

Generalized Sorting with Predictions*

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Abstract. Generalized sorting problem, also known as sorting with forbidden comparisons, was first introduced by Huang et al. [HKK11] together with a randomized algorithm which requires $\tilde{O}(n^{3/2})$ probes. We study this problem with additional predictions for all pairs of allowed comparisons as input. We propose a randomized algorithm which uses $O(n \log n + w)$ probes with high probability and a deterministic algorithm which uses $O(nw)$ probes, where w is the number of mistakes made by prediction.

1 Introduction

1.1 Generalized sorting

Sorting is arguably the most basic computational task, which is also widely used as an important component of many other algorithms. Pair-wise comparison is the core of most sorting algorithms. In the standard model, it is assumed that we can make comparison between all pairs of elements as we want and they are all of the same cost. However, this may not be the case in many applications. There might be some constraints which forbid us to compare some pairs of elements, or the costs for comparing different pairs of elements may be different. The non-uniform cost sorting model is studied in [CFG⁺02, GK01, KK03]. A special case called matching nuts and bolts problem is studied in [ABF⁺94, KMS98].

In this paper, we study the model introduced by Huang, Kannan and Khanna [HKK11], which is known as generalized sorting problem or sorting with forbidden pairs. In this model, only a subset of the comparisons are allowed, and each allowed comparison is of the same cost. We can view the elements as the vertices of a graph, each undirected edge in the graph represents a pair of elements which is allowed to be compared. A comparison of two elements a, b leads to the exposure of the direction of

edge (a, b) . It is guaranteed that the hidden directed graph is acyclic, and it contains a Hamiltonian path which represents the total order of all elements. Our goal is to adaptively probe edges in E to find out the Hamiltonian path.

For generalized sorting problem, the performance of an algorithm is measured by the number of edges it probes. In standard sorting problem where G is a complete graph, the minimum number of probes required is $\Theta(n \log n)$. For general graphs, one may need more probes. Huang et al. [HKK11] have proved an upper bound of $\tilde{O}(n^{1.5})$ on the number of probes by giving a randomized algorithm. When the graph is dense and the number of edges is as large as $\binom{n}{2} - q$, [BR16] proposes a deterministic algorithm which makes $O((n + q) \log n)$ probes together with a randomized algorithm which makes $O(n^2 / \sqrt{q + n} + n\sqrt{q})$ probes with high probability. Most part of the generalized sorting problem is still open.

1.2 Algorithms with predictions

Recently, there is an interesting line of research called algorithm design with predictions [MV20], which is motivated by the observation that by making use of predictions provided by machine learning, one may be able to design a more effective algorithm. Normally, the better the prediction, the better the performance. In this framework, we aim for algorithms which have near optimal performance when the predictions are good, and no worse than the prediction-less case when the predictions have large errors. The above two targets in algorithm design with predictions are called consistency and robustness respectively.

Take the classic binary search algorithm as an example, which can find the position of an existing element in a sorted list in $O(\log n)$ comparisons. It starts by querying the median of the list. However, if a machine learning algorithm can roughly estimate the position of the given element, it may not be always a good idea to start from the middle. Based on this idea, one designed an algorithm with query complexity of $O(\log w)$, where w is the distance between the true position of the element and the

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estimated one, which measures the accuracy of the prediction. This algorithm can be much better than $O(\log n)$ when w is much smaller than n , and no worse than the prediction-less binary search even if the estimation is terribly wrong because w is at most n .

Algorithms with predictions are studied for caching [LV18, Roh20], ski-rental, online scheduling [PSK18] and other problems. See the nice survey by Mitzenmacher and Vassilvitskii [MV20].

1.3 Our results

In this paper, we initiate the study of generalized sorting with predictions. The model is very natural, besides the undirected graph $G = (V, E)$, we are also given an orientation of G as input, which are predictions of the hidden direction of the edges. The number of mis-predicted edges is denoted by w . With the help of predictions, we hope to improve the bound of $\tilde{O}(n^{1.5})$ when w is small.

In section 3, we propose a randomized algorithm for the generalized sorting problem and prove that it probes at most $O(n \log n + w)$ edges with high probability. The description of the algorithm is simple while the analysis is quite subtle and involved.

