

Project Report

An Implementation Of: A Simple Randomized $O(n \log n)$ -Time Closest-Pair Algorithm in Doubling Metrics

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1 Introduction

Implementation of an algorithm helps us observe its efficiency and behavior in practice. In this report, I will briefly explain each part of the closest-pair doubling algorithm [1], show the program's implementation along with practical running time analysis, and some implementation techniques I used. The theoretical information in this report fully refers to the work of A. Maheshwari, W. Mulzer and M. Smid, see [1].

Given a metric space $(P, dist)$, with doubling dimension d , whose P is a set of n points. The closest pair of points is the two points with the distance δ_0 that satisfies $dist(p_1, p_2) \geq \delta_0$ for any point $p_1, p_2 \in P$. Also, the doubling dimension d of a metric space indicates that for every point p in P and every real number $R > 0$, the $ball_P(p, R)$ can be covered by at most 2^d balls in P of radius $R/2$, see [1, Section 2]. By using the definition and properties of the doubling metric space and its doubling dimension, the algorithm will find the closest-pair distance without the direct use of the points' coordinates in $O(n \log n)$ time.

The closest-pair algorithm consists of three smaller parts:

1. Computing a separating annulus, denoted $SEPANN(S, n, d, \mu, c)$
2. The refinement of $SEPANN(S, n, d, \mu, c)$, denoted $SPARSESEPANN(S, n, d, t)$
3. The main recursive closest-pair algorithm, denoted $CLOSESTPAIR(S, n, d)$

Throughout the paper, let:

- $(P, dist)$ be a finite metric space in which P is the set of all points, and $dist$ is the function that calculate the distance between any two points
- d be the space's doubling dimension
- S be a non-empty subset of P

Note: I will only mention 2D points because of the extremely high running time of the algorithm in the space more than 3D (in which the doubling dimension is approximately at least $\log_2 21$ that gives us the base case with more than 3 000 000 points).

2 Algorithm 1: Computing a separating annulus

An important part of the main closest-pair algorithm is finding a separating annulus in the subset S . I will briefly describe this algorithm in the next subsection, due to A. Maheshwari, W. Mulzer and M. Smid [1, Section 3.1].

2.1 The $\text{SEPANN}(S, n, d, \mu, c)$ algorithm

In this section, $\mu \geq 1$ is a real constant number, c is calculated based on μ (I would say that $c = 2(4\mu)^d$ [1, Remark 1] since μ is not an integer in this case [1, Section 3.2]).

This algorithm picks a uniformly random point p from the subset S then finds the smallest ball centered at p , denoted $\text{ball}_S(p, R_p)$, that contains at least n/c point. If the outer ball $\text{ball}_S(p, \mu R_p)$ contains at most $n/2$ points, it returns p and R_p . If not, this procedure is repeated until the condition is satisfied. This algorithm's pseudocode is given below, see [1, Section 3.1].

<p>Algorithm 1: $\text{SPARSESEPANN}(S, n, d, t)$</p> <p>Input: Let S be a subset of P, of size n, d be the metric space's doubling dimension, $\mu \geq 1$ and $c > 1$ be large enough real numbers.</p> <p>Output: A point $p \in S$ and a radius $R_p > 0$.</p> <p>repeat</p> <p> $p =$ a uniformly random point in S;</p> <p> $R_p = \min\{r > 0 : \text{ball}_S(p, r) \geq n/c\}$;</p> <p>until $\text{ball}_S(p, \mu R_p) \leq n/2$</p> <p>return p and R_p</p>

This pseudocode is from [1, Section 3.1]

2.2 Finding the K^{th} smallest element

One step in $\text{SEPANN}(S, n, d, \mu, c)$ algorithm is to find the smallest ball which contains at least n/c points. This ball is easy to find using the k^{th} smallest element algorithm. Particularly, in a list of distances between p and all other points in S , we pick the $\lceil n/c \rceil$ -th smallest element and let it be the radius of the ball we need to find. Thus, all points closer to p are inside this ball.

A very easy approach to find the k^{th} smallest element in a list is to sort it in ascending order, and then simply return the element at the k^{th} place. This sorting algorithm takes $O(n \log n)$ time complexity in the worst case. Fortunately, we can improve the running time to $O(n)$ using a common recursive technique called QuickSelect, which is similar to QuickSort.

Given an unordered list D which contains n numbers, and a positive integer k satisfies $1 \leq k \leq n$. Each element in D has an index from 0 to $n - 1$. Consider a sublist of D , denoted $D[a : b]$, which starts from index a and ends at b , inclusively. The base case is when $a = b$, the algorithm returns the only element in that sublist. If it is not the case, the algorithm will choose a random *pivot* in the list. The algorithm then rearranges the list so that all elements

smaller and larger than the pivot are to the left and right of it, respectively. Now, the pivot has a new index, says c . If $k = c - a + 1$, the chosen pivot is the k^{th} smallest element of D , the algorithm returns $D[c]$. If $k < c - a + 1$, the algorithm recurses on the sublist to the left of the pivot, $D[a : c - 1]$. If $k > c - a + 1$, the algorithm recurses on the sublist to the right of the pivot, $D[c + 1 : b]$, with a different input k (which is $k - c + a - 1$) since this is the k of the subset to the right, not the whole set of points, and the elements' indices starts from 0 again when we recurse. This algorithm's pseudocode is given below.

<p>Algorithm: KTHSMALLEST(D, a, b, k)</p> <p>Input: Let D be a list of double numbers, the integers a and b respectively be the starting and ending indices of a sublist and k be an integer refers to the k^{th} smallest element.</p> <p>Output: The k^{th} smallest element in D, a double number.</p> <pre> if $a == b$ then return $D[i]$ else p = a random element in D; rearranging: all elements smaller than p to the left of p; all elements larger than p to the right of p; end; c = the current index of p in D; if $k == c - a + 1$ then return p else if $k < c - a + 1$ then return KTHSMALLEST($D, a, c - 1, k$) else return KTHSMALLEST($D, c + 1, b, k - c + a - 1$) end end </pre>
--

Unlike the original QuickSort algorithm, the QuickSelect recurses only once and on one side after rearranging. This helps the algorithm remain $O(n)$ time complexity. By using the random selection, the pivot on average is close to the middle of the list. Therefore, with the input is an n -sized list D , the recursive call is on a sublist whose size is a half. Because the rearrangement takes $O(n)$ time, this algorithm takes at most $2n$ time which is $O(n)$. The running time is shown below:

$$O(n) + O(n/2) + O(n/4) + \dots \leq O(2n) = O(n)$$

The following image is one of many outputs using different numbers of elements and k values. I generated 100 000 000 numbers uniformly random in the range $[-100000, 100000]$ with $k = 76,543,210$.

```

Number of elements: 100,000,000
k: 76,543,210
Value range: -100,000 to 100,000
- O(nlogn) Result Using Sort: 53081.606002
  Time taken: 53.615235s

- O(n) Result Using QuickSelect: 53081.606002
  Time taken: 3.588446s

```

Figure 1: The outputs of the $O(n \log n)$ algorithm using regular sorting and the $O(n)$ QuickSelect algorithm that recurses only once.

