Efficient Approximate Maximum Inner Product Search over Sparse Vectors

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Abstract—The maximum inner product search (MIPS) problem in high-dimensional vector spaces has various applications, primarily driven by the success of deep neural network-based embedding models. Existing MIPS methods designed for dense vectors using approximate techniques like locality-sensitive hashing (LSH) have been well studied, but they are not efficient and effective for searching sparse vectors due to the nearorthogonality among the sparse vectors. The solutions to MIPS over sparse vectors rely heavily on inverted lists, resulting in poor query efficiency, particularly when dealing with largescale sparse datasets. In this paper, we introduce SOSIA, a novel framework specifically tailored to address these limitations. To handle sparsity, we propose the SOS transformation, which converts sparse vectors into a binary space while providing an unbiased estimator of the inner product between any two vectors. Additionally, we develop a minHash-based index to enhance query efficiency. We provide a theoretical analysis on the query quality of SOSIA and present extensive experiments on real-world sparse datasets to validate its effectiveness. The experimental results demonstrate its superior performance in terms of query efficiency and accuracy compared to existing methods.

Index Terms—Maximum Inner Product Search, Sparse Vectors

I. INTRODUCTION

In recent years, there has been a significant surge in the availability of high-dimensional vector data (i.e., vector representations of objects), primarily due to the proliferation of unstructured data and the remarkable success of deep neural network-based embedding models. These models have demonstrated their ability to embed various types of unstructured data, including videos [10], molecular structural information of drugs [8], and documents [29], into vectors. Subsequently, these vectors can be stored in a vector database to facilitate a wide range of applications, including personalized recommendations and information retrieval. As a result, the Maximum Inner Product Search (MIPS) problem, which aims to find a vector (i.e., a representation of an object) from a dataset D that has the maximum inner product with a given query q, plays a crucial role in these applications. As many vectors generated by embedding models exhibit sparsity (e.g., only 43 non-zero entries in the vector generated by the SPLADE model [15], [16]), the MIPS problem for sparse vectors has been widely studied [26], [37], [38], [47], [52], [54]. However, the "Dimensionality Curse" phenomenon makes exact MIPS computation

in high-dimensional spaces computationally expensive. In this paper, we study the problem of Approximate Maximum Inner Product Search (AMIPS) over sparse vectors, which identifies the top-k "approximately" most relevant objects for the query q over the collection D consisting of sparse vectors. Formally, given a query point $q \in \mathbb{R}^d$, a positive integer k, and an approximation ratio $c \in (0,1)$, the AMIPS over the sparse vectors D retrieves k points $x_i \in D$ $(1 \le i \le k)$ such that the inner product $q^{\top}x_i \ge c \cdot q^{\top}x_i^*$ where x_i^* is the i^{th} exact MIPS point of q over D.

The AMIPS problem with sparse vectors presents a significant challenge: most vectors are near-orthogonal, which makes it difficult for MIPS algorithms designed for dense vectors to effectively operate on sparse vectors. In contrast to methods for the MIPS problem in dense datasets, existing studies [2], [6], [7], [9], [13], [14], [20], [34] tackle the AMIPS problem for sparse datasets by considering only the non-zero indices and values of each sparse data. These studies directly organize this data using inverted index-based methods [6], [7], [13]. Several evaluation strategies [6], [34], [39] have been developed to filter out certain points by estimating the inner product of two vectors, aiming to reduce the number of inner product computations. However, most of these evaluation strategies are not computationally cheaper than computing the exact inner product, and all points that share common nonzero entries with the query must be considered, resulting in a high query cost of O(n) where n = |D|. Furthermore, these evaluation strategies often rely on the assumption that the vectors follow a specific distribution, which may not be applicable in the case of dynamic datasets commonly found in modern vector databases. Another limitation is that there is no theoretical guarantee that these methods offer a better query cost than O(n), which hampers their performance on large-scale datasets. As a result, designing an efficient and effective method for the AMIPS problem over large-scale and dynamic sparse datasets, without any underlying constraints or assumptions, remains a significant challenge.

To effectively address the AMIPS problem, we start by considering a simplified version of the problem on a binary-valued dataset in $\{0,1\}^{d'}$. This simplified version is equivalent to a set overlap search problem, where the overlap of two sets A and B (denoted as $|A \cap B|$) is equivalent to the inner product of their binary representations [45]. In particular, set overlap search is

a type of set similarity search problem that is related to other set search problems such as Jaccard similarity search [24], [25] and set containment search [1], [5]. These related problems have mature algorithms for approximate or exact solutions [42], [53], [56], [57], including inverted list-based methods [56] and minHash-based methods [57], which greatly improve the efficiency of set overlap search without any constraints on the dataset and enable sublinear cost solutions. With this in mind, we can explore transforming a real-valued sparse dataset into a binary-valued dataset while preserving the inner product relationships in their original space. Consequently, the AMIPS problem can be completely solved by addressing a set overlap search problem. While some sketch-based methods like BinSketch [39] attempt to transform a sparse vector in \mathbb{R}^d into a binary space for addressing the MIPS problem, they fail to preserve the inner product relationship without introducing significant distortion error.

Motivated by these observations, we propose a novel framework called SOSIA to address the AMIPS problem in a sparse vector space. In particular, we present the SOS (Sparse Vector to Set) transformation to convert the AMIPS problem on \mathbb{R}^d into the AMIPS problem on a binary space $\{0,1\}^{d'}$, which can be solved using minHash-based methods. The core of SOS is a transformation function $\mathcal{T}: \mathbb{R}^d \to \{0,1\}^{d'}$ that constructs a binary sketch for each vector in the dataset and query set. Theoretical results demonstrate that the inner product between $\mathcal{T}(x_1)$ and $\mathcal{T}(x_2)$ serves as an unbiased estimator of the inner product between x_1 and x_2 for any two points in \mathbb{R}^d . This property enables us to tightly control the error introduced by the transformation. Additionally, \mathcal{T} is a data-independent transformation and supports dynamic datasets. Moreover, SOS offers simplicity in deployment, making it compatible with disk-based, parallel, or distributed settings. This flexibility enhances its adaptability within modern vector databases, enabling efficient processing and storage. We further develop a novel minHash-based index, which can efficiently address the AMIPS problem in the transformed space and achieve high query performance in terms of efficiency and accuracy. In summary, we make the following contributions in the paper.

- We propose a novel framework called SOSIA to address the problem of AMIPS over sparse vectors. To address the challenges posed by sparsity, we develop the SOS transformation, which converts sparse vectors into a binary space while providing an unbiased estimator of the inner product between any two vectors.
- We further develop a novel minHash-based index that enhances the efficiency of answering queries in the AMIPS problem. This indexing method ensures a high query quality with a small query cost.
- We conduct extensive experiments on real-world sparse datasets to validate the effectiveness of SOSIA. The experimental results demonstrate that SOSIA outperforms competitors in terms of both query efficiency and accuracy.

The rest of the paper is organized as follows. Section II reviews related work on MIPS problem and set similarity

search problem. Section III presents the problem setting and basic concepts. Section IV shows the details of the SOS sketch and SOSIA framework. Section V provides proofs of lemmas and the theoretical guarantee of SOSIA. Section VI presents experimental studies and Section VII concludes the paper.

II. RELATED WORK

A. Maximum Inner Product Search over Dense Datasets

The MIPS problem has been extensively studied in the context of dense datasets. In low-dimensional spaces, exact solutions to the MIPS problem can be achieved using spacepartitioning trees, such as the M-tree [27] and cone-tree [40]. But their performance rapidly degrades as the dimensionality d increases due to the curse of dimensionality. In high-dimensional spaces, MIPS problem is mainly solved approximately by locality-sensitive hashing (LSH) methods and graph based method [36], [54]. LSH based methods are the mainstream approximation method for solving the NNS and MIPS problems [3], [12], [19], [26], [38], [44], [54]. Since the inner product is not a metric, asymmetric transformations are applied to convert the MIPS problem into NNS ahead [3], [44], [46]. Then, the MIPS problem is reduced to an NNS problem. However, these transformations either bring distortion errors or result in an imbalance in the transformed dataset, and thus limit the query performance. FARGO [54] proposes a novel transformation named random XBOX transformation that addresses these two issues. Graph-based [33], [36], [49] and learning-based [11], [17], [43] methods are also proposed for MIPS problem. However, both of these methods incur significantly high training or construction costs, which restrict their applicability to large-scale datasets.

These methods [11], [36], [38], [54] are mainly designed for dense vectors, the case where the entries of each vector are almost surely non-zero. In this case, the data points from a real-world dataset (not a random dataset) are usually well-clustered and the local intrinsic dimensionality (LID) [21], [31], [55] of the dataset are much less than its dimensionality d, which make it possible to either partition the nearby data points into several hash buckets or organize the datasets by proximity graphs. Unfortunately, when the points are sparse with very few non-zero entries, these methods hardly port over successfully because the points are near-orthogonal in a sparse high-dimensional space.