THEOREM 1.1 (AN $O(n \log n + w)$ RANDOMIZED ALGORITHM) *There is a polynomial time randomized algorithm which solves generalized sorting problem in $O(n \log n + w)$ probes with high probability.*

In section 4, we also propose a deterministic algorithm using $O(nw)$ probes, in order to show that when w is as small as a constant, the generalized sorting with prediction problem can be solved using only linear probes.

THEOREM 1.2 (AN $O(nw)$ DETERMINISTIC ALGORITHM) *There is a polynomial time deterministic algorithm which solves generalized sorting problem in $O(nw)$ probes.*

Note that in the query complexity model, if we have two algorithms \mathcal{A} and \mathcal{B} which use $O(f(n))$ and $O(g(n))$ queries respectively, we can simply merge them into an algorithm \mathcal{C} , which simulates \mathcal{A} and \mathcal{B} , and make \mathcal{A} 's queries and \mathcal{B} 's queries alternately. Then \mathcal{C} uses only $O(\min(f(n), g(n)))$ queries. Therefore by combining our algorithms with the one in [HKK11], both consistency and robustness can be achieved.

2 Preliminaries

The input of the generalized sorting with prediction problem is an undirected graph $G = (V, E)$ together with an orientation \vec{P} of E . There is another

orientation \vec{E} of E which represents the underlying total order and is unknown to us. The problem is formally stated as follows:

DEFINITION 2.1 (GENERALIZED SORTING WITH PREDICTION) An instance of generalized sorting with prediction problem can be represented as (V, E, \vec{E}, \vec{P}) , where

- $G = (V, E)$ is an undirected graph.
- \vec{P}, \vec{E} are two orientations of E , i.e. $\forall (u, v) \in E$, exactly one of $(u, v) \in \vec{P}$ and $(v, u) \in \vec{P}$ holds, and exactly one of $(u, v) \in \vec{E}$ and $(v, u) \in \vec{E}$ holds.
- $\vec{G} = (V, \vec{E})$ is the directed graph which represents the underlying total order, it is guaranteed that \vec{G} is acyclic and there is a directed Hamiltonian path in it.
- $\vec{G}_P = (V, \vec{P})$ is the predicted directed graph, there are no more guarantees about \vec{G}_P .

An edge (u, v) whose direction is different in \vec{P} and \vec{E} is called a *mis-predicted edge*. An in-neighbor of u in \vec{G}_P which is not an in-neighbor of u in \vec{G} is called a *wrong in-neighbor*.

We use $n = |V|$ to denote the number of vertices and $w = |\vec{P} \setminus \vec{E}|$ to denote the number of mis-predicted edges. The input and the required output of the problem are stated as follows:

- Input: (V, E, \vec{P})
- Output: (v_1, v_2, \dots, v_n) s.t. $\forall 1 \leq i < n, (v_i, v_{i+1}) \in \vec{E}$, which represents the directed Hamiltonian path in \vec{G} .

Note that \vec{E} is not given as input, but is fixed at the very beginning and does not change when the algorithm is executing.

An algorithm can adaptively probe an edge and know its direction in \vec{E} . The performance of an algorithm is measured by the number of edges it probes, which may be in terms of n and w .

Recall that $\vec{G} = (V, \vec{E})$ is acyclic, so any subset $\vec{E}' \subseteq \vec{E}$ naturally defines a partial order of V . Sometimes we focus on a vertex set V' , and only consider the partial order in the induced subgraph $\vec{G}[V']$:

DEFINITION 2.2 (PARTIAL ORDER $<_{\vec{E}'}$ AND $<_{V'}$)

- $\vec{E}' \subseteq \vec{E}$ defines a partial order of V on graph (V, \vec{E}') , which is referred to as $<_{\vec{E}'}$. For $a, b \in V$, $(a <_{\vec{E}'} b)$ iff there is a directed path from a to b which only consists of edges in \vec{E}' .

- $V' \subseteq V$ defines a partial order of V' on the induced subgraph $\vec{G}[V']$, which is referred to as $<_{V'}$. For $a, b \in V'$, $(a <_{V'} b)$ iff there is a directed path from a to b which only consists of edges in \vec{E} and only passes vertices in V' .