With the given number of elements, k and value range, both of the algorithms produced the same result, which is 53081.606002, but the running times of them have a significant difference (For the program's output, see Figure 1).

Since the QuickSelect takes at most $2n$ running time, the regular sorting takes at least $\log(n)/2$ times more than it, and the log is in base 2. In this case, there are 100 000 000 numbers, so:

$$\frac{\log(n)}{2} = \frac{\log(100\,000\,000)}{2} \approx 13.2877 \text{ times} \leq \frac{53.615235}{3.588446} \approx 14.9411 \text{ times}$$

Since 14.9411 is quite close to 13.2877, the running time of this QuickSelect algorithm in practice is reasonable compared to the theory.

3 Algorithm 2: The refinement of $\text{SEPANN}(S, n, d, \mu, c)$

With the use of the algorithm $\text{SEPANN}(S, n, d, \mu, c)$'s output, this refinement algorithm continues to find an annulus that separates the points in S into different smaller sets of points.

Let $\mu = e$ and $c = 2(4e)^d$, see [1, Remark 1]. The algorithm $\text{SEPANN}(S, n, d, e, c)$ returns an annulus's center $p \in S$ and its radius, $R' > 0$. An input of this refined algorithm is $t > 0$, a large enough constant calculated based on n . Let $R_i = (1 + 1/t)^i \cdot R'$, in which i is a uniformly random element inclusively from 1 to t . The algorithm then finds an annulus centered at p with the inner radius R_{i-1} and outer radius R_i , denoted $A_i = \text{annulus}_S(p, R_{i-1}, R_i)$. If the annulus contains at most n/t points then we are done. The algorithm returns p and R_{i-1} . This algorithm has the expected $O(cn)$ time complexity, see [1, Section 3.2]. Otherwise, the previous procedure is repeated until the condition is satisfied. The pseudocode of this refined algorithm is given below.

Algorithm 2: SPARSESEPANN(S, n, d, t)

Input: Let S be a subset of P , of size $n \geq 2(4e)^d + 1$, d be the space's doubling dimension, $t \geq 1$ be an integer.

Output: A point $p \in S$ and a radius $R > 0$.

$c = 2(4e)^d$;

let $p \in S$ and $R' > 0$ be the output of algorithm SEPANN(S, n, d, e, c);

repeat

i = a uniformly random element in $\{1, 2, \dots, t\}$;

$R_i = (1 + 1/t)^i \cdot R'$;

$R_{i-1} = (1 + 1/t)^{i-1} \cdot R'$;

$A_i = \text{annulus}_S(p, R_{i-1}, R_i)$;

$s = |A_i|$;

until $s \leq n/t$;

$R = R_{i-1}$;

return p and R

This pseudocode is from [1, Section 3.2]

4 Algorithm 3: The main closest-pair algorithm

Using the result of SPARSESEPANN(S, n, d, t), this main algorithm CLOSESTPAIR(S, n, d) recursively compute the closest distance.

Let S be an n -sized subset of P , d be the metric space (P, dist) 's doubling dimension. In the base case, that is when $n < 2(16e)^d$, the algorithm uses brute-force to compute the closest distance. If this is not the case, the algorithm sets $t = \lfloor \frac{1}{16e}(n/2)^{1/d} \rfloor$. This algorithm then runs SPARSESEPANN(S, n, d, t) with this t value, and gets the output $p \in S$ and $R > 0$ which is the inner radius of the sparse annulus. The outer radius of it is $(1 + 1/t)R$.

Let S_1 be the set of points contained in $\text{ball}_S(p, R)$, S_2 be the set of points contained in $\text{annulus}_S(p, R, (1 + 1/t)R)$ and S_3 be the set that contains all the points outside of the outer radius $(1 + 1/t)R$. This algorithm now recursively calls itself on two subsets of S which are points contained in $\text{ball}_S(p, (1 + 1/t)R)$ (which is $S_1 \cup S_2$) and outside of $\text{ball}_S(p, R)$ (which is $S_2 \cup S_3$). Finally, the smaller distance output from two recursive calls, $\delta_0 = \min(\delta_1, \delta_2)$, is returned. Below is the algorithm's pseudocode, see [1, Section 4.1].

Algorithm 3: CLOSESTPAIR(S, n, d)**Input:** Let S be a subset of P , of size $n \geq 2$, d be the space's doubling dimension.**Output:** A real number δ_0 satisfies the two properties in [1, Lemma 5].

```

if  $n < 2(16e)^d$  then
  |  $\delta_0$  = the closest-pair distance in  $S$  using brute-force;
else
  |  $t = \lfloor \frac{1}{16e}(n/2)^{1/d} \rfloor$ ;
  | let  $p \in S$  and  $R > 0$  be the output of algorithm SPARSESEPANN( $S, n, d, t$ );
  |  $S_1 = \text{ball}_S(p, R)$ ;
  |  $S_2 = \text{annulus}_S(p, R, (1 + 1/t)R)$ ;
  |  $S_3 = S \setminus (S_1 \cup S_2)$ ;
  |  $n' = |S_1 \cup S_2|$ ;
  |  $n'' = |S_2 \cup S_3|$ ;
  |  $\delta' = \text{CLOSESTPAIR}(S_1 \cup S_2, n', d)$ ;
  |  $\delta'' = \text{CLOSESTPAIR}(S_2 \cup S_3, n'', d)$ ;
  |  $\delta_0 = \min(\delta', \delta'')$ ;
end
return  $\delta_0$ 

```

This pseudocode is from [1, Section 4.1]

5 The Implementation

This implementation of the closest-pair doubling algorithm is written in C++ since it is a very common and fast programming language with high-level supports of object-oriented programming that can help us organize the program efficiently (see the implementation's *GitHub repository*¹ for the source code). The implementation and its components will be described and explain throughout this section. For some small class, I will combine both class code and implementation code into one block in this report.

To begin with, a set of points, which does not necessarily refer to **set** in Computer Science, is always one of the input parameters of each partial algorithm. In this implementation, I would choose **vector**, a very popular built-in data structure of C++. Since its size can automatically change when we add or remove an element, it is easy to work with and manage its behavior when implementing. Not only for storing points, **vector** will be used throughout the implementation to store some other information.

5.1 Utility functions and classes

5.1.1 Random number generator classes

Due to A. Maheshwari, W. Mulzer and M. Smid, the closest-pair doubling algorithm uses randomized technique in some parts to find the targets in linear time. Therefore, generating

¹GitHub repository of the implementation. <https://github.com/ThangMinhCao/closestpairdoubling>

random numbers is important. I created two random number generator classes, one for integer and one for double, to make the program more simple. The generators' code is given below.

```

1 #include <random>
2
3 class RandomInt{
4     public:
5         RandomInt(int start, int end)
6             : generator{std::random_device{}()}, int_dist(start, end) {}
7         int next() { return int_dist(generator); }
8
9     private:
10        std::mt19937 generator;
11        std::uniform_int_distribution<> int_dist;
12 };
13
14 class RandomDouble{
15     public:
16         RandomDouble(int start, int end)
17             : generator{std::random_device{}()}, double_dist(start, end) {}
18         double next() { return double_dist(generator); }
19
20     private:
21        std::mt19937 generator;
22        std::uniform_real_distribution<> double_dist;
23 };

```

Source ²: master branch → *closestpairedoubling/include/random_generator.h*

For each class, I create a uniform distribution with a given range that corresponds to the number type and an `mt19937` random generator with a `random_device` input. These two variables will be initialized when a random generator object is instantiated. Each class also has a `next()` function that can be called to return a random number.