B. Maximum Inner Product Search over Sparse Datasets

Traditional approaches for addressing the MIPS problem in sparse datasets heavily rely on inverted index [6], [7], [13], [32], [34], [35]. The inverted index are previously designed for answering set similarity search, especially set overlap search. An inverted index consists of many posting lists where each posting list represents a specific entry of a vector and stores the point IDs and their non-zero values in this entry. By considering the inner product as a weighted set overlap, the problem can be answered exactly by traversing the posting lists of query's non-zero entries because the inner product is the sum of contributions in all entries. Such a naive approach

is simple but inefficient for large-scale datasets since nearly all data points are required to be computed in the worst case. Therefore, many evaluation strategies are designed for filtering some points by quickly computing their inner product upper bounds to the query. These evaluation strategies can be categorized into Document-at-a-time (DAAT) strategies and Term-at-a-time (TAAT) strategies. DAAT strategies, such as WAND, BMW, and VBMW [6], [13], [34], estimate the upper bound of the inner product of a single point appearing in the posting list of a query term before moving to the next point. The point whose upper bound is less than the current found largest inner product will be filtered out; otherwise, we compute its real inner product to the query and update the current largest inner product. The efficiency of these methods depends on how tight an upper bound is. WAND [6] adopts the maximal nonzero values in each posting list as an upper bound of this posting list. Then, the inner product upper bound of a point is estimated by accumulating the sum of upper bounds in the common posting lists of this point and query point. To obtain a tighter upper bound, BMW [13] and VBMW [34] organize the terms in a posting list by several smaller-size blocks and define an upper bound for each block rather than the entire posting list, which greatly reduces the difference between the upper bound and the exact values and thus make it possible to filter out more points. In addition, some methods represent sparse vectors by the sketches such that the inner product of sketches approximates the inner product of original vectors [2], [9], [20]. Then, we estimate the upper bounds of the inner product quickly by computing the inner product between two sketches. TAAT strategies, such as SINNAMON [7], process query terms one by one and accumulate a partial inner product as the contribution of each query term is computed. To obtain a tight estimate of the exact inner product and facilitate filtering some points, SINNAMON accesses the posting lists with the decreasing order of the query term's value since a large value has a higher impact on the inner product. When accessing the posting lists, SINNAMON also adopts a low-dimensional sketch like that in BinSketch [39] for estimating the inner product of two points by accessing only a part of posting lists. Although these methods effectively reduce the number of full computations, the computational cost associated with computing these upper bounds remains substantial, often comparable to that of computing the exact inner product. Moreover, in most cases, it is necessary to compute the upper bounds for all points that share common non-zero entries with the query, resulting in a query cost that is not significantly better than O(n).

III. PRELIMINARIES

In this section, we introduce the AMIPS problem and preliminaries. Frequently used notations are in Table I.

A. Problem Definition

Definition 1 (Maximum Inner Product Search). Given a dataset $\mathcal{D} = \{x_1, x_2, \cdots, x_n\} \subseteq \mathbb{R}^d$ with $|\mathcal{D}| = n$ and a

TABLE I: List of Key Notations.

Notation	Description				
\mathbb{R}^d	d-dimensional vector space				
$\mathcal D$	The dataset				
n	The cardinality of dataset				
x	A data point				
l	The base of SOS transformation				
x.ind	x's non-zero indices				
x.val	x's non-zero values				
q	A query point				
$q^{ op}x$	The inner product between q and x				
$\mathcal{N}_I(q)$	The point that has the largest inner product to q in \mathcal{D}				
x	The L2 norm of a point x in \mathbb{R}^d				
$ x , x_b $	The L1 norm of a point x in \mathbb{R}^d or a point x_b in $\{0,1\}^d$				
A, B	A set				

query point $q \in \mathbb{R}^d$, the maximum inner product search (MIPS) returns a point $\mathcal{N}_I(q) = x^* \in \mathcal{D}$ such that

$$\mathcal{N}_I(q) = \arg\max_{x \in \mathcal{D}} q^{\top} x.$$
 (1)

Definition 2 (Approximate Maximum Inner Product Search). Given a dataset $\mathcal{D} \subseteq \mathbb{R}^d$ with $|\mathcal{D}| = n$, a query point q, an approximation ratio $c \in (0,1)$ and a positive integer k, the approximate maximum inner product search ((c,k)-AMIPS) returns k points x_1, \ldots, x_k that are sorted in descending order k w.r.t. their inner products to k of k is the k-th maximum inner product of k in k it satisfies k is k-th maximum inner product of k in k-th satisfies k-th maximum inner k-th maximum inn

Remark 1. We denote c-AMIPS problem as the (c, k)-AMIPS problem at k = 1.

For a sparse vector $x \in \mathbb{R}^d$, we only record its non-zero information as a tuple $x = \langle x.ind, x.val \rangle$ where $x.ind = \{j \mid x[j] \neq 0\}$ is the non-zero indices and $x.val = \{x[j] \mid x[j] \neq 0\}$. For convenience, all indices begin at 0 rather than 1. We use x.nnz to denote the size of x.ind and x.val and use x.ind[j] and x.val[j] to denote the j-th non-zero entry in x, i.e., x[x.ind[j]] = x.val[j]. When x is a binary-valued data, it can be represented by only x.ind. Since x.ind is a set, we call the vector x as the binary representation of the set x.ind and call x.ind as the sketch of x. The inner product of two binary vectors is related to the set overlap.

Definition 3 (Set Overlap). The overlap of two sets A and B is the size of their intersection, i.e., $s_O(A, B) = |A \cap B|$.

We employ the following fact to establish the connection between inner product and set overlap.

Fact 1. For two sets $A, B \subset U = \{e_1, e_2, \dots, e_d\}$, the binary representation of A and B are two vectors $a, b \in \{0, 1\}^d$ and $a^{\top}b = |A \cap B|$.

To further illustrate the above conceptions, let us consider the example 1.

Example 1. Consider the dataset D shown in Fig. 1 where $D = \{x_0 = (0, 0.7, 0, 0), x_1 = (0, 0.2, 0, 0, 0.3), x_2 = (0, 0.5, 0, 0, 0), x_3 = (0.6, 0, 0.1, 0, 0.3)\} \subset \mathcal{R}^5$. x_2 can be represented as $x_1 = \langle [1, 4], [0.2, 0.3] \rangle$, where $x_1.ind = [1, 4]$ and $x_1.val = [0.2, 0.3]$. Similarly, a query point q = [0.2, 0.3]

(0,0.2,0,0,0.5) can be represented as $q = \langle [1,4], [0.2,0.5] \rangle$. The exact MIPS result for this query is x_1 ($q^{\top}x_1 = 0.19$). When c = 0.5 and k = 2, the (c,k)-AMIPS returns any two points in x_1, x_2, x_3 as correct results. Then, given a binary vector $x_b = \{0,1,0,0,1\}$, we can represent it as a set $\{[1,4]\}$.

Since a binary vector x is totally equivalent to the set A = x.ind. We do not distinguish between a binary vector and a set longer in this paper.

B. Jaccard Similarity and MinHash

Definition 4 (Jaccard Similarity). The Jaccord Similarity of two sets A and B is defined as $s_J(A, B) = |A \cap B|/|A \cup B|$.

A basic observation between Jaccard similarity and set overlap is

$$s_J(A,B) = \frac{s_O(A,B)}{|A| + |B| - s_O(A,B)},\tag{2}$$

Therefore, given the Jaccard similarity between two sets, we can efficiently estimate their overlap. Since Jaccard similarity is normalized to the range [0, 1], it is easier to estimate using techniques like MinHash, as compared to directly estimating set overlap.

Definition 5 (MinHash). Given a universal collection of sets U, a family of minHash functions $\mathcal{H} = \{h : U \to Z\}$ maps the members of U into integers such that for any two sets $A, B \in U$, the collision probability of A and B equals to their Jaccard similarity, i.e.,

$$\Pr[h(A) = h(B)] = s_J(A, B). \tag{3}$$

The goal of minHash is to estimate the Jaccard similarity between a query set and the set in a collection $\mathcal{D} \subset U$ quickly without explicitly computing the Jaccard. By adopting multiple minHash functions $h_0, h_1, \cdots, h_{m-1}$ from H, we can compute an unbiased estimation of the Jaccard similarity of two sets by counting the number of their collisions among m minHash functions.

C. Locality Sensitive Hashing index

When answering set Jaccard similarity search problem via a number of MinHash functions, a common strategy is to build Locality Sensitive Hashing (LSH) index. LSH function is a kind of hash functions based on specific metric, which is defined as:

Definition 6 (Locality Sensitive Hashing (LSH) [19], [51]). Given a metric space $\mathcal{M} = (X, dist)$ where X is a collection of points and dist is a function $dist: X \times X \to \mathbb{R}$, a distance r and an approximation ratio c > 1, a family of hash functions $\mathcal{H} = \{h: X \to Z\}$ is called (r, cr, p_1, p_2) -locality-sensitive, if for $\forall o_1, o_2 \in X$, it satisfies both conditions below:

- If $dist(o_1, o_2) \le r$, $Pr[h(o_1) = h(o_2)] \ge p_1$;
- If $dist(o_1, o_2) > cr$, $Pr[h(o_1) = h(o_2)] \le p_2$,

where $h \in \mathcal{H}$ is chosen at random, p_1, p_2 are collision probabilities and $p_1 > p_2$.

MinHash functions are locality-sensitive based on the Jaccard distance, which is defined as:

Definition 7 (Jaccard Distance [28]). The Jaccord distance of two sets A and B is $\delta_J(A, B) = 1 - s_J(A, B)$.