DEFINITION 2.3 (VERTEX SETS $\mathcal{N}_{in}(G_{\vec{P}}, u), S_u, T_u$)

Denote by $\mathcal{N}_{in}(G_{\vec{P}}, u)$ the set of all in-neighbors of u in the prediction graph, i.e. $\mathcal{N}_{in}(G_{\vec{P}}, u) = \{v | (v, u) \in \vec{P}\}$.

Denote by S_u the set of real in-neighbors of u among $\mathcal{N}_{in}(G_{\vec{P}}, u)$, i.e. $S_u = \{v | (v, u) \in \vec{P} \cap \vec{E}\}$.

Denote by T_u (with respect to a specific moment) the set of in-neighbors of u , which are not known to be wrong at that moment, i.e. the corresponding edges are either correct or unprobed. $T_u = \{v | (v, u) \in \vec{P} \wedge (u, v) \notin \vec{Q}\}$ where \vec{Q} (also with respect to a specific moment) is the set of probed directed edges up to that moment.

Note by definition it always holds $S_u \subseteq T_u \subseteq \mathcal{N}_{in}(G_{\vec{P}}, u)$. S_u and $\mathcal{N}_{in}(G_{\vec{P}}, u)$ are fixed while T_u may change over time. Initially there are no probed edges, so $T_u = \mathcal{N}_{in}(G_{\vec{P}}, u)$. As the algorithm proceeds, some mis-predicted edges between $\mathcal{N}_{in}(G_{\vec{P}}, u)$ and u are found, the corresponding wrong in-neighbors no longer belong to T_u and T_u will finally shrink to S_u .

3 An algorithm using $O(n \log n + w)$ probes

3.1 Description

The algorithm maintains a set of vertices A satisfying $\forall u \in A$, direction of edges between $\mathcal{N}_{in}(\vec{G}_P, u)$ and u are all known to us (either probed or can be deduced from other probed edges). Notice that the direction of edges in the induced subgraph $G[A]$ must be all known. When $A = V$, the direction of all edges are known and we can easily find the desired Hamiltonian path.

We initialize A as \emptyset , then iteratively add ‘ideal vertices’ to A , which are defined as follows:

DEFINITION 3.1 (IDEAL VERTEX) A vertex $u \in V$ is called an *ideal vertex* if both of the following conditions are satisfied:

1. $T_u \subseteq A$.
2. The partial order $<_A$ restricted to T_u is a total order.

Before adding a vertex u to A , we need to determine the direction of edges between T_u and u (those between $\mathcal{N}_{in}(\vec{G}_P, u) \setminus T_u$ and u have been already known to be mis-predicted). For an ideal

vertex u , this can be done by using a straightforward strategy: repeatedly probe the edge (t, u) , where t is the largest vertex in T_u with respect to $<_A$. If the direction of this edge is correct, i.e. $t <_{\vec{E}} u$, we can conclude that the direction of all edges between T_u and u are correct by transitivity. We can end this phase and add u to A . Otherwise (t, u) is a mis-predicted edge, t is removed from T_u and we move on to probe the edge between the new largest vertex in T_u and u , and so on.

If there is an ideal vertex, we are in an ideal case: by probing only one edge, we either learn the direction of all edges between T_u and u and add u into A , or find a mis-predicted edge. Notice that each vertex is added to A once, and the wrong probes are charged to the w term of complexity. Therefore we can add all vertices to A in only $O(n + w)$ probes, assuming there is **always** an ideal vertex in each step.

However, the assumption does not always hold due to the existence of mis-predicted edges, i.e. there may be a time when there is no ideal vertex. We have to relax the conditions to define a new type of vertex to help, which always exists:

DEFINITION 3.2 (ACTIVE VERTEX) A vertex $u \in V$ is called an *active vertex* if both of the following conditions are satisfied:

1. $S_u \subseteq A$.
2. The partial order $<_A$ restricted to S_u is a total order.

LEMMA 3.1. *There is at least one active vertex in $V \setminus A$ if $A \neq V$.*

Proof. Suppose the Hamiltonian path in \vec{G} is (v_1, \dots, v_n) . Let k be the smallest index s.t. $v_k \notin A$, then v_k satisfies

1. $S_{v_k} \subseteq \{v_1, \dots, v_{k-1}\} \subseteq A$.
2. $<_A$ restricted to S_{v_k} is a total order.