5.1.2 *Point* and *PointList* classes

Since points are fundamental elements of the algorithm, and they are not just numbers (they have their own coordinates), creating a `Point` class is quite essential. Moreover, with the addition of the class function `distance_to`, the algorithm can calculate the distance between any two points easily without directly using the points' coordinates. This function works as the input function *dist* that is provided with the metric space $(P, dist)$. The implementation code of the class is given below.

²Path to the source file in the implementation's GitHub repository.

```

1 #include <vector>
2 #include <cmath>
3
4 class Point {
5     private:
6         std::vector<double> coordinate;
7     public:
8         explicit Point(std::vector<double> coordinate) {
9             this->coordinate = std::move(coordinate);
10        }
11        Point()= default;
12        // distance is calculated using Pythagorean Theorem
13        double distance_to(const Point& another_point) {
14            double total_square = 0;
15            for (int i = 0; i < coordinate.size(); i++) {
16                double iDiff = coordinate[i] - another_point.getCoordinate()[i];
17                total_square += (iDiff) * (iDiff);
18            }
19            return sqrt(total_square);
20        }
21        std::vector<double> getCoordinate() const {
22            return coordinate;
23        }
24    }

```

Source: master branch → *closestpairst doubling/include/point.h*

Because there is not only one method to generate points (see the description later in Section 6), so if all of them are implemented in the `main()`, the function will be quite huge and complicated. Thus, I created another utility class to make the code more organized, `PointList`. The `PointList` class consists of a `vector` named *points* that store all the points in a subset of P and some point generating functions. This class is used as subsets of P and to initialize all the points in P at the start of the program.

```

1 #include "Point.h"
2 #include "RandomGenerator.h"
3
4 class PointList {
5     public:
6         std::vector<Point> points;
7
8         // initializes all points uniformly random, see Section 6 for details
9         void random_initializer(int dimension, int point_num,
10                                int coor_start_range, int coor_end_range)

```



```

11 {
12     RandomDouble random_double(coor_start_range, coor_end_range);
13     for (int i = 0; i < point_num; i++) {
14         std::vector<double> coor;
15         // number of coordinates is the dimension value
16         for (int j = 0; j < dimension; j++) {
17             coor.push_back(random_double.next());
18         }
19         points.push_back(Point(coor));
20     }
21 }
22
23 // initializes points in grid, see Section 6 for details
24 void grid_initializer(double dist, int xFactor, int yFactor, int
position)
25 {
26     // the grid starts at coordinates (position, position)
27     int y = 0;
28     while (y < yFactor) {
29         int x = 0;
30         while (x < xFactor) {
31             std::vector<double> coor;
32             coor.push_back(position + (x++) * dist);
33             coor.push_back(position + y * dist);
34             points.push_back(Point(coor));
35         }
36         y++;
37     }
38 }
39 }

```

Source: master branch → `closestpairedoubling/include/point_list.h`

5.1.3 K^{th} smallest element finder

The K^{th} smallest algorithm has its own class. In this class, I define a shorter type name `DVect` which is `std::vector<double>`. The QuickSelect algorithm is divided into three functions, the main is `get`, and two assistances `swap` and `partition`. The function `get` returns the k^{th} smallest element using QuickSelect, and the other using the normal sort. Although I only use the QuickSelect technique, I create both to compare their time complexity, see Section 2.2.

```

1 #include <vector>
2
3 typedef std::vector<double> DVect;
4
5 class KthSmallest {
6     private:
7         static void swap (double *a, double *b);
8         static int partition(DVect& distances, int start, int end);
9
10    public:
11        // get k-th smallest element with QuickSelect
12        static double get(DVect& distances, int start, int end, int k);
13        // get k-th smallest element with normal sorting
14        static double get_with_sorting(DVect distances, int k);
15 };

```

Source: master branch → closestpairedoubling/include/kth_smallest.h

Due to its simplicity and irrelevant uses (only for testing and analyzing), the implementation of the normal sorting technique will not be listed in this report, see the *GitHub repository* for the code. The following is the implementation code of the QuickSelect algorithm.

```

1 #include "KthSmallest.h"
2 #include "RandomGenerator.h"
3
4 void KthSmallest::swap (double *a, double *b) {
5     double temp = *a;
6     *a = *b;
7     *b = temp;
8 }
9
10 int KthSmallest::partition(DVect& distances, int start, int end) {
11     double pivot = distances[end];
12     int slow = start;
13     int fast = start;
14     // moving the smaller elements to the left and larger elements to the
15     // right
16     while (fast < end) {
17         if (distances[slow] > pivot) {
18             while (fast < end and distances[fast] > pivot) {
19                 fast++;
20             }
21             if (distances[fast] < pivot) {
22                 swap(&distances[slow], &distances[fast]);

```

```

22     } else {
23         break;
24     }
25 }
26 fast++;
27 slow++;
28 }
29 // swapping the pivot from end back to its original place
30 swap(&distances[slow], &distances[end]);
31 return slow;
32 }
33
34 double KthSmallest::get(DVect& distances, int start, int end, int k) {
35     if (start == end) {return distances[start];}
36     // swapping the random pivot to the end
37     RandomInt random_int_gen = RandomInt(start, end);
38     swap(&distances[random_int_gen.next()], &distances[end]);
39     int cur_pivot = partition(distances, start, end);
40     if (k == cur_pivot - start + 1) {
41         return distances[cur_pivot];
42     } else if (k < cur_pivot - start + 1) {
43         return get(distances, start, cur_pivot - 1, k);
44     } else {
45         return get(distances, cur_pivot + 1, end, k - cur_pivot + start - 1);
46     }
47 }

```

Source: master branch → closestpairstoubling/src/kth_smallest.cpp

The function `swap(double *a, double *b)` swaps two elements in a vector by swaps the values at the addresses that the pointers `*a` and `*b` point to. Thus, their values in a vector are swapped while their identities are still the same.

The function `partition(DVect& distances, int start, int end)` takes the last element in `distances` to be the pivot. Then, it rearranges the vector so that every element smaller and larger than pivot is to the left and right of it. The technique I use in this function is creating two pointers `fast` and `slow`. Both pointers move towards the end step by step together until `distances[slow] > pivot`, then it stops. The `fast` keep moving until `distances[fast] < pivot`, then the value at two indices are swapped. This procedure is repeated until the `fast` reach the end of the vector. The `pivot` is now swapped back to the `slow` position, where it is supposed to be.

