 22 Species and 22 Strains in total. This dataset can be found in the QIIME database under the study identifiers 1685.

Bokulich_3 - Similar to Bokulich_2 except that it was sequenced in different environment and tool. The dataset was prepared with the TruSeq v1 paired-end library kit at Illumina Cambridge. Available at QIIME database under the study ID 1686.

Bokulich_6 - A 16S rRNA dataset which was sequenced at Washington University School of Medicine. It has 1 sample which contains evenly distributed mock microbial community. This dataset contains 13 taxonomic Families, 23 Genera, 44 Species and 48 Strains in total.

All these datasets from Bokulich et al.[38] are available at QIIME database⁴ under their respective ID's. Since these are mock datasets their taxonomic profile of microbiome are known.

Genuine Datasets

Canadian Soil⁵ - This dataset contains genomic data of Canadian soil spanning from Arctic Tundra to Agricultural soil suitable for different agricultural products like soi, wheat, corn etc. in different regions of Canada.

Body Sites - This dataset contains composition of bacterial communities from up to 27 body sites in healthy adults. A collection of 602 samples acquired from different body sites of human subjects are provided with meta-data.

Global Soil - Ramirez et al.[41] combined data representing a range of biomes from Alaska to Antarctica from two previous studies of Dinsdale EA, et al.[42] and Lauber CL et al.[43] to compare with below-ground diversity in New York City's Central Park.

All of these datasets have been used in previous studies [38],[44]. Any analysis on these datasets requires appropriate preprocessing which can significantly affect results of clustering and taxonomic classification based on them [38]. Performance of different open source sequence clustering methods were assessed and compared in [44] on these datasets and they have also provided preprocessed and filtered using qiime and provided together (URL: https://github.com/ekopylova/otu-clustering).

⁴http://qiime.org/home_static/dataFiles.html

⁵http://www.cm2bl.org/

B. Evaluation Metrics

We have used some standard and popular performance measure to assess our proposed clustering method and compare with other clustering methods.

Faiths phylogenetic diversity metric (PD) - Originally described in [45] also known as PD is based on the Phylogeny tree. It combines all the branch lengths of the tree as a measure of diversity. So, if a new OTU is found and it is closely related to another OTU in the sample, it will be a small increase in diversity. However, if a new OTU is found and it comes from a totally different lineage than anything else in the sample, it will contribute a lot to increasing the diversity:

 $\begin{tabular}{lll} \textbf{Shannon} & \textbf{Entropy} & \textbf{-} & \textbf{Shannon-Wiener diversity index} \\ \textbf{is defined as:} & \end{tabular}$

$$H = -\sum_{i=1}^{s} (p_i \log_2 p_i)$$
 (5)

where s is the number of OTU and p_i is the proportion of the community represented by OTU i.

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

$$1 - \sum p_i^2 \tag{6}$$

where where p_i is the proportion of the community represented by OTU i.

F-measure - For 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 or expected taxonomic composition. The following definitions were used:

$$precision = \frac{TP}{(TP + FP)} \tag{7}$$

$$recall = \frac{TP}{(TP + FN)} \tag{8}$$

$$FScore = \frac{2 \times precision \times recall}{(precision + recall)} \tag{9}$$

C. Experimental Details

For kmers, the value of parameter k was set to 4. Which means occurrences of all possible combination of 4 letter nucleotides were generated from the reads of datasets. Since k=4 and there can be 4 possible nucleotide representatives (A,C,G and T), total number of features became $4^4=256$. This way we were able to convert the string representation of nucleotide data in standard FASTA format into numeric. After finding the kmers, each numeric representation of data reads were normalized to range [0-1].

For Locality Sensitive Hashing the number of Hyperplanes (parameter d) was set to 150. One of the motivations of using LSH was fast approximation of distances from one data read to another while reducing feature dimension. Since total number of features became 256 after numeric representation using kmer, we wanted to reduce the number of features while maintaining a relatively accurate yet fast distance approximation from LSH. So the value of parameter d should be less than the actual number of feature but not too small because that might lead to wrong distance approximation.