Therefore v_k is active at the moment. \square

An ideal vertex is always active since $S_u \subseteq T_u$ holds, but there may be some wrong in-neighbors not identified yet (the vertices in $T_u \setminus S_u$) to prevent an active vertex from being ideal. By cleverly identify the wrong in-neighbors and remove them from T_u , an active vertex u would become an ideal one and we can use the above strategy again.

As S_u is invisible to us, we know neither which vertices are active, nor which in-neighbors of a vertex are wrong, so we turn to focus on the in-neighbors of u which prevent it from being ideal:

1. $v \in T_u$ s.t. $v \notin A$.
2. $v_1, v_2 \in T_u$ s.t. $(v_1, v_2 \in A) \wedge (v_1 \not\prec_A v_2) \wedge (v_2 \not\prec_A v_1)$.

Now consider an active vertex u . If such v in case 1 exists, then (v, u) must be a mis-predicted edge. If such v_1, v_2 in case 2 exists, there is at least one mis-predicted edge in $(v_1, u), (v_2, u)$. By probing (v, u) or both $(v_1, u), (v_2, u)$ repeatedly, we can keep removing its wrong in-neighbors from T_u and finally make u ideal.

For an inactive vertex u , the direction of (v, u) or both of $(v_1, u), (v_2, u)$ may be correct, but that tells us u is not active hence is not the vertex we are looking for. In this case the vertex v or the pair of vertices (v_1, v_2) is called a *certificate* for u , which proves that u is currently not active.

DEFINITION 3.3 (CERTIFICATE) For a vertex $u \in V$,

1. A type-1 certificate is a vertex $v \in S_u$ s.t. $v \notin A$.
2. A type-2 certificate is a pair of different vertices $v_1, v_2 \in S_u$ s.t. $(v_1, v_2 \in A) \wedge (v_1 \not\prec_A v_2) \wedge (v_2 \not\prec_A v_1)$.

Once a certificate for u is found, we turn to check the activeness of other vertices and do not need to probe any other incoming edges for u until the next vertex is added to A . For a fixed set A , both activeness of vertices and validity of certificates are determined and do not change when new probes are made. Only when A is extended and the current certificate of u is no longer valid do we need to look for a new certificate for u . A type-1 certificate v becomes invalid when v is added into A , while a type-2 certificate (v_1, v_2) becomes invalid when $(v_1 \prec_A v_2) \vee (v_2 \prec_A v_1)$ happens as A expands.

In the worst case, one may need to update the certificates again and again and thus probe too many edges. By checking the validity of certificates in a random order, the worst case is avoided with high probability. We prove that our algorithm uses only $O(n \log n + w)$ probes with high probability in the next subsection, where the term $n \log n$ comes from the probes used in re-searching for valid certificates.

Our algorithm works by repeatedly choose a vertex u which does not have a valid certificate.

1. If it is an ideal vertex, we use the strategy mentioned above to determine the direction of edges between T_u and u , then add u to A .
2. Otherwise there must be a vertex $v \in T_u$ s.t. $v \notin A$, or a pair of vertices $v_1, v_2 \in T_u$ s.t. $(v_1, v_2 \in A) \wedge (v_1 \not\prec_A v_2) \wedge (v_2 \not\prec_A v_1)$. We

randomly choose such a vertex v or such a pair of vertices (v_1, v_2) and probe the edge(s) between u and them. Then either at least one mis-predicted edge is found, or a valid certificate for u is found.

Since there is always an active vertex u , after finding some mis-predicted edges and removing the corresponding wrong in-neighbors from T_u , u must become an ideal vertex, that is how the algorithm makes progress.