A generic call is `get(DVect& distances, int start, int end, int k)`. It first checks if the sub-vector has only one element (as well as `start == end`) then returns it. Otherwise, it selects a random pivot in the vector then swaps it with the last element. Now, the `partition` function is called to return the index of the pivot after rearranging. The

base case is when there are exactly k elements from index 0 to the current pivot in the original vector (which is `cur_pivot - start + 1`), so it is the k^{th} smallest element. If there are more than k elements, the function is recursively called on the sub-vector to the left of the pivot. If there are less than k elements, the function is called on the sub-vector to the right, but now the k should be decreased by `cur_pivot - start + 1`. Due to Section 2.2, this procedure takes $O(n)$ time complexity.

5.1.4 The 2D algorithm for testing

In the project, I also implement the 2D closest-pair algorithm using Divide-and-conquer algorithm due to Bentley and Shamos, see [5]. This algorithm is very helpful in testing and debugging the closest-pair doubling algorithm because of its short running time. I will briefly describe the algorithm below.

Given a set of point P , the base case is when P contains less than 2 points. If there is only one, the algorithm returns positive ∞ . If there are two points, the algorithm basically returns the distance between them. Otherwise, P is sorted in ascending order based on the points' x -coordinate. Then, it chooses a vertical line, ℓ , that divides the set P into two halves, P_1 contains $\lfloor n/2 \rfloor$ points and P_2 contains $\lceil n/2 \rceil$ points. The algorithm now recursively calls itself on P_1 and P_2 , so the minimum of the two outputs would be the temporary closest-pair distance, $\delta_0 = \min(\delta_1, \delta_2)$. It is obvious that there could be a pair of points (one from P_1 and the other from P_2) which has a closer distance than the temporary δ_0 .

To combine the two set P_1 and P_2 , we create a strip centered at ℓ with its left end and right end are the lines ℓ_1 and ℓ_2 so that both lines are δ_0 away from ℓ . All the points in the strip are now sorted based on their y -coordinate. The algorithm then can loop through the strip and check if there is any smaller distance than δ_0 . Fortunately, for each point in the strip, the algorithm just needs to check the distances between the current point with its 7 successive points, that have y -coordinates \geq the current point's y -coordinate. If there is any closer distance, it is assigned to δ_0 . Finally, δ_0 is returned.

To keep the report on point and less complicated, the code of this 2D algorithm will not be listed. See the implementation on my *GitHub repository*³.

5.2 The closest-pair doubling algorithm

The implementation of the closest-pair algorithm still consists of three main functions, `sep_ann`, `sparse_sep_ann` and `closest_pair`. Moreover, the `brute_force` function is added to be used in the base case. The following is the algorithm's class.

```

1 #include "PointList.h"
2
3 typedef std::vector<double> DVect;
4
5 class ClosestPairDoubling {

```

³The implementation of the 2D algorithm with merge sort algorithm. <https://github.com/ThangMinh-Cao/closestpairdoubling/tree/master/utlis>

```

6 private:
7     static std::tuple<Point, double, DVect>
8         sep_ann(PointList &S, int n, double mu, double c);
9     static std::pair<Point, double>
10        sparse_sep_ann(PointList &S, int n, double d, int t);
11 public:
12     ClosestPairDoubling() = default;
13     static double brute_force(PointList &S);
14     static double closest_pair(PointList &S, double d, int recursion=0);
15 };

```

Source: master branch → `closestpairdoubling/include/closest_pair_doubling.h`

5.2.1 The $\text{SEPANN}(S, n, d, \mu, c)$ algorithm

The `sep_ann` function starts by defining some important variables to be used in its loop. Let `p` be the chosen random point in S , `Rp` be the radius of the smallest ball containing at least n/c points, `distances_from_p` be the vector stores the distances between p and all points in S and `outer_ball_count` be the number of points in the outer ball, $ball_S(p, \mu R_p)$. To select a random point p , I initialize an integer random generator `int_gen` with the range from 0 to $n - 1$, possible indices of points in S .

Since c is calculated based on d which could be done outside the function, I removed the input d of the `sep_ann` function.

```

1 std::tuple<Point, double, DVect>
2     ClosestPairDoubling::sep_ann(PointList &S, int n, double mu, double c
3     )
4 {
5     Point p;
6     double Rp = -1.0;
7     DVect distances_from_p;
8     int outer_ball_count = (n / 2) + 1;
9     RandomInt int_gen(0, n - 1);
10
11     while (Rp == -1.0 or outer_ball_count > std::floor(n / 2)) {
12         distances_from_p.clear();
13
14         // selecting the random point p and calculate all the distances
15         int random_index = int_gen.next();
16         p = S.points[random_index];
17         for (const Point& point: S.points) {
18             distances_from_p.push_back(p.distance_to(point));
19         }
20         Rp = KthSmallest::get(distances_from_p, 0,
                               (int)distances_from_p.size(), ceil(n / c));

```

```

21
22 // counting the number of point in the outer ball
23 outer_ball_count = 0;
24 for (double dist: distances_from_p) {
25     if (dist <= mu * Rp) {
26         outer_ball_count++;
27     }
28 }
29 }
30 return std::make_tuple(p, Rp, distances_from_p);
31 }

```

Source: master branch \rightarrow closestpairstdoubling/src/closest_pair_doubling.cpp

Each time the *while* loop is repeated, `distances_from_p` is cleared to get rid of the previous loop's result. The function uses `int_gen` to generate a random point in S , then it stores distances between p and every point in S including itself into `distances_from_p`. The QuickSelect function from `KthSmallest` class is now used to get the $\text{ceil}(n/c)$ -smallest element in `distances_from_p`, then it is assigned to `Rp`. The function loops through the distance vector to count the number of points inside the outer ball and assigns it to `outer_ball_count`. If `outer_ball_count` $< \lfloor n/2 \rfloor$ then we are done, the function returns a tuple containing `p`, `Rp` and `distances_from_p`. If this is not the case, the procedure is repeated until the condition is satisfied. Note that, I return `distances_from_p` here to use it in SPARSESEPANN without spending time to loop through the vector again.

5.2.2 The refined algorithm SPARSESEPANN(S, n, d, t)

Let `sep_ann_res` be the output of the function `sep_ann(S, n, e, c)`, `Ai_size` be the number of points $\text{annulus}_S(p, R_{i-1}, R_i)$ contains. Let `p`, `R_prime` and `distance_from_p` be the variables store the values which are derived from `sep_ann_res`. The implementation code of SPARSESEPANN is given below.

```

1 std::pair<Point, double>
2 ClosestPairDoubling::sparse_sep_ann(PointList &S, int n, double d, int t)
3 {
4     const double e = std::exp(1.0); // the Euler constant
5     double c = 2 * pow(4 * e, d);
6     std::tuple<Point, double, DVect> sep_ann_res = sep_ann(S, n, e, c);
7
8     // followings are results derived from sep_ann_res
9     Point p = std::get<0>(sep_ann_res);
10    double R_prime = std::get<1>(sep_ann_res);
11    DVect distances_from_p = std::get<2>(sep_ann_res);
12

```

```

13  int Ai_size = n / t + 1;
14  RandomInt range_t_random(1, t);
15
16  double R;
17  while (Ai_size > n / t) {
18      int random_i = range_t_random.next();
19      double Ri = pow(1 + 1.0 / t, random_i) * R_prime;
20      R = pow(1 + 1.0 / t, random_i - 1) * R_prime;
21      Ai_size = 0;
22      for (double dist: distances_from_p) {
23          if (R <= dist and dist <= Ri) {
24              Ai_size++;
25          }
26      }
27  }
28  return std::pair<Point, double> {p, R};
29 }

```

Source: master branch → `closestpairdoubling/src/closest_pair_doubling.cpp`

The algorithm first selects a random value i in the range from 1 to t using the random generator `range_t_random`, then it calculates the inner and outer radius of $annulus_S(p, R_{i-1}, R_i)$ in which $R_i = (1 + 1/t)^i$. An inner `for` loop goes through `distance_from_p` to count the number of points in the annulus and then stores the number to `Ai_size`. If `Ai_size` $> n/t$, the right annulus is found, then it returns `p` and `R`. Otherwise, the `while` loop repeats the procedure to find the annulus that satisfies `Ai_size` $\leq n/t$.