The 2 parameters of Canopy clustering namely T1 (loose threshold) and T2 (tight threshold) based on the reduced feature space provided by LSH and normalized hamming distance measure define which data reads will be grouped initially before the preferred expensive clustering begins. For this experiment and comparison we have chosen T1 = 0.46and T2 = 0.34. For parameter d = 150, canopy threshold T1 = 0.46 refers that a data read can belong to a canopy if the hamming distance is $150 \times 0.46 = 69$ or less. And threshold T2 = 0.34 refers that if the hamming distance between randomly chosen canopy centroid and data read is $150 \times 0.34 = 51$ or less then the read should not be included in any other canopy assuming that we have found a stable canopy membership for the read. Once canopy memberships of all reads were found we created multiple smaller data files in standard FASTA format from actual dataset to be used by expensive cluster.

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 [46] 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 multi-core CPU based computing systems we utilized Python's multiprocessing module instead of threading module. According to Python's documentation⁶ multiprocessing module allows the programmer to fully leverage multiple processors on a given machine by spawning subprocesses instead of threads. In our experiment these subprocesses are independent Python programs running more accurate and expensive clustering on smaller datasets created by canopy clusterings. In our experiment the number of parallel processes was set equal to the number of CPU cores (which is 8 in our case) with an expectation that OS will deploy each of those processes in separate cores unlike threading. This is convenient since it utilizes 100% of the CPU in most cases.

After receiving output from all canopy clustering we merged the outputs to make a single clustering result. Some

⁶https://docs.python.org/2/library/multiprocessing.html

performance metric used in this study like Faiths Phylogenetic Diversity (PD) metric require sequence alignment. We used PyNast⁷ [47] open source sequence aligner for aligning clustered output.

V. RESULTS DISCUSSION

A. Clustering Performance Comparison

Performance of UCLUST [28], SUMACLUST [31], SWARM [29] and their respective versions with our proposed approach Locality Sensitive Hashing based Canopy Clustering (LSH-Canopy) is provided in Table II. Datasets used in this study contain multiple samples. Performance metrics like Phylogenetic Diversity (PD), Shannon entropy and Simpson index are generated per sample basis for any dataset. To show comparative performance of the clustering methods used in this study we showed the range of values the performance metrics in [min-max] where min and maxrepresent the minimum and maximum values respectively observed in the whole dataset regardless of samples for a particular clustering method. This represents the range of values a performance metric can take over all samples in a dataset. Performance of 2 clustering methods on same dataset can be assumed similar in terms of biodiversity if their minimum and maximum values for performance metrics are similar. Metrics representing biodiversity can significantly change over clustering methods, their respective data pre-processing pipeline, post processing, range of OTU lengths, number of OTU's etc. Nevertheless, [min - max]range of a clustering method and it's counterpart with our proposed LSH-Canopy pipeline should be roughly similar. F-measure and Pearson Correlation Coefficient (ρ -value) can take only one value each for a clustering method on a dataset. And the correlation metric (ρ -value) is applicable for clustering methods with our proposed approach only since this is a correlation between the output of a standard clustering method and it's respective LSH-Canopy version. Thus we can interpret a better outcome of this experiment as (i) similar range of biodiversity (PD, Shannon and Simpson values), (ii) higher F-measure and Correlation (ρ -value) with respect to corresponding clustering without our proposed LSH-Canopy pipeline and finally (iii) lower running time.

From Table II we can see that UCLUST and LSH-Canopy-UCLUST provide similar range for PD, Shannon and Simpson values for both synthetic and real world datasets. Moreover, LSH-Canopy-UCLUST provided higher F-score than UCLUST for all 3 synthetic datasets used in this study. The lowest F-score observed by UCLUST was 0.39 for Bokulich_2 dataset. with LSH-Canopy-UCLUST we achieved 0.42 for the same dataset. For synthetic datasets taxonomic profiles were known as prior. This taxonomic distribution can be taken as ground truth and it can be used to

identify whether the cluster representatives actually support this ground truth. A higher F-score is an indicator that relevant and important cluster representatives are retained as they support ground truth. Higher F-scores of our proposed approach is due to the nature of how canopy is formed. A single data instance can belong to multiple canopies due to the nature of soft threshold T1. As a result a single read can contribute to multiple Operational Taxonomic Units (OTU) in metagenome data. As a result even after removing singletons, expensive clustering inside a canopy can bring higher numbers of useful OTU's. For real world datasets LSH-Canopy-UCLUST provided similar range of values for PD, Shannon and Simpson. Correlation of LSH-Canopy-UCLUST with navie UCLUST at the Genus level of assigned taxonomy was observed to be highly correlated. The lowest ρ -value was 0.8177 for Canadian Soil metagenomic data which is the largest dataset used in this study containing more than 9 million reads.