Here is the pseudo code of the algorithm:

Algorithm 1 A randomized algorithm using $O(n \log n + w)$ probes

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1: set  $A := \emptyset$ 
2: while  $A \neq V$  do
3:   pick  $u \in V \setminus A$  s.t.  $u$  does not have a valid
   certificate (if there are multiple ones, pick one
   with the smallest index)
4:   if  $T_u \not\subseteq A$  then
5:     randomly pick  $v \in T_u \setminus A$ 
6:     probe  $(v, u)$ 
7:   else if  $\exists$  different  $v_1, v_2 \in T_u$  s.t.  $(v_1 \not\prec_A$ 
    $v_2) \wedge (v_2 \not\prec_A v_1)$  then
8:     randomly select such a pair  $v_1, v_2$ 
9:     probe  $(v_1, u), (v_2, u)$ 
10:  else
11:    let  $t$  be the largest vertex in  $T_u$  w.r.t.  $<_A$ 
12:    probe  $(t, u)$ 
13:    if the direction of  $(t, u)$  is correct, i.e.
    $t <_{\vec{E}} u$  then
14:      add  $u$  to  $A$ 

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3.2 Analysis

We first prove that the algorithm can always proceed, and always terminates.

LEMMA 3.2. *The algorithm can always proceed, and will terminate in finitely many steps.*

Proof. We know from Lemma 3.1 that there is always at least one active vertex outside A if $A \neq V$. Since active vertices have no valid certificates, in each execution of line 3, there is always at least one vertex which meets our requirements.

In each execution of the ‘while’ loop, exactly one of the following happens:

1. We find a certificate for u which doesn’t have a valid one previously.
2. We find a mis-predicted edge.
3. We add a vertex u into A .

When case 1 happens, we find a new certificate for u . Only when this certificate becomes invalid do we need to look for a new one for u . An invalid certificate will never become valid again since A is always enlarging. Therefore this case happens for finitely many times.

Case 2 happens for finitely many times because once we find a mis-predicted edge, we remove a vertex from T_u , the size of which is initially finite and non-negative all the time.

Case 3 also happens for finitely many times because each vertex is added into A only once. \square

Now we proceed to analyze the total number of probes made by the algorithm. First we introduce some important lemmas:

LEMMA 3.3. *All vertices are added into A in a fixed order regardless of the randomness.*

Proof. Recall that an ideal vertex is always active, so we only add active vertices to A . Whether a vertex u is active or not only depends on the current A . Therefore $\forall 1 \leq i \leq n$, when $|A| = i - 1$, the i -th vertex added into A is always the one with the smallest index among all active vertices in $V \setminus A$ regardless of randomness. \square

COROLLARY 3.4. *Let $C_{(a,b)}(a, b \in V)$ denotes the event that the vertex pair (a, b) becomes comparable in $<_A$, i.e. $(a, b \in A) \wedge ((a <_A b) \vee (b <_A a))$. As A expands, all $\binom{n}{2}$ such events happen in a fixed order regardless of the randomness (breaking ties in an arbitrarily fixed manner).*

Remark. Lemma 3.3 and Corollary 3.4 use the fact that we determine the order among vertices in T_u not according to all edges probed till now, but only according to the edges in the induced subgraph $G[A]$. It seems more efficient if we use all the information instead of the restricted portion, but it will create subtle correlations and we do not know how to analyze. The above fixed-order properties are crucial in the proof of our main theorem.

LEMMA 3.5. *For a random permutation $\{Y_1, \dots, Y_n\}$ of $\{1, \dots, n\}$, the number of elements Y_i s.t. $Y_i = \max_{j \leq i} Y_j$ does not exceed $6 \ln n + 6$ w.p. at least $1 - \frac{1}{2n^2}$.*

Proof. A random permutation can be built in such a way:

- $\forall 1 \leq i \leq n$, randomly and independently pick Z_i from $\{1, \dots, i\}$.

- take the unique permutation $\{Y_1, \dots, Y_n\}$ s.t. $\forall i, Y_i$ is the Z_i -th largest element in $\{Y_1, \dots, Y_i\}$.

It's easy to see a random permutation built in this way is uniformly distributed.