5.2.3 The main closest-pair algorithm $CLOSESTPAIR(S, n, d)$

Let `n` be the number of points contained in S , `e` be the Euler constant, `delta0` be the closest-pair distance in S . Furthermore, I combine the three subsets S_1 , S_2 and S_3 (see Section 4) to only two that are `S1orS2` and `S2orS3` that represent the subset $S_1 \cup S_2$ and $S_2 \cup S_3$.

```

1  double
2      ClosestPairDoubling::closest_pair(PointList &S, double d, int recursion
3      )
4  {
5      int n = (int)S.points.size();
6      const double e = std::exp(1.0);
7      double delta0; // the closest-pair distance in S
8
9      if (n < 2 * pow(16 * e, d)) {
10         return brute_force(S);
11     } else {

```

```

11  int t = floor((1 / (16 * std::exp(1.0))) * pow((double)n / 2, 1 / d));
12  std::pair<Point, double> ssann_result = sparse_sep_ann(S, n, d, t);
13  Point p = ssann_result.first;
14  double R = ssann_result.second;
15  PointList S1orS2 = PointList();
16  PointList S2orS3 = PointList();
17  for (const Point& point: S.points) {
18      double d_to_p = p.distance_to(point);
19      if (d_to_p <= R) {
20          S1orS2.points.push_back(point);
21      } else if (d_to_p > R and d_to_p <= (1 + 1.0/t) * R) {
22          S1orS2.points.push_back(point);
23          S2orS3.points.push_back(point);
24      } else {
25          S2orS3.points.push_back(point);
26      }
27  }
28  std::vector<Point>().swap(S.points); // free the memory that stores S
29  double delta1 = closest_pair(S1orS2, d, ++recursion);
30  double delta2 = closest_pair(S2orS3, d, ++recursion);
31  delta0 = std::min(delta1, delta2);
32  }
33  return delta0;
34  }

```

Source: master branch → `closestpairedoubling/src/closest_pair_doubling.cpp`

With $t = \lfloor \frac{1}{16e}(n/2)^{1/d} \rfloor$, `sparse_sep_ann(S,b,d,t)` is called, and the outputs of it are assigned into `p` and `R`. The `for` loop goes through S to build the two sets by adding corresponding points to them using the distance from p to each point. Let `d_to_p` be the distance from p to the current point in the `for` loop. If $d_{to_p} \leq R$, the current point is in the $ball_S(p, R)$, so it is added to `S1orS2`. If $R < d_{to_p} \leq (1 + 1/t)R$, the point is in the $annulus_S(p, R, (1 + 1/t)R)$, then it is added to both `S1orS2` and `S2orS3`. Otherwise, the point is appended into only `S2orS3` since it is outside the outer radius.

Because every point in S is now in `S1orS2` and `S2orS3`, it is not necessary to keep the vector `S` as well as its memory usage, which could be stacked up after many recursive calls and cause memory overload. Thus, after storing all the points into `S1orS2` and `S2orS3`, I clear the vector `S` and its data in the memory using the function `vector::swap()` to swap `S` with an empty vector. Note that, I will make sure to pass the full set of points **by value** or pass a copy of it to the first call of the algorithm, so the original one will not be changed. Therefore, it can not affect the execution of `brute_force` for testing later.

After that, the outputs of two recursive calls, one on `S1orS2` and one on `S2orS3`, will be stored in `delta1` and `delta2` variables. Finally, the smaller one of them will be assigned into `delta0` and returned.

5.3 The closest-pair distance of non-Euclidean spaces

The algorithm runs correctly in a special case in which the given metric space is non-Euclidean of doubling dimension 1, see [3, Section 2.1] for more details. Given a metric space $S = \{p_1, p_2, \dots, p_n\}$ that contains n “points”, denoted p_m with $1 \leq m \leq n$. Specifically, the value of p_m represents an integer between a given range. Let i and j be any number that satisfies $1 \leq i \leq j \leq n$, the distance between p_i and p_j is defined:

$$\text{dist}(p_i, p_j) = \begin{cases} 0 & \text{if } i = j \\ 4^{\max(i,j)} & \text{if } i \neq j \end{cases}$$

Using the definition, the closest-pair algorithm finds the closest distance in this metric space normally without changing any core detail.

Although the points are just integers, I still treat them as `Point` objects with 1D coordinates. Therefore, I do not need to make many small changes which may lead to programming errors that can take a lot of time to fix, and I still keep the main idea of the algorithm that works with the `Point` class.

The only problem is that the distance between any two points could be extremely large. For instance, let the upper bound of the points’ value be a big number, say 100, so $p_i \leq p_j \leq 100$. Let $p_i = 100$, then $\text{dist}(p_i, p_j) = 4^{100} > 1.6 \times 10^{60}$ which is much larger than the maximum 32-bit integer that C++ can handle. Therefore, I decided to use a library named *Boost*⁴ which can handle numbers with precisions up to 1024 bits. When changing the current algorithm to use *Boost*, I only need to replace the `double` type of normal distances by `cpp_int` type which can automatically choose the corresponding precision for each distance. Also, I needed to make some minor changes, so the algorithm can be able to run correctly. Since there are not many changes and they do not affect the main theory of the algorithm, the implementation for this non-Euclidean will not be listed in this report. See the `strange_metric` branch⁵ of my GitHub repository for the implementation.

6 The practical running time and behavior

After doing many tests on normal Euclidean spaces, I found out that the algorithm’s behavior is different based on the way the points are arranged on the metric space. There are two approaches to generate the points that give me the most obvious differences.

1. **Generating uniformly random** points spread throughout the space. To do this, we can just use any of the random number generators to get the points’ random coordinates in a given range, see Section 5.1.1 and Section 5.1.2 for more details.
2. **Generating points evenly in a grid.** Let n be the number of points, a and b be any two factors of n that satisfies $a \times b = n$, d be the distances between any two points which are close together. Start at any position, the points are set into the $a \times b$ grid which means each point is on a corner of a d -sided square, see Figure 2.