Jaccard distance is a metric that satisfies the triangle inequality [28]. In the metric space $\mathcal{M}=(U,\delta_J)$, a minHash function is (r,cr,1-r,1-cr)-locality-sensitive for any 0 < r < 1/c. We can adopt LSH index to answer an approximate Jaccard similarity search, like other nearest neighbor search (NNS), such as in Euclidean space. An LSH function is insufficient to answer NNS well since p_2 is too large in an LSH function such that there are many false negative points, *i.e.*, the points are too far away from the query, but still have the same hash value. (K,L)-LSH index is a common approach to address this problem [19], [51]. In the (K,L)-LSH index, to reduce the false negative ratio p_2 , we usually concatenate K LSH functions $h_0,h_1,\cdots,h_{K-1}\in\mathcal{H}$ as

$$g(o) = \langle h_0(o), h_1(o), \cdots, h_{K-1}(o) \rangle$$

and the points that agree on all K LSH functions are regarded as collisions and mapped into the same hash bucket. In the query phase, we only check the points that collide with the query point q. In this strategy, the false negative ratio are decreased as p_2^K , which greatly reduce the number of points to be checked. On the other hands, the true positive ratio are decreased from p_1 into p_1^K , which will also reduce the probability that a real nearest neighbor are found and degrade the query accuracy. A practical approach to address this issue is to repeat such a procedure L times independently to obtain enough candidate points in total L hash buckets. By properly choosing the values of K and L, an approximate nearest neighbor search can be solved in sublinear query cost with quality guarantee.

IV. THE SOSIA FRAMEWORK

In this section, we present our framework called SOSIA to address the (c, k)-AMIPS problem in a sparse vector space, as depicted in Fig 1. SOSIA involves several key steps. Firstly, SOSIA employs the SOS transformation (refer to Section IV-A) to convert points in the sparse dataset into binary vectors. These binary vectors are considered as sets, which are mapped into a hash bucket in each of m hash tables using m independent MinHash functions (refer to Section IV-B). When a query q is received, it is also transformed into a binary vector q_b using the SOS transformation. Subsequently, it performs a nearest neighbor search based on Jaccard distance by utilizing the hash tables to obtain T candidates for the query q (refer to Section IV-C). During the query processing, we notice that the existing minHash LSH indexes cannot handle the issue that the collision probability varies greatly with the query points so that some queries obtain too many candidates while other queries collide with a few points. To address this problem, a novel counting and refinement strategy is designed to find adaptively no more than T candidates for different query points efficiently.

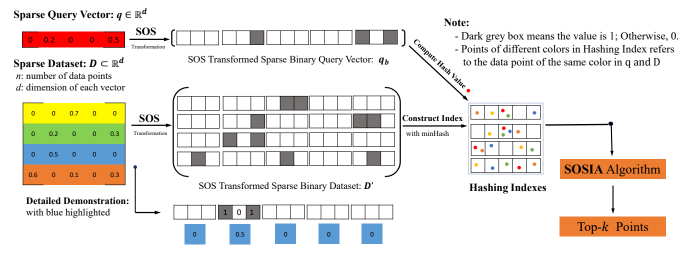


Fig. 1: SOSIA Framework

A. SOS Transformation

Given a base l, SOS transformation converts a point x in the dataset $\mathcal{D} \subset \mathbb{R}^d$ into a binary vector x_b in $\{0,1\}^{d'}$ where $d'=d\cdot l$, i.e.,SOS is a transform $\mathcal{T}:\mathbb{R}^{+d} \to \{0,1\}^{d'}$. Each entry x[i] in \mathbb{R}^d is mapped to l entries at the range $x_b[il:il+l-1]$ in $\{0,1\}^{d'}$ independently. The larger x[i] is, the more likely the entries of $x_b[il:il+l-1]$ are to be 1. To be simplified, we assume the maximal entries in \mathcal{D} is 1; otherwise, we normalized \mathcal{D} to make its maximal entries

$$M = \max_{x \in \mathcal{D}} \max_{0 \leq i < d} x[i]$$

to be 1, which does not effect the MIPS result of any query point. The detail of SOS is shown in Alg. 1. For a non-zero entry x[x.ind[j]] = x.val[j], each of l entries in $x_b[il:il+l-1]$ will be set as 1 with the probability x.val[j] (Line 5). The output of Alg. 1 is a binary vector in $x_b \in \{0,1\}^{d\cdot l}$. We take the datasets shown in Fig. 1 as an example to illustrate the details of the SOS transformation.

Algorithm 1: SOS Transformation

```
Input: A normalized sparse point x \in \mathbb{R}^d and a positive integer l

Output: A binary vector x_b \in \{0,1\}^{d \cdot l}

1 S \leftarrow \varnothing;

2 for j \leftarrow 0 to x.nnz do

3 for j \leftarrow x.ind[j] \cdot l to x.ind[j] \cdot l + l do

4 Genarate a random variable r from the uniform distribution U(0,1);

5 if r < x.val[j] then

6 S \leftarrow S \cup \{j\};
```

Example 2. Let us consider the SOS transformation with the base l=3 for the points shown in Fig. 1. We transform

 $x_0 = (0,0,0.7,0,0) \in \mathcal{D}$ into $\mathcal{T}(x_0) \in \{0,1\}^{15}$. Specifically, we loop only through non-zero entries $(x_0.ind)$ of x_0 , which is 2 in this simple case. In index 2 of x, for each of its three corresponding entries 6,7 and 8 in $\mathcal{T}(x_0)$, we generate a random variable r following an uniform distribution U(0,1). If the r is less than its entry value (0.7 in this case), we add the index to S, which is the output set containing all non-zero index for $\mathcal{T}(x)$. For example, if we generate three random numbers 0.8, 0.3 and 0.4 for these three indices 6,7,8, respectively. 7 and 8 will be put into S since 0.8 > 0.7, 0.3 < 0.7 and 0.4 < 0.7. Finally, the output of SOS is $\mathcal{T}(x_0) = \{7,8\}$, which is equivalent to the vector (0,0,0,0,0,0,1,1,0,0,0,0,0,0). Similarly, the point $x_1 = (0,0.2,0,0,0.3)$ can be transformed into $\mathcal{T}(x_1) = \{5,12,13\}$.

By adopting SOS to process the query q and each point x in \mathcal{D} , we can answer the AMIPS problem in \mathcal{D} by considering the AMIPS of $\mathcal{T}(q)$ in a binary-valued dataset $\mathcal{D}' = \{\mathcal{T}(x) | x \in \mathcal{D}\}$ since $\mathcal{T}(q)^{\top}\mathcal{T}(x)$ satisfies the following lemma:

Lemma 1. Given a query point q and a data $x \in \mathcal{D}$ where $q_b = \mathcal{T}(q)$ and $x_b = \mathcal{T}(x)$, then $q_b^{\top} x_b/l$ is an unbiased estimation of $q^{\top} x$.

The proof of this lemma is provided in Section V-A. This lemma guarantees that SOS will not incur any distortion error in terms of the inner product relationship of points in the original space, based on which we can estimate the value of $q^{\top}x$ as tight as desired, i.e., make $q_b^{\top}x_b/l$ be as close to $q^{\top}x$ as possible. Intuitively, enlarging l forces $q_b^{\top}x_b/l$ to be closer to $q^{\top}x$. We formulate this conclusion by computing the variance of $q_b^{\top}x_b/l$.

Lemma 2. The variance of $q_b^{\top} x_b/l$ is

$$\operatorname{Var}[q_b^{\top} x_b/l] = \frac{1}{l} \sum_{j=0}^{d-1} q[j]x[j](1 - q[j]x[j]).$$

The proof of this lemma is provided in Section V-A. Since the value of $\sum_{j=0}^{d-1}q[j]x[j](1-q[j]x[j])$ depends only on the

q and x themselves, the variance of $q_b^{\top} x_b / l$ can be reduced by enlarging l, which allows us to estimate the value of $q^{\top} x$ more tightly by choosing a larger base l.

Algorithm 2: Indexing in SOSIA

```
Input: A normalized sparse dataset \mathcal{D}, a query point q
             and an approximate ratio c
   Output: Hash Tables
1 Determine the value of m based on Eq. 4;
2 Generate m minHash functions h_1, h_2, \ldots, h_m;
\mathfrak{B}[1:m] \leftarrow \varnothing;
4 for x \in \mathcal{D} do
        x_b = \mathcal{T}(x);
        id \leftarrow x_b's point id;
        for j \leftarrow 0 to m-1 do
            Insert \langle h_j(x_b), \langle id, |x_b| \rangle \rangle into \mathcal{B}[j];
9 for j \leftarrow 0 to m-1 do
        for a hash bucket B in \mathcal{B}[j] do
10
             Sort the tuples \langle id, |x_b| \rangles in B by the
               descending order of |x_b|;
```

B. Indexing in SOSIA

12 return \mathcal{B} ;