Our proposed approach also improves the outcomes of SUMACLUST. From Table II we can see that SUMACLUST provided higher F-score than UCLUST for Bokulich_2 and Bokulich_3 and same value for Bokulich_6. After running SUMACLUST with LSH-Canopy we observed same F-score as SUMACLUST for Bokulich_2 and higher for other 2 synthetic datasets. SUMACLUST with our proposed approach outperforms the naive SUMACLUST and UCLUST. Ranges of PD, Shannon and Simpson were similar to naive SUMACLUST for both synthetic and real world datasets. ρ -values were higher indicating a strong positive correlation with naive SUMACLUST specially for synthetic datasets. Lowest ρ -value was 0.7859 which was observed for larger Canadian Soil real world metagenome dataset.

Finally SWARM with our proposed LSH-Canopy pipeline also showed better results than naive SWARM. We observed the highest F-score of 0.56 in this study for Bokulich_6 dataset from both naive and LSH-Canopy version of SWARM. For the other 2 synthetic datasets our proposed LSH-Canopy version of SWARM outperforms the naive counterpart. Ranges for PD, Shannon and Simpson were similar and correlation values were observed to be high. The lowest ρ -value for LSH-Canopy-SWARM was 0.7736 for Canadian Soil dataset.

B. Runtime Comparison

Table III shows the runtime in minutes of UCLUST, SUMACLUST, SWARM and their respective version with our proposed LSH-Canopy 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 was one of the major motivations of our proposed approach - fast distance approximation using LSH, using this approximation for initial canopy clustering to reduce pairwise comparison

⁷http://biocore.github.io/pynast/

Table II
PERFORMANCE COMPARISON

Methods	Comparison Metric	Datasets						
		Bokulich_2	Bokulich_3	Bokulich_6	Body Sites	Canadian Soil	Global Soil	
	F-Measure	0.39	0.40	0.51	N/A	N/A	N/A	
UCLUST	PD	[171.95 - 221.85]	[186.90 - 212.84]	[2.98 - 3.29]	[1.46 - 46.79]	[0.30 - 1352.73]	[0.30 - 1352.73]	
UCLUST	Shannon	[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	[0.55 - 0.75]	[0.55 - 0.76]	[0.96 - 0.96]	[0.049 - 0.98]	[0.8 - 0.99]	[0.0 - 0.98]	
	F-Measure	0.42	0.44	0.54	N/A	N/A	N/A	
!	ρ–value	0.9911	0.9935	0.9787	0.9581	0.8177	0.9824	
LSH-Canopy-UCLUST	PD	[162.49 - 206.72]	[168.01 - 194.66]	[109.33 - 109.33]	[2.37 - 44.52]	[0.52 - 1319.31]	[3.02 - 3.26]	
	Shannon	[2.26 - 3.94]	[2.71 - 4.86]	[6.2 - 6.2]	[0.75 - 7.06]	[3.16 - 11.94]	[4.36 - 11.48]	
	Simpson	[0.64 - 0.77]	[0.58 - 0.83]	[0.96 - 0.96]	[0.097 - 0.99]	[0.88 - 0.99]	[0.11 - 0.99]	
	F-Measure	0.40	0.41	0.51	N/A	N/A	N/A	
SUMACLUST	PD	[106.00 - 162.78]	[142.85 - 174.19]	[89.22 - 89.22]	[0.93 - 39.47]	[0.59 - 1279.29]	[2.98 - 3.29]	
	Shannon	[2.00 - 3.01]	[2.19 - 3.28]	[5.48 - 5.48]	[0.16 - 7.43]	[2.32 - 10.32]	[1.00 - 8.03]	
	Simpson	[0.52 - 0.73]	[0.54 - 0.75]	[0.95 - 0.95]	[0.027 - 0.98]	[0.8 - 0.99]	[0.40 - 0.98]	
	F-Measure	0.40	0.43	0.53	N/A	N/A	N/A	
LSH-Canopy-SUMACLUST	ρ–value	0.9909	0.9950	0.9726	0.9575	0.7859	0.8812	
	PD	[112.36 - 177.37]	[157.55 - 183.41]	[91.96 - 91.96]	[0.74 - 39.51]	[0.86 - 1281.66]	[1.05 - 5.14]	
	Shannon	[3.15 - 4.02]	[2.6 - 3.65]	[5.77 - 5.77]	[0.18 - 7.92]	[2.52 - 11.62]	[2.73 - 9.91]	
	Simpson	[0.61 - 0.75]	[0.57 - 0.79]	[0.97 - 0.97]	[0.05 - 0.99]	[0.90 - 0.99]	[0.31 - 0.98]	
	F-Measure	0.47	0.48	0.56	N/A	N/A	N/A	
SWADM	PD	[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	[2.98 - 3.91]	[2.01 - 3.04]	[5.03 - 5.03]	[0.28 - 7.63]	[1.0 - 9.69]	[1.66 - 8.30]	
	Simpson	[0.70 - 0.82]	[0.53 - 0.74]	[0.95 - 0.95]	[0.05 - 0.98]	[0.5 - 0.99]	[0.00 - 0.99]	
LSH-Canopy-SWARM	F-Measure	0.54	0.55	0.56	N/A	N/A	N/A	
	ρ–value	0.9997	0.9980	0.9463	0.9769	0.7736	0.9136	
	PD	[17.53 - 25.79]	[17.48 - 25.92]	[32.17 - 32.17]	[1.34 - 27.65]	[0.10 - 745.40]	[2.17 - 7.41]	
	Shannon	[2.66 - 4.12]	[2.19 - 3.62]	[5.15 - 5.15]	[0.63 - 7.39]	[1.12 - 11.31]	[3.14 - 12.17]	
	Simpson	[0.76 - 0.86]	[0.49 - 0.78]	[0.97 - 0.97]	[0.08 - 0.99]	[0.64 - 0.99]	[0.38 - 0.99]	