⁴The *Boost* library’s website. <https://www.boost.org/>

⁵The implementation of the algorithm on non-Euclidean metrics. https://github.com/ThangMinhCa/o/closestpairstree/tree/strange_metric

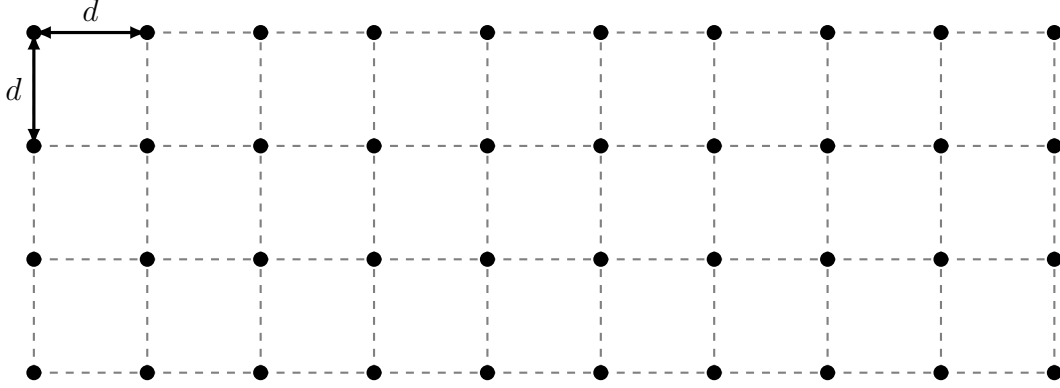


Figure 2: An example of points in a grid on a Euclidean space. A set of 40 points that gives us a 4×10 grid. Points are placed at corners of the squares whose sides are equal to d that could be any number.

Since the points in non-Euclidean spaces are simply integers, they do not have properties as many as the ones on normal Euclidean ones. Therefore, different ways to generate points do not change its behavior which is already as expected. I will only show its behavior with random points, see Section 6.2.

To generate random points in a non-Euclidean space, I use a `RandomInt` object. With a given range and number of points, I can easily call `next()` to get a value to initialize a `Point` and store it into a `PointList` object.

6.1 The practical running time

To correctly compare the algorithm's running time, we can use brute-force which takes $O(n)$ time or the 2D algorithm for only 2D points. However, to have the running time that is close to the theory, the number of points in P must be extremely large since the base case of the algorithm is when $n < 2(16e)^d$, which is already a quite large number (approximately 79550 points in 2D and more than 3000000 points in 3D, for example). The algorithm could take days to finish calculating the whole metric space, and in this case, brute-force would take a lot more than that amount of time, which is not very reasonable for me to keep the program running until it finish. Fortunately, since the running time of the algorithm is $O(n \log n)$ which is in theory and also expected in practice, the ratio of the actual running time (in seconds) to the theoretical should be approximately the same for different large enough numbers of points. Thus, we can use this information to prove that the running time in practice is actually $O(n \log n)$.

Particularly, let n be the number of points in a metric space, denoted $|P|$, m be a number which is much larger than the base case of the algorithm ($2(16e)^{\log_2 7} \approx 79550$ points in 2D metric spaces), r_n be the ratio of the actual running time to the theoretical one that is calculated using n with the log of base 2. I would initialize and run the algorithm on different metric spaces P with $n \in \text{sizes} = \{m, 2m, 4m, 8m, 16m\}$. For each size, I would run the algorithm several times to find the average running time, denoted $T(n)$.

Having n and $T(n)$, the ratio is calculated as below:

$$r_n = \frac{T(n)}{n \log n}$$

As expected, the ratios should be approximately the same. Denoted:

$$r_m \approx r_{2m} \approx r_{4m} \approx r_{8m} \approx r_{16m}$$

If this holds, the practical running time actually represents $O(n \log n)$ time. This is obvious because they have the same time complexity, so when the actual amount of time changes by an amount, the theoretical one should change by the same amount that is also based on the number of points.

After running the algorithm with different kinds of inputs, the tables and graphs below are the results that I come up with.

1. **Random generated Euclidean metric spaces:** I choose $m = 500\,000$ points since it is much bigger than the base case, and as I estimated, the algorithm would take a reasonable time to finish all cases.

n	The running time (in seconds)				$r_n = \frac{T(n)}{n \log n}$
	T_1	T_2	T_3	$T(n)$	
500 000	2176.5	2237.33	2211.71	2208.51	2.333×10^{-4}
1 000 000	3047.94	3063	3091.8	3067.58	1.539×10^{-4}
2 000 000	6280.73	6040.06	5844.34	6055.04	1.446×10^{-4}
4 000 000	12888.7	12761.5	12632.7	12760.97	1.454×10^{-4}
8 000 000	33400.4	32850.6	31038.6	32429.8	1.767×10^{-4}

Figure 3: The table of random points metric spaces' data of running time and the ratio r_n .

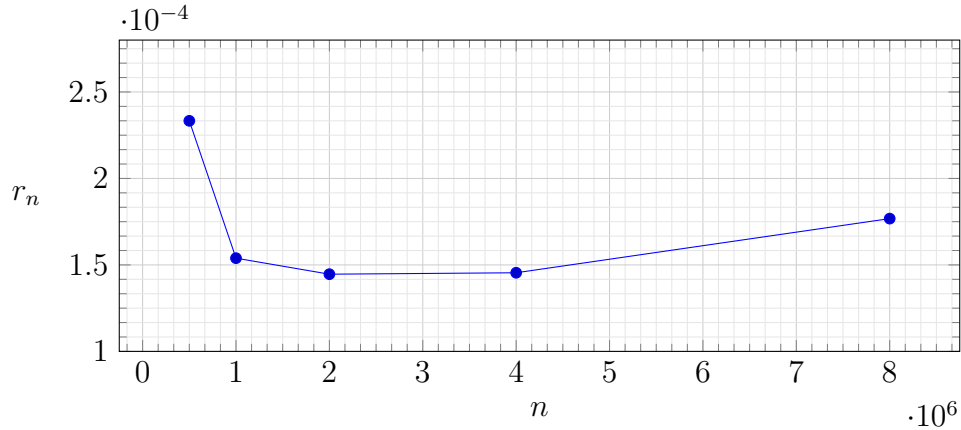


Figure 4: The graph of ratios r_n versus different values of n of random points with $n \in \{500\,000, 1\,000\,000, 2\,000\,000, 4\,000\,000, 8\,000\,000\}$ (with y-scale = 0.5×10^{-4}).

As you can see in the table (see Figure 7), for random points, the ratios r_n do not have any significant differences. By assigning them the same base and exponent of their scientific notation (10^{-4}), their coefficients are mostly the same except for the 500 000 points, which is still less than doubled of the lowest ratio. This is reasonable since there are many factors that can affect the running time of the program. Also, this is the lowest value of n in the set, so its running time probably has the highest error. The graph also shows that the points are roughly on a horizontal line, so the ratios have acceptable differences, see Figure 8.