After obtaining transformed dataset $\mathcal{D}' = \{x_b = \mathcal{T}(x) | x \in \mathcal{D}\}$, we choose m minHash functions and compute the hash values for each points x_b in \mathcal{D} . Then, m independent minHash indexes are built for the subsequent search in \mathcal{D}' . Unlike (K, L)-LSH indexes that build L numbers of K-dimensional hash tables, SOSIA builds m one-dimensional hash tables to facilitate determining the collision conditions based on different query points adaptively and dynamically. Algorithm 2 describes the indexing phase of SOSIA. First, we need to determine the number of hash functions, m, for answering the (c,k)-AMIPS problem. A larger m enables us to find higher-quality candidates but brings a higher query cost and space consumption. Based on our query strategy, we choose the smallest m that guarantees a correct (c,k)-AMIPS result with high probability, m is computed by

$$\begin{cases}
t = \left(\frac{\sqrt{c}+1}{2}\right)^2 \\
m = \frac{3c(1-c\gamma)(\sqrt{t/c}-t\gamma)^2}{\gamma(t-\sqrt{tc})^2} \cdot \ln\frac{2n}{T},
\end{cases}$$
(4)

where $\gamma \in (0,1)$ is a parameter to control the minimal inner product to be searched in our index and T is the maximal number of points to be checked. Such an m guarantees that SOSIA index returns a correct c-AMIPS with a constant probability 1/2 - 1/e when $q^{\top}x^* \geq \gamma(|q| + |x|)$ where $x^* = \mathcal{N}_I(q)$ is the maximal inner product result of q in \mathcal{D} , as analyzed in Section V-B. The reason why introducing a γ is as following: Let $s_I^* = q^{\top}\mathcal{N}_I(q)$ be the maximal inner product, when s_I^* is too small compared to the norm of q, any point in the dataset will not be similar to the query point q in

Algorithm 3: Search in SOSIA

```
Input: A dataset \mathcal{D}, the SOSIA index \mathcal{B}, T, c and k
    Output: At most k points in \mathcal{D}
 1 q \leftarrow \frac{q}{\max q[i]}, q_b \leftarrow \mathcal{T}(q);
 2 l \leftarrow the base of SOS; (see Alg. 1)
 3 Determine the value of t based on Eq. 4;
 4 I_0 \leftarrow \sqrt{\|q\|}, I_t \leftarrow 0, I \leftarrow I_0;
 5 Compute q_b's minHash values h_1(q_b), \ldots, h_m^*(q_b);
 6 H \leftarrow \varnothing, R \leftarrow \varnothing, cost \leftarrow 0;
 7 while I_t < cI and cost < T + k do
         x_b \leftarrow the first unseen point with largest norm in
           \mathcal{B}[j].at(h_i(q_b)), (1 \leq j \leq m);
         if no points is unseen then
 9
          Break;
10
         \alpha \leftarrow |\{j|h_j(q_b) = h_j(x_b)\}|;
11
        s_O \leftarrow \frac{|q_b| + |x_b|}{1 + m/\alpha};
12
13
14 while I_t < cI and cost < T + k do
15
         if H is empty then
          Break;
16
         \langle x, s_O \rangle \leftarrow H.top();
17
         if s_O/l < tI then
18
          I \leftarrow cI, continue;
19
         H.pop();
20
         x \leftarrow the point reprsented by x_b in \mathcal{D};
21
         I_x = q^{\top} x;
22
         cost \leftarrow cost + 1;
23
         e \leftarrow \{x, I_x\};
24
         R \leftarrow R \cup \{e\};
25
         if |R| = k + 1 then
26
              Remove the element with the smallest I_x in R;
27
              I_t \leftarrow \min\{e.I_x | e \in R\};
29 return R;
```

terms of the inner product, making it impossible to distinguish the points in the dataset via minHash. So, we only guarantee a correct (c,k)-AMIPS result when Then, m minHash LSH indexes are adopted for indexing the points x_b transformed by SOS. For a term $\langle key, value \rangle$ stored in a hash table, the key is the hash value of the point x_b and the value is a tuple consisting of the x_b 's identity and norm. Storing the norm $|x_b|$ is for restoring the value of $s_O(q_b, x_b)$ by $s_J(q_b, x_b)$ since minHash is designed for estimating $s_J(q_b, x_b)$. As such,

$$s_O(q_b, x_b) = \frac{|q_b| + |x_b|}{1 + 1/s_J(q_b, x_b)}.$$
 (5)

This equation also indicates that a larger $|x_b|$ are more likely to yield a larger $s_O(q_b, x_b)$. So, in each hash bucket *i.e.*, the hash terms with the same keys, we will sort the points by the descending order of their norms to quickly find the points with a large norm at first in the query phase, as shown in Line 11.

```
Algorithm 4: Update(x, s_O)
```

```
Input: x_b, q, l, t, I, cost, c, k, H, and R
   Output: k points in \mathcal{D}
 1 if s_O/l > tI then
         x \leftarrow the point reprsented by x_b in \mathcal{D};
 3
         I_x = q^{\top} x;
         cost \leftarrow cost + 1;
 4
         e \leftarrow \{x, I_x\};
 5
         R \leftarrow R \cup \{e\};
7 else
         x \leftarrow the point reprsented by x_b in \mathcal{D};
        H \leftarrow H \cup \{\langle x, s_O \rangle\};
10 if |R| = k + 1 then
         Remove the element with the smallest I_x in R;
         I_t \leftarrow \min\{e.I_x | e \in R\};
```

C. Search in SOSIA

When a query q comes, we also transform it into the binary vector q_b via SOS and then compute its m hash values. To find the candidate points with a large set overlap to q_b , a novel search algorithm based on collision counting is designed to quickly find enough number of high-quality candidates with a small size of indexes. The search strategy in SOSIA is quite different compared to the (K, L)-LSH indexes. In (K, L)-LSH indexes, there are L numbers of K-dimensional hash tables and the points that collide with the query point in at least one of L hash tables will be considered as a candidate and will be checked. For each data point, the collision condition in each K-dimensional hash tables is that its K hash functions are all evaluated to be the same with the hash value of query points. While such a strategy is efficient to answer an approximate nearest neighbor search problem in Euclidean space, it is inefficient and infeasible to answer an approximate Jaccard similarity problem due to the difference between the minHash function and the LSH functions under Euclidean space.

A typical LSH defined in the Euclidean space \mathbb{R}^d is as follows [12]:

$$h^{L2}(x) = \left| \frac{a^{\top} x + b}{w} \right|, \tag{6}$$

where a is a d-dimensional vector whose entries are chosen independently from the standard normal distribution, w is a pre-defined integer and b is a real number chosen uniformly from [0, w). The flexibility of such a hash function is that we can tune the collision probability for any two points by varying the value of w. A larger w enables a greater number of points agreeing on a hash value and enlarging the collision probability for any two points. Moreover, regardless of the distance between two points, they can be mapped into the same hash bucket by selecting a sufficiently large value for the parameter w. This enables LSH functions in Euclidean space to solve the approximate nearest neighbor search problem with varied distances between query points and their exact nearest neighbor.

However, these conditions are not satisfied by a minHash function. A minHash function h, defined for the a collections of sets \mathcal{D}' , is usually based on a random permutation of the elements of the universal set $U = \bigcup_{A \in \mathcal{D}'} A$. Let $P = \bigcup_{A \in \mathcal{D}'} A$ $\{e_1, e_2, \dots, e_d\}$ be a random permutation of U, h(A) is the minimal position of A's elements in P, i.e., $h(A) = \min\{j \mid$ $P[j] \in A$. Since no hyper-parameter like w in h^{L2} is used for tuning the collision probability of two points, we are unable to enlarge the collision probability of two sets when their Jaccard similarity is fixed. So, it becomes extremely difficult to choose a static K and L for varied query points. Specifically, denote $\mathcal{N}_{I}(q)$ as the set that has the maximal Jaccard similarity to the query q in the dataset \mathcal{D} , a large K enables us to find the good results for those queries q having a large $s_J(q, \mathcal{N}_J(q))$ but make the queries q having a small $s_J(q, \mathcal{N}_J(q))$ hard to find any candidate. While a small K enables us to find the good results for those queries q having a small $s_J(q, \mathcal{N}_J(q))$ but make the queries q having a small $s_J(q, \mathcal{N}_J(q))$ find too many candidates and increase the query cost. To address this dilemma, we adopt a dynamic counting based strategy for adaptively addressing varied query point.

Alg. 3 describes the search processing in SOSIA. Although the LSH method cannot solve a c-AMIPS problem directly, it can solve the problem by resolving a sequence of threshold-based (I,c)-MIPS problem by decreasing the inner product threshold I [23], [54]. Therefore, to answer a c-AMIPS problem, we execute a sequence of (I,c)-MIPS problems as defined below.

Definition 8 ((I, c)-MIPS [30]). Given a threshold I and an approximation factor c < 1, the (I, c)-MIPS for a query point q over dataset \mathcal{D} returns the following result:

- 1) If at least one point x exists in \mathcal{D} such that $q^{\top}x > I$, it returns such a point x' in \mathcal{D} that $q^{\top}x' > c \cdot I$;
- 2) If no point x exists in \mathcal{D} such that $q^{\top}x > c \cdot I$, it returns nothing.

So, we execute a sequence of (I,c)-MIPS problems by beginning with $I=\sqrt{\|q\|}$ (Line 4), which is the maximal value of $q^{\top}\mathcal{N}_I(q)$ since all entries have been normalized into [0,1] in \mathcal{D} . I_t is used for recording the current found k-th largest inner produce value. We initialize a max-heap H to store the all counted points and a min-heap R with size no more than k to store the found best k results so far. We divide the remaining search processing into two stages – the *counting stage* and *refinement stage*.