Let $\{X_1, \dots, X_n\}$ be a sequence of 0-1 random variables indicating whether $Z_i = i$. The number mentioned in the lemma is just $X = \sum_{i=1}^n X_i$ since $Z_i = i \Leftrightarrow Y_i = \max_{j \leq i} Y_j$. We have

$$\Pr[X_i = 1] = 1 - \Pr[X_i = 0] = \frac{1}{i}$$

Note that $\mu = \mathbb{E}[X] = H_n = \sum_{i=1}^n \frac{1}{i} \approx \ln n + \gamma$ as $n \rightarrow \infty$ where γ is the euler constant, and

$$\ln n + \gamma < H_n \leq \ln n + 1, \forall n \geq 1$$

Plugging $\epsilon = 5$ into a Chernoff bound: $\Pr[X > (1 + \epsilon)\mu] \leq \exp\left(-\frac{\mu\epsilon^2}{2+\epsilon}\right)$, we have

$$\begin{aligned} \Pr[X > 6(\ln n + 1)] &\leq \exp\left(-\frac{25}{7}(\ln n + \gamma)\right) \\ &\leq \frac{1}{2n^2} \end{aligned}$$

\square

THEOREM 1.1 (AN $O(n \log n + w)$ RANDOMIZED ALGORITHM) *There is a polynomial time randomized algorithm which solves generalized sorting problem in $O(n \log n + w)$ probes with high probability.*

Proof. The correctness of the algorithm directly follows from Lemma 3.2. We only need to bound the number of probes it uses.

Follow the proof of Lemma 3.2 we know in each execution of the 'while' loop, exactly one of the three cases happens:

- We find a certificate for u which doesn't have a valid one previously.
- We find a mis-predicted edge.
- We add a vertex u into A .

The algorithm makes no more than two probes in each loop, so it's sufficient to bound the number of occurrences of each cases.

Case 2 and case 3 happen for at most $n + w$ times in all, because the number of mis-predicted edges is at most w and each vertex is added into A exactly once.

We now focus on case 1. Once a valid certificate for u is found, we won't make any further probes for u until its current certificate becomes invalid.

According to Lemma 3.3, all vertices are added into A in a fixed order, which means all possible type-1 certificates for u (the set of which is S_u initially) become invalid in a fixed order.

Each time we find a uniformly random type-1 certificate for u among its currently valid ones. In the analysis it can be equivalently viewed as that for each u , a random permutation $\{P_1^{(u)}, \dots, P_{|S_u|}^{(u)}\}$ of S_u is chosen and fixed at first, and all type-1 certificates used in the process are identified according to this permutation: when a new valid type-1 certificate is found, let it be $P_i^{(u)}$, where i is the smallest index s.t. $P_i^{(u)}$ is currently valid as a type-1 certificate for u .

Let $\{Y_1^{(u)}, \dots, Y_{|S_u|}^{(u)}\}$ represents the fixed order of $\{P_1^{(u)}, \dots, P_{|S_u|}^{(u)}\}$ to become invalid, i.e. $P_i^{(u)}$ is the $Y_i^{(u)}$ -th earliest to become invalid. $\{Y_1^{(u)}, \dots, Y_{|S_u|}^{(u)}\}$ is a uniformly random permutation as well as $\{P_1^{(u)}, \dots, P_{|S_u|}^{(u)}\}$, and the total number of valid type-1 certificates found for u equals to the number of $Y_i^{(u)}$ s.t. $Y_i^{(u)} = \max_{j \leq i} Y_j^{(u)}$.

From Lemma 3.5 we know w.p. at least $1 - \frac{1}{2n^2}$, this number does not exceed $6 \ln n + 6$. Take union bound over all u , w.p. at least $1 - \frac{1}{2n}$, no vertex uses more than $(6 \ln n + 6)$ type-1 certificates, hence total number of valid type-1 certificates the algorithm finds does not exceed $6n \ln n + 6n$.

The above analysis is exactly the same for type-2 certificates, since according to Corollary 3.4, the possible type-2 certificates for u also become invalid in a fixed order. The only difference is that the number of valid type-2 certificates for each u may be up to n^2 , while it is at most n for type-1 certificates. Again use Lemma 3.5 and take union bound, w.p. at least $1 - \frac{1}{2n}$, the total number of valid type-2 certificates the algorithm finds does not exceed $12n \ln n + 6n$.

Combining all cases we can conclude that w.p. at least $1 - \frac{1}{n}$, the algorithm uses no more than $O(n \log n + w)$ probes in total. \square

4 An algorithm using $O(nw)$ probes

Here we briefly introduce a deterministic algorithm using $O(nw)$ probes, to show that the generalized sorting with prediction problem can be solved in only linear probes when w is as small as a constant.