2. **Grid generated Euclidean metric spaces:** I choose $m = 200\,000$ points in this case because the algorithm SEPANN will repeat several times most of the times it is called, that leads to much longer running time. Also, that is still a good number which is about 2.5 times larger than the base case, so it probably return a good result of running time.

n	The running time (in seconds)				$r_n = \frac{T(n)}{n \log n}$
	T_1	T_2	T_3	$T(n)$	
200 000	3467.18	3191.12	3242.48	3304.22	0.938×10^{-3}
400 000	8107.99	8298.77	8787.8	8398.18	1.128×10^{-3}
800 000	19771.3	18744.3	19015.7	19177.1	1.222×10^{-3}
1 600 000	38205.9	36618.35	39131.24	37993.08	1.152×10^{-3}
3 200 000	33400.4	32850.6	31038.6	32429.8	1.767×10^{-4}

Figure 5: The table of grid points metric spaces' data of running time and the ratio r_n .

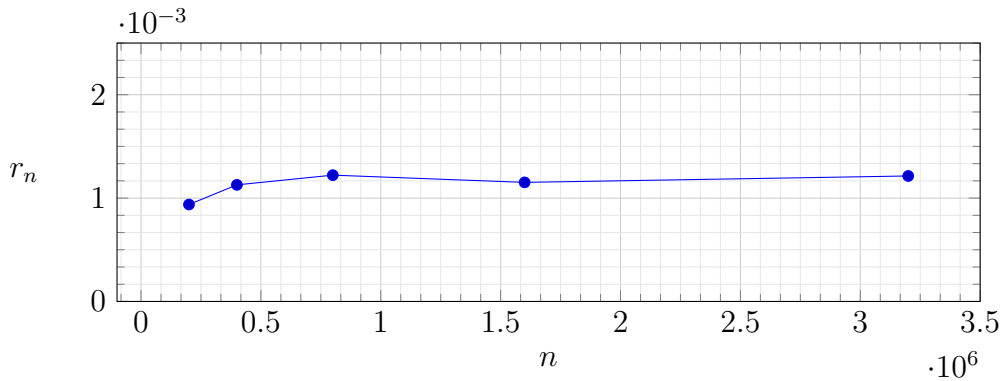


Figure 6: The graph of ratios r_n versus different values of n of grid points with $n \in \{200\,000, 400\,000, 800\,000, 1\,600\,000, 3\,200\,000\}$ (with y-scale = 10^{-3}).

The relationship is clearer in this example of grid generated points. All the ratios are very close to each other, see r_n column of Figure 5. The graph also shows that the differences between the ratios are very small since all the points are almost on a horizontal line.

3. **Random generated non-Euclidean metric spaces:** Since handling extremely large numbers makes the program significantly slower, I start with a much smaller value of m (2000 points) compared to the Euclidean spaces. However, for non-Euclidean spaces, the doubling dimension is 1, so the base case is approximately 87 points. Thus, 2000 is a pretty large number of point compared to the base case.

n	The running time (in seconds)				$r_n = \frac{T(n)}{n \log n}$
	T_1	T_2	T_3	$T(n)$	
2000	10.3557	8.83033	6.09795	8.428	3.843×10^{-4}
4000	38.6867	47.721	34.8211	40.4096	8.443×10^{-4}
8000	209.766	257.704	217.096	228.1886	2.2×10^{-3}
16000	1439.05	1202.57	1173.77	1271.4633	5.69×10^{-3}
32000	7692.18	7825.26	7718.64	7745.36	1.617×10^{-2}

Figure 7: The table of random points non-Euclidean metric spaces' data of running time and the ratio r_n .

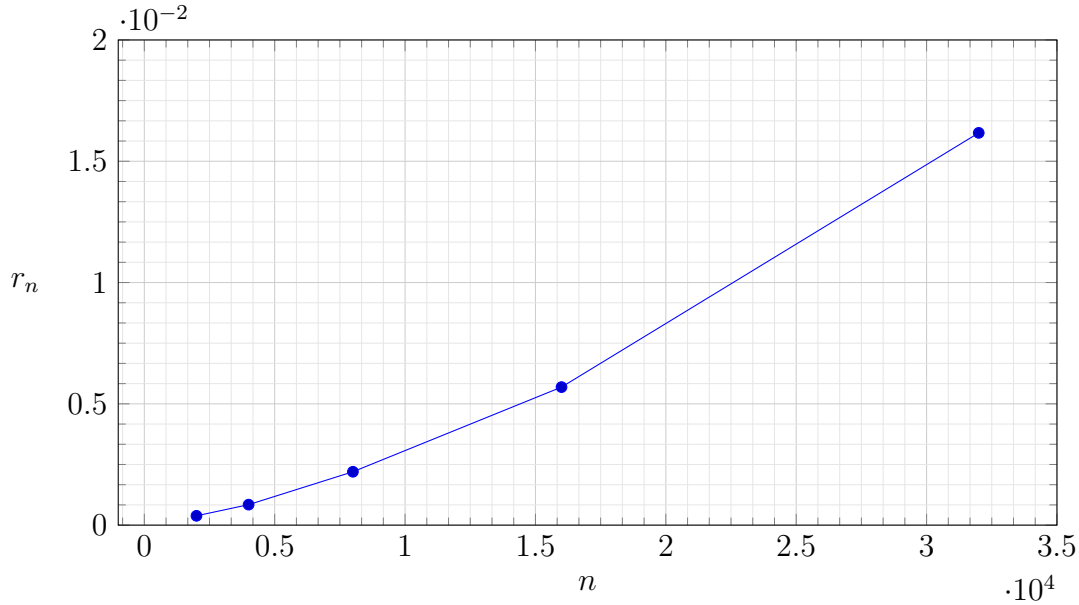


Figure 8: The graph of ratios r_n versus different values of n of random points on non-Euclidean spaces with $n \in \{2000, 4000, 8000, 16\,000, 32\,000\}$ (with y-scale = 0.5×10^{-2}).

With non-Euclidean metric spaces, the algorithm now has a different behavior with the running time in seconds. The more points the space has, the more the running time increase. As you can see, from 2000 points to 4000 points, the ratio r_n increases a little, but from 16 000 points to 32 000 points, there is a significant increase. From this, I can confidently conclude that the increase is from the handling of multiple-precision integers of *Boost* library.

Since the number of points increases, the maximum distance increases, so it takes more time to handle each instance.

Since the value of increase cannot be exactly identified, I decided to change from calculating running time in **seconds** to the **number of *dist()* function calls**. This works because distance between two points is the principle information that the algorithm uses, and it is also the algorithm's result. Furthermore, every part of the algorithm has at least one loop that access distance in each iteration, even the brute-force function. Importantly, the number of *dist()* function calls does not relate to the handling of large integer, only the value *dist()* returns.

Note that, the function `sparse_sep_ann()` does not call *dist()* after the `sep_ann()` call. However, I also increase the counter in the inner loop that iterates through the `distances_from_p` vector since the procedure in the function also accesses and uses the distances to calculate.

Below information is the table of running time by the number of *dist()* function calls and the graph represents it.

n	The running time (in <i>dist()</i> calls)				$r_n = \frac{T(n)}{n \log n}$
	T_1	T_2	T_3	$T(n)$	
2000	101104	118118	114577	111266.33	5.07334
4000	252964	253449	247274	251299	5.25036
8000	525386	498586	555710	526560.67	5.07644
16000	1157266	1137811	1146926	1147334.33	5.13457
32000	2778423	2445893	2615109	2613141.67	5.45649

Figure 9: The table of random points non-Euclidean metric spaces' data of running time and the ratio r_n .