Counting stage (Lines 7-13). At this stage, we count the number of collisions between $q_b = \mathcal{T}(q)$ and points that lie in the same hash buckets as q_b . And in each hash bucket, we sort points in decreasing order by theirs' norm (Line 7). Since the points in a hash bucket has been sorted based on their norms, it only takes $O(\log n)$ time to obtain the first unseen point x_b that has the largest norm in q_b 's m hash buckets by maintaining a max-heap that stores the first unseen point with largest norm in q_b 's each hash bucket. For such a point x_b that collides α times with q_b , its estimated Jaccard similarity to q_b is α/m and we will compute its set overlap s_O to q_b based

on Eq. 5. Next, we updating the results by inserting x_b into either H or R based on the condition $s_O/l > tI$ where t is a threshold computed by Eq. 4. As shown in Lemma 3 defined below, s_O/l is an unbiased estimator of $q^{\top}x$. The proof of Lemma 3 is provided in Section V-B.

Lemma 3. Given a set space X and a family of MinHash functions $\mathcal{H} = \{h : X \to Z\}$, $h_0, h_1, \ldots, h_{m-1}$ are m min-Hash functions that are drawn independently from \mathcal{H} . For two points $q, x \in \mathbb{R}^d$, denote $\alpha = |\{j|h_j(\mathcal{T}(q)) = h_j(\mathcal{T}(x))\}|$. If $\alpha > 0$, then

 $s_O/l = \frac{|\mathcal{T}(q)| + |\mathcal{T}(x)|}{(1 + m/\alpha)l},\tag{7}$

is an unbiased estimator of $q^{\top}x$ where \mathcal{T} is the SOS transformation with the base l.

Alg. 4 describes the details of updating the results: When $s_O/l > tI$, $q^\top x$ is believed to be larger than cI with a high probability. We then compute the value of $q^\top x$ and put it in the result set R. Otherwise, $q^\top x$ is believed to be smaller than cI with a high probability and we insert it into H without computing $q^\top x$ in this stage. We keep record the k-th largest found result so far as I_t when R has more than k points. The counting stage terminates when one of the following three conditions satisfies:

- **T1.** All points, which lie in the same hash bucket (in each of m hash tables) as q_b , are counted (Line 9).
- **T2.** I_t is no smaller than cI (Line 7).

T3. There are T + k points x that $q^{\top}x$ is computed (Line 7). When T2 is satisfied, the algorithm successfully returns kcorrect (I, c)-MIPS results. Since the current $I = \sqrt{\|q\|}$ is the maximal possible inner product between q and any point in \mathcal{D} , the found results are correct (c, k)-AMIPS results. When T3 is satisfied, the algorithm has checked T+k points but less than k satisfied (I,c)-MIPS result are found. In this case, let I_k^* be the k-th exact largest inner product between q and the point in \mathcal{D} , I_k^* is believed to be less than I_t/c^2 with high probability, i.e., the algorithm has found correct (c^2, k) -AMIPS results. We will prove this statement in Section V-B. When T1 is satisfied, the algorithm neither checks enough number of candidates, i.e., T + k candidates, nor has found k correct (I, c)-MIPS result. In this case, we requires to conduct another (cI, c)-MIPS result with decreasing the threshold from I to cI in the next refinement stage.

Refinement stage (Lines 14-28). In this stage, we attempt to extract more candidates x from H whose estimated inner product to q, s_O/l , exceed cI. If no such point x exists but the termination conditions T2 and T3 are not satisfied, we keep decreasing the inner product threshold from I to cI. The counting stage terminates when T2, T3 and T4 satisfies. T2 and T3 are same as that in the counting stage and T4 is:

T4. The max-heap H becomes empty (Line 15).

When the *refinement stage* is terminated by **T4**, it indicates we fail to find enough number of candidates from m minHash indexes. In this case, it is uncertainty whether we have found correct (c^2, k) -AMIPS results. But by setting m based on

Eq. 4, we ensure this case will happen only when $I_k^* < \gamma \sqrt{\|I_0\|}$ where γ is a parameter defined in Eq. 4 with a high probability. In other word, when I_k^* is larger than this threshold, we would have found it with a high probability in one of two stages. When the *refinement stage* is terminated by **T2** or **T3**, we guarantee a correct (c^2, k) -AMIPS result as analyzed in the *counting stage*. The correctness of these statements will be proven in Section V-B.

Example 3. We take the dataset \mathcal{D} shown in Fig. 1 as an example to illustrate the indexing of SOSIA and how to answer a (2,0.5)-AMIPS in SOSIA. There are m=4 minHash functions h_0, \dots, h_3 that are designed for the transformed dataset $\mathcal{D}' = \{x_b = \mathcal{T}(x) | x \in \mathcal{D}\}$ and 4 hash tables are built based on their hash values. When the query q comes, we compute its hash values and find the points that collide with it among these 4 hash tables for answering an (I,c)-MIPS problem with $I = \sqrt{\|q\|} = 0.54$. In the counting stage, the first point to be checked is x_{b3} since it has the largest norm among all unseen points. To update the results in H and R, we find the collision number between q_b and x_{b3} is $\alpha = 1$ and the computed $s_O = 1.2$. Assume t = 0.8, $s_O/l = 0.4 \ge tI = 0.43$ is not satisfied and thus we store x_3 into H. Similarly, for the second point x_{b2} , $\alpha = 1$ and the computed s_O is 1.0. So, we store x_2 into H. For the third checked point is x_{b1} with $\alpha = 2$. The computed $s_O = 2.0$ and $s_O/l \ge tI$ is satisfied. We store x_1 into R. At this moment, all points that collide with q are counted and no enough point is found. Therefore, the query processing continues in the refinement stage with decreasing I from $I_0 = 0.54$ to $cI_0 = 0.27$. Then, x_3 are popped from H and we update it into $R = \{x_1\}$. Then, |R| = 2 and $I_t = \min\{q^{\top}x_1, q^{\top}x_3\} = 0.15$. Hence, $I_t < cI$ is violated and the query terminates based on the condition **T2**. The returned points are x_1 and x_3 , which are the correct (2,0.5)-AMIPS for q.

V. THEORETICAL ANALYSIS

In this section, we provide the proofs of lemmas in Section IV and then demonstrate the quality guarantee of SOSIA.

A. Some Proofs

We first prove Lemma 1 that reveals $q_b^{\top} x_b / l$ is an unbiased estimator of $q^{\top} x$.

Proof. The expectation of $q_b^{\top} x_b/l$ is

$$E[q_b^{\top} x_b/l] = 1/l \cdot E\left[\sum_{j=0}^{dl-1} q_b[j] x_b[j]\right]$$
$$= 1/l \cdot E\left[\sum_{j=0}^{d-1} \sum_{m=0}^{l-1} q_b[jl+m] x_b[jl+m]\right].$$

Since all the entries of q_b and x_b are set as 1 independently, we have:

$$E[q_b^{\top} x_b/l] = 1/l \cdot \sum_{j=0}^{d-1} \sum_{m=0}^{l-1} E[q_b[jl+m]] E[x_b[jl+m]].$$

 $q_b[jl+m]=1$ incurs when the random number $r \sim U(0,1)$ is less than q[j]. Since $0 \leq q[j] \leq 1$, $\mathrm{E}[q_b[jl+m]] = \mathrm{Pr}[q_b[jl+m]=1] = q[j]$. Similarly, $\mathrm{E}[x_b[jl+m]] = x[j]$. So,

$$\mathrm{E}[q_b^\top x_b/l] = 1/l \cdot \sum_{j=0}^{d-1} \sum_{m=0}^{l-1} q[j]x[j] = \sum_{j=0}^{d-1} q[j]x[j] = q^\top x,$$

which indicates $q_b^{\top} x_b / l$ is an unbiased estimation of $q^{\top} x$. \square

Then, we prove Lemma 2.

Proof. The variance of $q_b^{\top} x_b/l$ is

$$Var[q_b^{\top} x_b/l] = 1/l^2 \cdot Var \left[\sum_{j=0}^{dl-1} q_b[j] x_b[j] \right]$$
$$= 1/l^2 \cdot \sum_{j=0}^{d-1} \sum_{m=0}^{l-1} Var \left[q_b[jl+m] x_b[jl+m] \right].$$

The variable $q_b[jl+m]x_b[jl+m]$ follows a binomial distribution and $\Pr[q_b[jl+m]x_b[jl+m]=1]=\Pr[q_b[jl+m]=1]$ $\Pr[x_b[jl+m]=1]=q[j]x[j]$. So, $\operatorname{Var}[q_b[jl+m]x_b[jl+m]]=q[j]x[j](1-q[j]x[j])$ and

$$\operatorname{Var}[q_b^{\top} x_b/l] = 1/l^2 \cdot \sum_{j=0}^{d-1} \sum_{m=0}^{l-1} q[j]x[j](1 - q[j]x[j])$$
$$= 1/l \cdot \sum_{j=0}^{d-1} q[j]x[j](1 - q[j]x[j]).$$

Next, we prove Lemma 3 to demonstrate that we can obtain an unbiased estimator of q^Tx by using minHash.