Table shows values of F-measure for mock datasets for which taxonomy is known, Phylogeny Diversity (PD) value, Shannon and Simpson Coefficient intervals in the format [min - max]. Most of these datasets contain multiple samples and PD, Shannon and Simpson values are generated for each of these samples separately. So the values are represented as intervals from minimum to maximum that covers all performance metric values for the whole dataset regardless of number samples.

and finally utilize parallelism in multi-core CPU. We can see from Table III that LSH-Canopy-UCLUST outperforms UCLUST for all datasets. In fact LSH-Canopy-UCLUST took nearly half the time or less of naive UCLUST for all datasets. This improvement in runtime is due to efficient parallelization of expensive clustering in canopies. For Canadian Soil dataset which is the largest dataset used in this study and contains more than 9 million reads, UCLUST takes 72.47 minutes whereas LSH-Canopy-UCLUST takes only 41.92 minutes.

For SUMACLUST we can see from Table III that LSH-Canopy-SUMACLUST outperforms naive SUMACLUST in most cases. Major improvement in runtime was observed in large scale real world datasets specially Canadian Soil and Global Soil datasets. The highest amount of time taken by naive SUMACLUST is 510.92 minutes where LSH-Canopy-SUMACLUST performs a similar job in just 64.58 minutes. Which is almost 8 times faster than the naive version. But LSH-Canopy-SUMACLUST took more time than SUMACLUST for Bokulich_6 dataset which is the smallest dataset used in this study with only 250,903 number of reads. This gives the intuition that for relatively smaller datasets applying Locality Sensitive Hashing followed Canopy clustering might be unnecessary and may incur more time than the naive version of the clustering methods. A greedy clustering is enough and faster than a modified or added pipeline.

We can see a similar footprints for LSH-Canopy-

SWARM from Table III. SWARM took 128.12 minutes for Bokulich_2 dataset whereas LSH-Canopy-SWARM took 37.87 minutes only. SWARM was relatively faster for Bokulich_3, Bokulich_6 and Body Site datasets than SUMACLUST and LSH-Canopy-SWARM was faster than the corresponding naive version. But again major improvement in runtime was observed for larger real world datasets namely Canadian Soil and Global Soil datasets. LSH-Canopy-SWARM was nearly 3 times faster than naive SWARM for Canadian Soil dataset and ?? faster than Global Soil dataset. These observations refers to the fact that our proposed LSH-Canopy pipeline scale well with larger datasets and performs better in such case comparing to medium or smaller datasets.