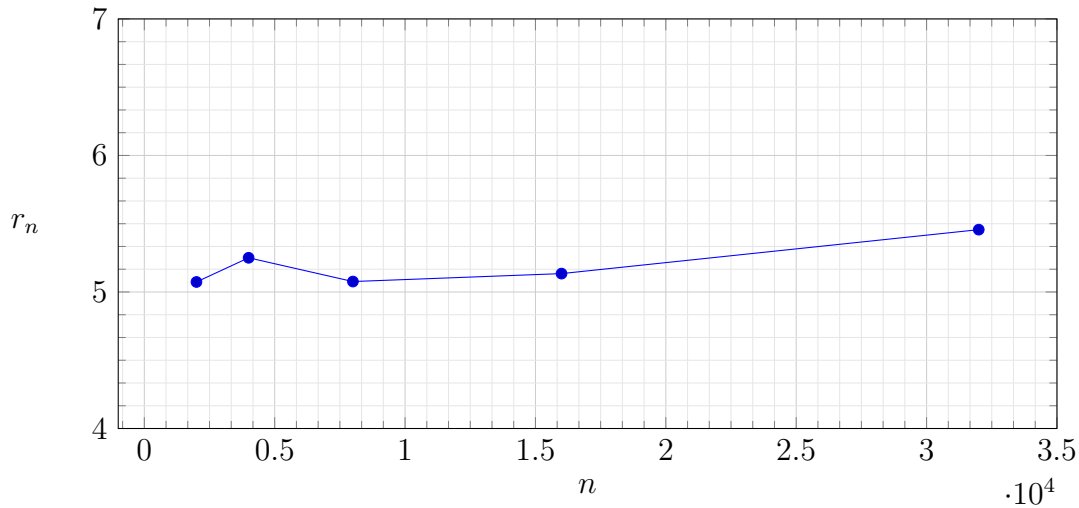


Figure 10: The graph of ratios r_n versus different values of n of random points on non-Euclidean spaces with $n \in \{2000, 4000, 8000, 16\,000, 32\,000\}$ (with y-scale = 1).

Now, it is more obvious to say that the running time of the algorithm on non-Euclidean spaces is also $O(n \log n)$. All the ratio r_n are close together and do not have the increment relationship as the outcome of run-time in seconds, see r_n column of Figure 9.

6.2 Probability to select a “good” point of $\text{SEPANN}(S, n, d, \mu, c)$

Again, the goal of algorithm $\text{SEPANN}(S, n, d, \mu, c)$ is to find a “good” point p in S . A “good” point p implies that with R_p (the radius of the smallest $\text{ball}_S(p, R_p)$ that contains at least n/c points), the $\text{ball}_S(p, \mu R_p)$ contains at most $n/2$ points. Due to A. Maheshwari, W. Mulzer and M. Smid [1, Lemma 3], the algorithm has probability at least $1/c$ (about $1/1623.4$ in 2D space) to select a good point uniformly random from S . Like the practical running time, the number of times the algorithm $\text{SEPANN}(S, n, d, \mu, c)$ repeats until it gets a good point also depends on the way how the input points are generated. This only applies the normal Euclidean metric spaces.

Generating point randomly produces a surprising behavior of this algorithm. Every time $\text{SEPANN}(S, n, d, \mu, c)$ is called, it only repeats *once* until it gets to the base case, that means the probability to get a good point is 100%, see algorithm’s `sep_ann` data files on *GitHub repository*⁶. Since the points are generated randomly, so they change every time, there are no explanations about specific properties of the points in this case. However, since 100% satisfied the probability at least $1/c$, this result is reasonable.

On the other hand, when using the grid way to generate points, the algorithm $\text{SEPANN}(S, n, d, \mu, c)$ now repeats multiple times which is its expected behavior, see an example output in Figure 11. Again, the lowest probability in most of the trials in this case is about $1/20$ which satisfies probability at least $1/c$. For the full grid input data, see the `sep_ann` data files on the *GitHub repository*⁷.

n	Repeat times	n	Repeat times
200000	6
193836	7	101943	4
179534	2	96367	2
172501	1	96154	2
158381	13	83621	6
156531	3	83288	13
...	...	82515	2

Figure 11: Given an input of 200 000 points generated in a grid. This is a portion of the data about the number of times the algorithm $\text{SEPANN}(S, n, d, \mu, c)$ repeats.

⁶Random points SepAnn data. https://github.com/ThangMinhCao/closestpairedoubling/blob/master/report/images_%26_data/closest_pair/random_generation/

⁷Grid points SepAnn data. https://github.com/ThangMinhCao/closestpairedoubling/blob/master/report/images_%26_data/closest_pair/grid_generation/

For non-Euclidean spaces, the algorithm behaves exactly as expected even with random points. Since the doubling dimension 1 which gives us the base case approximately 87, several thousands of points are large enough for an input that shows the correct running time and behavior. With different input, the algorithm produces expected answers which are calculated using brute-force but much faster, see folder `report` in my GitHub repository for more details about the outputs. With random inputs as well as any other type, the `sep_ann` function repeats several times, see Figure 12. The table below represents the number of times `sep_ann` repeats with different number of points in S .

n	Repeat times	n	Repeat times
200000	6
193836	7	101943	4
179534	2	96367	2
172501	1	96154	2
158381	13	83621	6
156531	3	83288	13
...	...	82515	2

Figure 12: Given an input of 200 000 points generated in a grid. This is a portion of the data about the number of times the algorithm $\text{SEPANN}(S, n, d, \mu, c)$ repeats.

7 Concluding remarks

I have briefly described the theory of the closest-pair doubling algorithm, and the implementation of it. Even we do not have the ability to run the algorithm with a huge number of points for more accurate running time outputs, the algorithm works exactly as we expected in practice:

1. For each example with different number of points in P , the closest-pair distance is correctly returned (the same as the brute-force's) without the use of the points' coordinates. Its running time is obviously much less than the result from brute-force.
2. The algorithm can correctly find the closest-pair distances on non-Euclidean spaces of doubling dimension 1 without changing any core details. Also, its behavior is the same as when the input is a normal metric space.
3. The practical running time of the algorithm is certainly $O(n \log n)$ which is proved by the close relationship between the practical and theoretical running time.
4. For any way that the points are arranged on the metric space, the number of times the algorithm $\text{SEPANN}(S, n, d, \mu, c)$ repeats satisfies the probability $1/c$ in theory. Therefore, in practice, this procedure runs in linear time.

References

- [1] A. Maheshwari, W. Mulzer and M. Smid. *A Simple Randomized $O(n \log n)$ -Time Closest-Pair Algorithm in Doubling Metrics*, 2020. <https://arxiv.org/abs/2004.05883>
- [2] J. L. Bentley and M. I. Shamos. Divide-and-conquer in multidimensional space. In *Proceedings of the 8th ACM Symposium on the Theory of Computing*, pages 220–230, 1976.
- [3] M. Smid. *The Weak Gap Property in Metric Spaces of Bounded Doubling Dimension*. <https://people.scs.carleton.ca/~michiel/weakgap.pdf>