The basic idea is to find a mis-predicted edge in $O(n)$ probes and correct it in the predicted graph. We use $\vec{G}_C = (V, \vec{P}_C)$ to denote the predicted graph after correction: $\vec{P}_C = \{(v, u) \in \vec{P} \mid (u, v) \notin \vec{Q}\} \cup \vec{Q}$ where $\vec{Q} \subseteq \vec{E}$ is the set of directed edges probed till now.

If there is a directed cycle in \vec{G}_C , there must be at least one mis-predicted edge on the cycle since the actual \vec{G} is acyclic. We just probe all edges on a simple directed cycle, update \vec{G}_C and loop again.

If \vec{G}_C is acyclic, consider running topological sort on it. If the direction of all edges in G_C are correct, each time there should be exactly one vertex whose in-degree is 0. If not, i.e. there are two vertices v_1, v_2 with in-degree 0 at the same time, this can only happen when there are some mis-predicted edges either adjacent to v_1, v_2 or on the path produced by the topological sort before. We probe all such edges and loop again.

The pseudo code of the algorithm is stated as follows:

Algorithm 2 A deterministic algorithm using $O(nw)$ probes

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1: while there is no Hamiltonian path consisting of
   only probed edges do
2:   if there is a simple directed cycle in  $\vec{G}_C =$ 
      $(V, \vec{P}_C)$  then
3:     probe all the edges on the cycle
4:   else
5:     run topological sort on  $\vec{G}_C$  and stop when
      $\exists v_1, v_2$  both with in-degree 0
6:     let  $(a_1, \dots, a_k)$  be the (partial) topological
     order
7:     if  $k = n$  (i.e. no such  $v_1, v_2$  found) then
8:        $\forall 1 \leq i < k$ , probe the edge  $(a_i, a_{i+1})$ 
9:     else
10:       $\forall 1 \leq i < k$ , probe the edge  $(a_i, a_{i+1})$ 
11:      probe all edges adjacent to  $v_1, v_2$ 

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THEOREM 1.2 (AN $O(nw)$ DETERMINISTIC ALGORITHM) *There is a polynomial time deterministic algorithm which solves generalized sorting problem in $O(nw)$ probes.*

Proof. In each execution of the ‘while’ loop, exactly one of the following three cases happens, and we analyze them separately:

1. *There is a simple directed cycle in \vec{G}_C .* Since the actual \vec{G} is acyclic, at least one edge on the cycle in \vec{G}_C is mis-predicted. By probing all edges on it we will find at least one mis-predicted edge.
2. *\vec{G}_C is acyclic and $k = n$ in line 7, i.e. the topological sort terminates normally.* By probing all edges between the adjacent vertices in the topological order, we either find a mis-predicted edge, or can know that the resulting path is just the desired Hamiltonian path.

3. \vec{G}_C is acyclic and there are two different vertices v_1, v_2 with in-degree 0 during the topological sort.

In this case, a mis-predicted edge must be found in line 10 or line 11. Prove it by contradiction, assume all edges probed in line 10 and line 11 are correct, if $k > 0$, (a_k, v_1) and (a_k, v_2) must both lie in \vec{G}_C since the topological sort stops just after handling a_k , so $\forall 1 \leq i \leq k, (a_i <_{\vec{E}} v_1) \wedge (a_i <_{\vec{E}} v_2)$ holds due to transitivity. W.l.o.g assume $v_1 <_{\vec{E}} v_2$. Consider the directed path from v_1 to v_2 in the actual graph \vec{G} , let it be (b_1, \dots, b_l) where $b_1 = v_1$ and $b_l = v_2$. Note that $a_1 <_{\vec{E}} \dots <_{\vec{E}} a_k <_{\vec{E}} b_1 = v_1 <_{\vec{E}} \dots <_{\vec{E}} b_l = v_2$. Therefore the edge $(b_{l-1}, b_l) \in \vec{G}_C$ and $b_{l-1} \notin \{a_1, \dots, a_k\}$, which contradicts the fact that the in-degree of v_2 is 0 at that moment.

Therefore in each loop, we either find at least one mis-predicted edge in \vec{G}_C or find the correct Hamiltonian path. It's obvious that the number of probes we make is $O(n)$ in each loop, so the total number of probes does not exceed $O(nw)$. \square

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