C. Effect of Varying Number of Multi-processes

Figure 2a-2f shows how the number of multiprocess can affect runtime of the LSH-Canopy version of UCLUST, SUMACLUST and SWARM on 6 datasets that we have used in this study. We can see that increasing the number of multiprocess to perform expensive clustering inside canopies in parallel can significantly reduce total runtime. The steepest curve showing reduction in runtime can be found in Figure 2e for LSH-Canopy-SUMACLUST on Canadian Soil dataset. For single process version LSH-Canopy-SUMACLUST is slower in most cases except Bokulich_2 dataset where it's runtime is better comparing to SWARM with LSH-Canopy. For Body Site dataset (Figure 2d) the

Table III
RUNTIME COMPARISON (IN MINUTES)

	Datasets		Methods									
				UCLUST			SUMACLUST			SWARM		
Type	Title	# of Reads	UCLUST	with	Speed Up	SUMACLUST	with	Speed Up	SWARM	with	Speed Up	
				LSH-Canopy			LSH-Canopy			LSH-Canopy		
	Bokulich_2	6,938,836	10.08	4.82	2.09x	87.53	33.03	2.65x	128.12	37.87	3.38x	
Synthetic	Bokulich_3	3,594,237	7.85	3.65	2.15x	11.73	7.89	1.48x	9.10	6.93	1.31x	
	Bokulich_6	250,903	1.73	0.95	1.82x	1.28	1.34	0.96	1.97	1.29	1.53x	
	Body Sites	886,630	2.06	0.98	2.10x	15.42	7.15	2.16x	3.64	1.17	3.11x	
Real World	Canadian Soil	2,966,053	9.55	5.32	1.80x	363.96	79.45	4.58x	97.5	31.34	3.11x	
	Global Soil	9,252,764	72.47	41.92	1.73x	510.92	64.58	7.91x	269.07	53.71	5.00x	

LSH-Canopy version of SWARM and UCLUST have closer runtime with respect to increasing number of multiprocess. Figure 2f shows reduction in runtime for the largest dataset used in this study namely Global Soil with more than 9 million reads. Even though there are significant difference in the runtime of single process version, LSH-Canopy reduces runtime to closer interval for UCLUST, SWARM and SUMACLUST with increasing number of multiprocess.

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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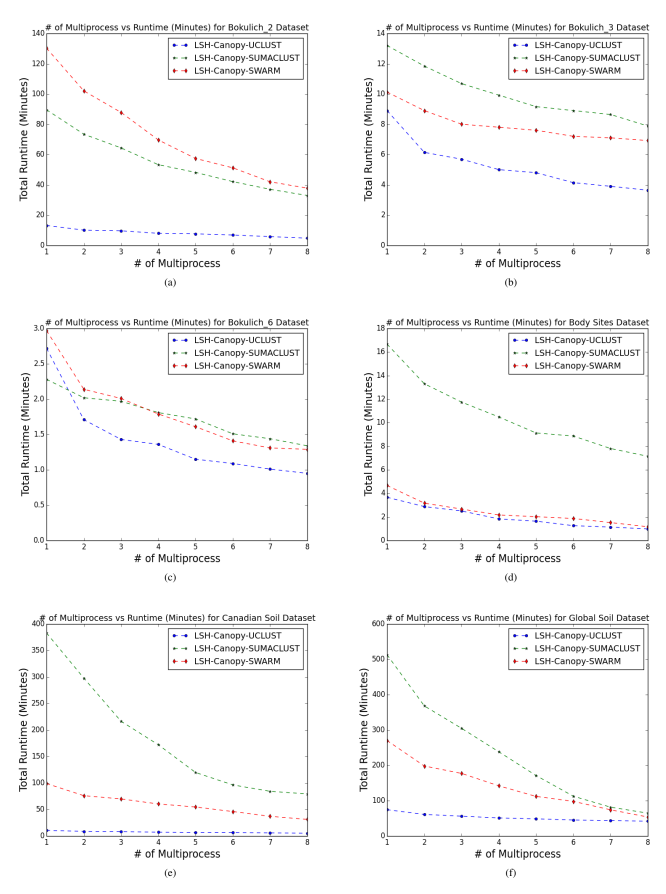


Figure 2. Runtime comparison of LSH-Canopy framework with UCLUST, SUMACLUST and SWARM on Bokulich_2, Bokulich_3, Bokulich_6, Body Sites, Canadian Soil and GLobal Soil datasets.