Proof. Let ν_j be a 0-1 random variable and $\nu_i=1$ indicates $h_j(\mathcal{T}(q))=h_j(\mathcal{T}(x)).$ Then, we have $\mathrm{E}\,\nu_j=\mathrm{Pr}[\nu_j=1]=\mathrm{Pr}[h_j(\mathcal{T}(q))=h_j(\mathcal{T}(x))]=s_J(\mathcal{T}(q),\mathcal{T}(x)).$ Since $\alpha=\sum_{j=0}^{m-1}\nu_j$ and ν_1,\cdots,ν_j are independent, $\mathrm{E}\,\alpha=\sum_{j=0}^{m-1}\mathrm{E}\nu_j=m\cdot s_J(\mathcal{T}(q),\mathcal{T}(x)).$ And $\mathrm{E}\left[\frac{1}{(1+m/\alpha)l}\right]=\frac{1}{(1+m/\mathrm{E}\,\alpha)l}=(1+1/s_J(\mathcal{T}(q),\mathcal{T}(x)))/l.$ Hence,

$$E[s_O/l] = E\left[\frac{|\mathcal{T}(q)| + |\mathcal{T}(x)|}{E[(1+m/\alpha)l]}\right]$$
$$= 1/l \cdot E\left[\frac{|\mathcal{T}(q)| + |\mathcal{T}(x)|}{1+1/s_J(\mathcal{T}(q), \mathcal{T}(x))}\right]$$
$$= E\left[\frac{s_O(\mathcal{T}(q), \mathcal{T}(x))}{l}\right],$$

which equals to $q^{\top}x$ based on Lemma 1.

B. Theoretical Guarantee of SOSIA

To demonstrate the reasonableness to adopt $s_O/l \ge tI$ as the condition for generating the candidates (Line 1 in Alg. 4), we provide the following lemma:

Lemma 4. Given a query q, a point x, a threshold $0 < \gamma < 1$, m minHash functions, an inner product threshold I, an approximation ratio 0 < c < 1, and a parameter c < t < 1, let

 $q_b = \mathcal{T}(q)$ and $x_b = \mathcal{T}(x)$. We define $s_O(q_b, x_b) = \frac{|q_b| + |x_b|}{1 + m/\alpha}$, where α is the counting numbers between q_b and x_b in m minHash functions. For the following two events:

- **E1:** If $q^{\top}x \geq I$, then $s_O(q_b, x_b) > ltI$,
- E2: The number of point x, where $s_O(q_b, x_b) > ltI$ but $q^\top x < cI$, is less than T

by setting t and w based on Eq. 4 and setting $l = \frac{4 \cdot \max\{3c,2\}}{(1-\sqrt{c})^2} \cdot \ln n$, the probability that E1 occurs is at least 1-1/e, and the probability that E2 occurs is at least 1/2 when $q^{\top} \mathcal{N}_I(q) \geq \gamma(|q| + \mathcal{N}_I(q))$.

Proof. (Sketch.) We prove that $\Pr[s_O(q_b, x_b) > ltI \mid q^\top x < cI] < \frac{T}{2n}$ according to the Chernoff bound ¹. Then the expected number of points x, which satisfies $s_O(q_b, x_b) > ltI$ but $q^\top x < cI$, is $N_f \le n \cdot \frac{T}{2n} = T/2$. Based on the Markov's inequality, the probability that E2 occurs is at most $1 - T/N_f = 1/2$. Similarly, the probability that E1 occurs can be directly bounded by the Chernoff bound. Due to the space limitation, we provide the detailed proof in our technical report [41].

Theorem 1. When $q^{\top} \mathcal{N}_I(q) \geq \gamma(|q| + \mathcal{N}_I(q))$, by setting parameters as that in Lemma 4, Alg. 3 returns a correct c^2 -AMIPS result with a constant probability of at least 1/2-1/e.

Proof. We prove the theorem by assuming E1 in Lemma 4 holds, whose probability is at least 1-1/e. When $q^{\top}\mathcal{N}_I(q) \geq$ $\gamma(|q| + \mathcal{N}_I(q))$, the probability that $\mathcal{N}_I(q)$ does not collide with q in m hash tables is at most 1 - 1/e, based on E1 in Lemma 4. In this case, Alg. 3 will not terminate by **T4**. If Alg. 3 terminates by **T2** $(I_t \ge cI)$ when conducting (I_1, c) -MIPS search, it indicates the last round of (I, c)-MIPS with $I = I_1/c$ does not find correct results. So, $q^{\top} \mathcal{N}_I(q) < I_1/c$ based on E1 in Lemma 4; otherwise, $\mathcal{N}_I(q)$ must have been found when conducting (I_1, c) -MIPS search. In addition, since $I_t \geq cI_1 \geq c^2 \mathcal{N}_I(q)$, a correct c^2 -AMIPS result have been found. If Alg. 3 terminates by **T3** when conducting (I_1, c) -MIPS search, it also indicates $q^{\top} \mathcal{N}_I(q) < I_1/c$, based on E1 in Lemma 4. In addition, T + k number of points x fulfill $s_O(q_b, x_b) > lt \cdot I_1$ but none of them satisfies $q^{\top}x > cI$, which conflicts with E2. Since $Pr[E2] \ge 1/2$, the probability that Alg. 3 terminates by T3 is at most 1/2.

C. Query Time and Space Consumption

As shown in Eq. 4 and Lemma 4, $m = O(\log n)$ and $l = O(\log n)$. So the extra space consumption for storing the transformed dataset and minHash indexes are $O(n\log n)$. For the query time, the inner product of at most T+k points are computed with respect to q, where T is a constant. Each inner product computation cost O(q.nnz) time, so the computing cost is O(q.nnz). In the worst case, all points are counted O(m) times, thus the counting cost is $O(nm) = O(n\log n)$. Therefore, the total query time is $O(n\log n + q.nnz)$.

¹https://en.wikipedia.org/wiki/Chernoff_bound

TABLE II: Summary of Datasets

Datasets	n	φ_d	φ_q	d	Size (GB)
SPLADE1M*	1,000,000	126.3	49.1	30,109	0.97
SPLADE-Full*	8,841,823	126.8	49.1	30,109	8.42
Rand1M [†]	1,000,000	127.3	49.0	30,000	0.97
Rand10M [†]	10,000,000	127.8	49.0	30,000	9.52

^{*}Real-world Datasets; †Synthetic Datasets

Remark 2. Although the query time of SOSIA is $O(n \log n + q.nnz)$, which seems to be worse than a brute force method. However, such a cost is the bound of the worst query complexity and in practice, the query time is still dominated by the time of computing the real inner products, i.e., the computation cost. Fortunately, the computation cost is bounded by O(q.nnz), much better than O(n). Many LSH methods, under Euclidean space, have O(n) or even $O(n \log n)$ query time, such as C2-LSH [18], QA-LSH [22], SRS [48] and PM-LSH [55], but still outperform many other LSH methods that have a sublinear query cost, such as LSB-Forest [50]. We will further demonstrate the efficiency of SOSIA in the next section by experiments.

VI. EXPERIMENTAL STUDY

In this section, we conduct extensive experiments on real world data and synthesized datasets for a comprehensive evaluation and analysis on SOSIA. We implement the SOSIA ² and competitors in C++ and compiled with g++ using -Ofast optimization. All experiments are conducted on a Ubuntu server with 4 Intel(R) Xeon(R) Gold 6218 CPUs (160 threads) and 1.5 TB RAM.

A. Experimental Settings

<u>Datasets.</u> Four sparse datasets are used in our experiments, as shown in Table II. The former two datasets are obtained by embedding MS Marco Passage Retrieval v1 dataset [4] via the SPLADE model [15], [16], since the SPLADE model has been proven to be the most promising sparse embedding model for document retrieval. SPLADE1M contains 1M points chosen randomly from SPLADE-Full. The latter two datasets Rand1M and Rand10M are randomly generated based on the datasets SPLADE-Full. In Table II, φ_d and φ_q are the average number of non-zero values in the dataset and query set, respectively. Compared to the dimensionality d, φ_d and φ_q are much smaller.

Competitors. To demonstrate the indexing and query performance of our SOSIA Framework, we compare it with the WAND [6] and the SINNAMON [7]. WAND is the most representative inverted index based method using DAAT (Document-at-a-time) query strategy. For the SINNAMON [7], proposed by Pinecone, it is a good representative for TAAT (Term-at-a-time) strategy-based method, where it utilizes BinSketch [39] to derive the approximation of inner products. Therefore, we choose WAND and SINNAMON as our competitors.

Parameter Settings. We consider the (c,k)-AMIPS queries with c=0.5 and k=50 for all algorithms as the default settings. Following settings in the paper or source code of our competitors, we adopt the fixed parameters on each algorithm to facilitate using them. The parameter settings of all algorithms are as follows: For SOSIA, the base of SOS, l=40, the number of minHash functions m=150, the maximal number of points to be checked, T=10000 for small datasets (SPLADE1M, Rand1M), and T=20000 for large datasets (SPLADE-Full and Rand10M). For SINNAMON, the sketch dimensionality $m=0.25\varphi_d$, the number of mappings h=2, and the time budget T is set as twice of the running time of SOSIA. For WAND, we set the factor F=2.0.

Evaluation Metrics. We evaluate the performance of our SOSIA framework and the compipitors with respect to three aspects: index size, query efficiency, and query accuracy. Specifically, we use the running time to evaluate indexing time and query time; and the recall is used to measure the query accuracy. For an algorithm, given a query q, we denote the result of a (c, k)-AMIPS as $R = \{x_1, x_2, \cdots, x_k\}$. Given a query q, denote $R^* = \{x_1^*, x_2^*, \cdots, x_k^*\}$ as the result of exact k MIPS. The recall are computed as following:

$$Recall = \frac{|R \cap R^*|}{|R^*|} \tag{8}$$

B. Self Evaluation

In this subsection, we demonstrate the effects of the base of SOS, l, and the number of minHash functions, m, on the query performance of our framework SOSIA.

- 1) Parameter study on l: To further study the SOSIA framework, we evaluate the impact of l, the base of SOS transformation, on the query performance of SOSIA by setting l in the range of $\{1,10,20,40,80\}$. We show our results on SPLADE1M and SPLADE-Full datasets for brevity. As shown in Fig. 2, the query time of SOSIA remains nearly unchanged as l increases. The reason is that l does not affect the collision probability of two transformed vectors and thus does not affect the query time. As for the recall, it keeps increasing with l, since a larger recall brings us a better estimation for the exact inner product based on Lemma 2 and thus enhance the query accuracy. While the marginal increase of recall is small after l=40, we choose l=40 as the default value.
- 2) Parameter study on m: In this experiment, we evaluate the effect of the m, the number of hash functions, on the query performance of SOSIA by setting m in the range of $\{25, 50, 100, 150, 200\}$ on the dataset SPLADE1M and SPLADE-Full. As shown in Figure 3, the query time continuously increases when we increase m on both datasets. This is because a larger m requires a longer time for counting the collision numbers. Also, the recall increases as m increases, as more minHash functions can yield a more accurate estimation on Jaccard similarity between two sets and thus yield a more accurate estimation on inner product. The marginal increase of the recall is small after m=150, while the query time would increase significantly, we choose m=150 as the default value.

²https://github.com/Jacyhust/SOSIA

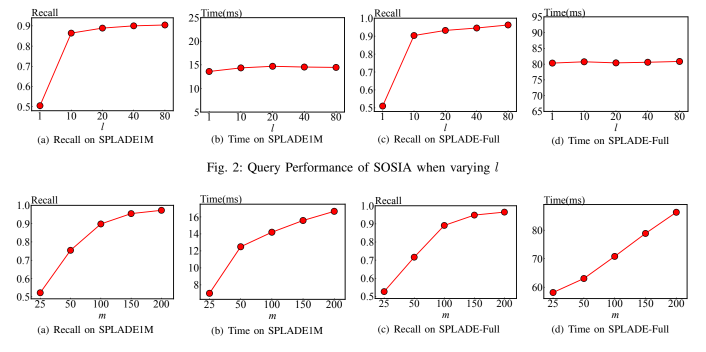


Fig. 3: Query Performance of SOSIA when varying m

TABLE III: Performance Overview

		SOSIA	SINNAMON	WAND
	Time (ms)	15.58	33.28	91.69
SPLADE1M	Recall	0.9548	0.9526	0.9498
	Index Size (GB)	1.1	1.2	0.94
	Time (ms)	82.41	180.2	778.7
SPLADE-Full	Recall	0.9414	0.8538	0.9658
	Index Size(GB)	9.9	10.3	8.4
Rand1M	Time (ms)	21.37	42.77	86.04
	Recall	0.9492	0.6284	0.5386
	Index Size (GB)	1.1	1.2	0.95
	Time (ms)	137.3	289.6	849.3
Rand10M	Recall	0.9168	0.3428	0.435
	Index Size (GB)	11.2	11.8	9.5

C. Performance Overview

In this experiment, we provide an overview of the average query time, recall and index size of all algorithms with default parameter settings on all datasets, as shown in Table III. When comparing the query performance of the different algorithms, we observe that SOSIA consistently achieves the highest recall on most of datasets, except for SPLADE-Full where it is slightly lower than WAND. Additionally, SOSIA exhibits significantly faster query times compared to its competitors. In fact, the query time of SOSIA is smaller half of that of the second-best algorithm. The reason is that SOSIA only considers points with a large collision number to the query, which greatly reduces the cost of exactly computing inner products. When comparing the index size of the different algorithms, we found that all algorithms have the close index size and SINNAMON always has the largest one. This is because SINNAMON requires storing an inverted list and nsketches, where the length of the sketch for each point is $m \approx 30$. Compared to WAND that only stores an inverted list, SINNAMON has a larger index size. The index size of SOSIA depends on the number of hash tables m. Each hash table only stores a point once and the total index size is close to the size of dataset itself.

D. Evaluation of Query Performance

1) Effect of k: We compare the query performance of all algorithms as k varies in the range $\{1,10,20,\ldots,100\}$, and present the results in Figures 4. As k increases, the recall of SOSIA and SINNAMON tend to decrease and their query time remains steady. The reason is that SOSIA and SINNAMON check nearly the same number of points for answering (c,k)-AMIPS at different k. So, k has little impact on the query time. However, the query accuracy of SINNAMON decreases rapidly as k increases, indicating the scoring strategy in SINNAMON has a limited performance for a large k. On the contrary, the recall and query time of WAND increases with k. This is because the pruning condition in WAND is to compare the upper bound of a point with the current found k-th largest results. A larger k enables a smaller number of points to be pruned and thus brings a higher recall and query time.

2) Recall-Time Curve: In these experiments, we analyze the trade-off between recall and query time through the Recall-Time curve for all algorithms. The algorithm with the smaller query time for a given target recall is considered to have better query performance. The results on all datasets are shown in Figure 5. From the figures, we make the following observations: (1) Among all algorithms, SOSIA spends the shortest query time to reach the same recall, indicating that it achieves the best trade-off between query quality and efficiency. The reason is that by using SOS transformation and

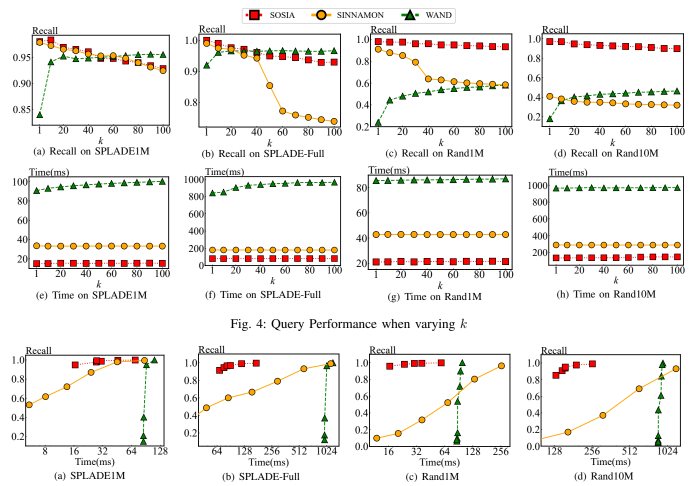


Fig. 5: Recall-Time Curves

minHash indexes, SOSIA has a better estimator than SINNA-MON and WAND. For SINNAMON, the cost of completely scoring a vector x is even higher than directly computing $q^{\top}x$. Although the scoring phase can be terminated if the time budget T is exhausted, the query accuracy decreases significantly while using the partial scores for ranking and it is difficult to find a proper T. The efficiency of WAND depends on the number of pruned points. When the pruning ratio is low, WAND has a large query time. (2) As query time increases, all algorithms achieve higher recall, which aligns with the principle of AMIPS methods, where query accuracy is traded for query efficiency. Additionally, the ratio recall/timedecreases greatly as the query time increases, especially when the recall is close to 1. This reveals the difficulty and computationally expensive nature of finding exact MIPS results. (3) When comparing the recall-Time curves among these four datasets, we find the difference of query performance between SOSIA and the other two algorithms varies greatly. On SPLADE1M and SPLADE-Full datasets, SOSIA achieves about 2-4× speedup of query time to reach the same recall compared to the second best algorithm. While on Rand1M and Rand10M datasets, the speedup reaches 5-10×, indicating the competitors performs poorly on the random datasets. The reason is that the estimating accuracy and pruning efficiency of SINNAMON and WAND is affected by the data distribution. Moreover, the speedup of SOSIA to its competitors on large dataset such as SPLADE-Full and Rand10M is much larger than that on the small dataset, enhancing the adaptability of SOSIA on large-scale datasets.

VII. CONCLUSION

In this paper, we propose a novel transformation called SOS, which converts sparse vectors into binary vectors while providing an unbiased estimator of their inner product. Building upon SOS, we design an approach named SOSIA that utilizes minHash to address the MIPS problem over sparse vectors. SOSIA incorporates a counting-based query strategy, enabling the efficient and adaptive generation of candidates for different query points. This approach allows us to achieve high query accuracy while greatly reducing computational costs. Theoretical analysis demonstrates that SOSIA guarantees a correct *c*-AMIPS result with constant probability by verifying only a limited number of points. Extensive experiments reveal that SOSIA outperforms its strongest competitor, SINNAMON, by achieving a query time speedup of 2-10× while maintaining the same recall.

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APPENDIX I: PROOFS

In this the appendix, we provide the detailed proofs of Lemma 4 that establishes the theoretical basic of SOSIA. Before proving it, we first introduce some required conclusions:

Lemma 5. Given a query q and a point x, let $q_b = \mathcal{T}(q)$ and $x_b = \mathcal{T}(x)$, we denote α_i as an indicator of the event that q_b and x_b collide in i-th minHash functions occurs, i.e., $\alpha_i = \Pr[h_i(q_b) = h_i(x_b)].$ Then, $\mathbf{E} \alpha_i = \frac{q^\top x}{|q| + |x| - q^\top x}.$

Proof. We have

$$\alpha_i = \Pr[h_i(q_b) = h_i(x_b)] = s_J(q_b, x_b)$$
$$= s_J(q_b, x_b) = \frac{q_b^{\top} x_b}{|q_b| + |x_b| - q_b^{\top} x_b}.$$

So,

$$\begin{split} \mathbf{E} \, \alpha_i &= \mathbf{E} \left[\frac{q_b^\top x_b}{|q_b| + |x_b| - q_b^\top x_b} \right] \\ &= \frac{\mathbf{E}[q_b^\top x_b]}{\mathbf{E}[|q_b|] + \mathbf{E}[|x_b|] - \mathbf{E}[q_b^\top x_b]} \end{split}$$

 $\mathrm{E}[|q_b|] = \mathrm{E}[\sum_{j=0}^{dl-1} x_b[j]] = l \cdot |q|$ and $E[|x_b|] = l \cdot |x|$. Based on Lemma 1, $\mathrm{E}[q_b^{\top} x_b] = l \cdot q^{\top} x$. Therefore, $\mathrm{E} \, \alpha_i = \frac{q^{\top} x}{|q| + |x| - q^{\top} x}$

Lemma 6 (Chernoff Bound ³). Let X_1, X_2, \ldots, X_n be independent 0-1 random variables and $X = \sum_{i=1}^n X_i$ with

- For any $0 \le \epsilon \le 1$, $\Pr[X \le (1 \epsilon)\mu] \le \exp(-\frac{\mu\epsilon^2}{2})$. For any $0 \le \epsilon \le 1$, $\Pr[X \ge (1 + \epsilon)\mu] \le \exp(-\frac{\mu\epsilon^2}{3})$.

Based on the Chernoff bound, we have the following corollary:

Corollary 1. By setting the base of SOS $l = \frac{4 \cdot \max\{3c, 2\}}{(1 - \sqrt{c})^2} \cdot \ln n$, given a query q and a point x, let $q_b = \mathcal{T}(q)$ and $x_b = \mathcal{T}(x)$, we denote $L_b = |q_b| + |x_b|$ and $L_0 = |q| + |x|$, then $l \cdot \sqrt{t} L_0 \le$ $L_b \leq l \cdot \sqrt{t/cL_0}$ holds with the probability at least 1 - 2/nwhere $t = (1 + \sqrt{c})/2$.

Proof. Since $E[L_b] = l \cdot L_0$ and L_b is the sum of $2d \cdot l$ 0-1 random variables $q_b[i]$ s and $x_b[i]$ s, we have the following bounds for L_b :

•
$$\Pr[L_b \le l \cdot \sqrt{t}L_0] \le \exp(-\frac{l \cdot L_0(1-\sqrt{t})^2}{2}).$$

• $\Pr[L_b \ge l \cdot \sqrt{t/c}L_0] \le \exp(-\frac{l \cdot L_0(\sqrt{t/c}-1)^2}{3}).$

As q has been normalized to make $\max\{q[i]\} = 1, L_0 \ge |q| \ge$ 1, then $\Pr[L_b \leq l \cdot \sqrt{t}L_0] \leq 1/n$ and $\Pr[L_b \geq l \cdot \sqrt{t/c}L_0] \leq 1/n$ when $l = \frac{4 \cdot \max\{3c,2\}}{(1-\sqrt{c})^2} \cdot \ln n$. Proven!

³https://en.wikipedia.org/wiki/Chernoff_bound

We prove the Lemma 4 based on the conclusions above:

Proof. Based on the definition of $s_O(q_b, x_b)$,

$$\Pr[s_O(q_b, x_b) > ltI] = \Pr[\alpha > \frac{m \cdot ltI}{L_b - ltI}]$$

where $L_b = |q_b| + |x_b|$.

Proving E1. For E1, when $q^{\top}x=I$, $\operatorname{E}\alpha_i=\frac{q^{\top}x}{|q|+|x|-q^{\top}x}=\frac{I}{L_0-I}$ where $L_0=|q|+|x|$. Consider $I\geq \gamma L_0$, $\mu_1=\operatorname{E}\alpha=\operatorname{E}[\sum_{i=0}^{m-1}\alpha_i]=\frac{mI}{L_0-I}\geq \frac{m\gamma}{1-\gamma}$. We denote ϵ_1 to make

$$(1 - \epsilon_1)\mu_1 = \frac{m \cdot ltI}{L_b - ltI}. (9)$$

Then, based on Corollary 1, $L_b \ge l \cdot \sqrt{t} L_0 \ge lt L_0$ and thus

$$0 \le (1 - \epsilon_1)\mu_1 \le \frac{m \cdot ltI}{ltL_0 - ltI} = \frac{mI}{L_0 - I} = \mu_1, \tag{10}$$

indicating $0 \le \epsilon_1 \le 1$. According to the Chernoff bound (Lemma 6),

$$p_{f1} = \Pr[\alpha \le \frac{m \cdot ltI}{L_b - ltI}] = \Pr[\alpha \le (1 - \epsilon_1)\mu_1]$$
$$\le \exp(-\frac{\mu_1 \epsilon_1^2}{2}).$$

Based on Eq. 9,

$$\begin{split} \epsilon_1 = & 1 - \frac{m \cdot ltI}{L_b - ltI} \cdot \frac{1}{\mu_1} \\ \geq & 1 - \frac{m \cdot ltI}{l \cdot \sqrt{t}L_0 - ltI} \cdot \frac{1 - \gamma}{m\gamma} \\ = & \frac{\sqrt{t} - t}{\sqrt{t} - \gamma t}. \end{split}$$

$$\begin{split} \frac{\mu_1 \epsilon_1^2}{2} &\geq \frac{(\sqrt{t} - t)^2}{2(\sqrt{t} - \gamma t)^2} \cdot \frac{m\gamma}{1 - \gamma} \\ &\geq \frac{t(1 - \sqrt{t})^2}{2(\sqrt{t} - \gamma t)^2} \cdot \frac{3c\gamma(1 - c\gamma)(\sqrt{t/c} - t\gamma)^2}{\gamma(1 - \gamma)(t - \sqrt{tc})^2} \cdot \ln \frac{2n}{T} \\ &= \frac{3(1 - \sqrt{t})^2(1 - c\gamma)(\sqrt{t} - t\gamma\sqrt{c})^2}{2(1 - \gamma)(1 - \gamma\sqrt{t})^2(t - \sqrt{tc})^2} \cdot \ln \frac{2n}{T} \\ &= \frac{3(1 - c\gamma)(1 - \gamma\sqrt{tc})^2}{2(1 - \gamma)(1 - \gamma\sqrt{tc})^2} \cdot \ln \frac{2n}{T} , (1 - \sqrt{t} = \sqrt{t} - \sqrt{c}) \\ &\geq \frac{3}{2} \cdot \ln \frac{2n}{T} > 1 , (0 \leq c \leq 1) \end{split}$$

So, $p_{f1} \leq \exp(-\frac{\mu_1 \epsilon_1^2}{2}) < 1/e$. Therefore, the probability that E1 occurs is at least $1-p_{f1}=1-1/e$.

Proving E2. For E2, when $q^{\top}x = cI$, $\operatorname{E}\alpha_i = \frac{q^{\top}x}{|q| + |x| - q^{\top}x} = \frac{cI}{L_0 - cI}$. Consider $I \geq \gamma L_0$, $\mu_2 = \operatorname{E}\alpha = \operatorname{E}[\sum_{i=0}^{m-1}\alpha_i] = \frac{mcI}{L_0 - cI}$. We denote ϵ_2 to make

$$(1 + \epsilon_2)\mu_2 = \frac{m \cdot ltI}{L_b - ltI}.$$
 (11)

Then, based on Corollary 1, $L_b \leq l \cdot \sqrt{t/c}L_0$ and thus

$$(1 + \epsilon_1)\mu_2 \ge \frac{m \cdot ltI}{l \cdot \sqrt{t/c}L_0 - ltI},\tag{12}$$

Similarly, $0 \le \epsilon_2 \le 1$. According to the Chernoff bound (Lemma 6),

$$p_{f2} = \Pr[\alpha \ge \frac{m \cdot ltI}{L_b - ltI}] = \Pr[\alpha \ge (1 + \epsilon_2)\mu_2]$$

$$\le \exp(-\frac{\mu_2 \epsilon_2^2}{3}).$$

Based on Eq. 11,

$$\begin{split} \epsilon_2 \mu_2 = & \frac{m \cdot ltI}{L_b - ltI} - \mu_2 \\ \geq & \frac{m \cdot ltI}{l \cdot \sqrt{t/c}L_0 - ltI} - \frac{mcI}{L_0 - cI} \\ = & \frac{m(t - \sqrt{tc})IL_0}{(\sqrt{t/c}L_0 - tI)(L_0 - cI)} \end{split}$$

So,

$$\frac{\mu_2 \epsilon_2^2}{3} = \frac{(\mu_2 \epsilon_2)^2}{3\mu_2} \ge \frac{m(t - \sqrt{tc})^2 I L_0^2}{3c(\sqrt{t/c}L_0 - tI)^2 (L_0 - cI)}$$
$$\ge \frac{\gamma(t - \sqrt{tc})^2}{3c(1 - c\gamma)(\sqrt{t/c} - t\gamma)^2} m$$
$$\ge \ln \frac{2n}{T}$$

So, $p_{f2} \leq \exp(-\frac{\mu_2 \epsilon_2^2}{3}) \leq \frac{T}{2n}$. Then the expected number of points x, which satisfies $s_O(q_b, x_b) > ltI$ but $q^\top x < cI$, is $N_f \leq n \cdot \frac{T}{2n} = T/2$. Based on the Markov's inequality, the probability that E2 occurs is at most $1 - T/N_f = 1/2$. \square