Weighted Distinct Sampling: Cardinality Estimation for SPJ Queries

(Full Version)

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ABSTRACT

SPJ queries form the backbone of many SQL queries used in practice. Accurate size estimation of these queries is thus an important problem, with applications in query optimization, approximate query processing, and data analytics. However, this problem has not been rigorously addressed in the literature, despite the fact that size estimation techniques of the three relational operators, selection, projection, and join, have each been extensively studied (but not when used in combination) in the past 30+ years. The major technical difficulty is that (distinct) projection seems to be difficult to combine with the other two operators when it comes to query size estimation.

In this paper, we give the first formal study of cardinality estimation for SP queries. While it was studied in a prior work in 2001, there is no guarantee on its optimality. We define a class of algorithms, which we call *weighted distinct sampling*, for estimating SP query sizes, and show how to find a near-optimal sampling strategy that is away from the optimum only by a lower order term. We then extend it to handling SPJ queries, giving the first non-trivial solution for SPJ query size estimation. We have also performed an extensive experimental evaluation to complement our theoretical findings.

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1 INTRODUCTION

Query size estimation (*a.k.a.* cardinality estimation) is a fundamental problem in database systems that requires little justification for its importance. Size estimation techniques for the three basic relational operators: Selection (σ), Projection¹ (π), and (natural) Join (\bowtie), have each been extensively studied in the literature. However, most existing methods do not compose, which becomes a serious

 1 In this paper, the projection operator π_A uses its relational algebra semantics, *i.e.*, duplicates in A are removed after the projection. If not, this operator need not be considered as it does not affect the query size. This corresponds to SELECT DISTINCT A in SQL. Furthermore, GROUP BY A yields a result size identical to that of π_A , so our techniques also apply to queries where the projection is replaced by a group-by.

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problem to their practical adoption as these relational operators rarely appear on their own in a query. While a number of techniques can handle SJ queries [8, 25, 32, 42], how to deal with P jointly with SJ largely remains an unsolved problem. There is one early work investigating size estimation for SP queries [19], which is reviewed in more detail in Section 1.3, but no (non-trivial) techniques can handle SPJ queries.

When P is the only operator applied on a table R, the size of $\pi_A(R)$ is the number of distinct values (NDV) in attribute A. The problem of estimating the NDV using small space has been studied for more than 30 years by both the database and the algorithms community [5, 6, 11, 13–15, 30, 41]. However, P by itself does not yield many interesting queries. Instead, it is often combined with S and J, as illustrated by the following examples on the TPC-H schema:

SELECT count(distinct o_custkey) FROM orders
WHERE o_orderdate > 2020-01-01
 AND o_orderpriority = "1-URGENT"

SELECT count(distinct o_custkey) FROM orders, lineitem WHERE o_orderdate > 2020-01-01 AND l_extendedprice > 100 AND o_orderpriority = "1-URGENT"

AND o_orderkey = 1_orderkey

The first is an SP query that counts the number of distinct customers who have ever placed an urgent order this year, while the second is an SPJ query that adds one more constraint that the order must contain one item priced at least 100.

The current practice for dealing with such queries decouples P and SJ, i.e., it just returns the smaller of the NDV of projection attribute and the estimated size of SJ query. This over-simplistic approach ignores any interaction between P and SJ, which often leads to a gross overestimation. As an extreme case, imagine that there are many customers, but only one has placed urgent orders this year, and it happens that s/he placed many urgent orders.

In this paper, we first design a new technique for estimating the size of SP queries, significantly improving over the results from [19] in both theory and practice. Then we consider SPJ queries, advancing the state of the art of cardinality estimation one step closer to general SQL queries.

1.1 Problem Definition

Any SPJ query can be expressed by the following normal form [1]:

$$Q := \pi_A(\sigma_\phi(R_1 \bowtie R_2 \bowtie \cdots \bowtie R_m)). \tag{1}$$

When m = 1, this becomes a SP query. For simplicity, we only consider the case where the projection is on a single attribute A. For SP queries, this is not a restriction, since we can (conceptually) concatenate all the projection attributes into a single one. This does

not work for SPJ queries, though, and we leave SPJ queries with multiple projection attributes to future studies.

We assume that all the relations R_1,\ldots,R_m and the projection attribute A are known in advance. However, the selection condition ϕ is only given at query time and can be arbitrary. In particular, ϕ may involve attributes from different relations, so our formulation also incorporates theta-joins with arbitrary join conditions given at query time. The goal is to construct a small synopsis on R_1,\ldots,R_m , such that given any ϕ , we can estimate |Q| from the synopsis with small error.

Our definition reflects how cardinality estimation is needed in practical database systems: The join structure of the queries has only limited forms, often confined by the database schema; the projection attribute also has limited possibilities. So it is affordable to build a synopsis for each possible query in the form of (1) that the database expects to receive. On the other hand, the selection condition ϕ may use arbitrary expressions (even UDFs) over any attributes, which is not possible to be given in advance.

We introduce some more notations. For SP queries, we use R to denote the only relation in (1), whose size is N. For SPJ queries, we use $N_{\bowtie} = |R_1 \bowtie \cdots \bowtie R_m|$ to denote the join size. When the projection attribute A is clear from the context, we use D to denote the NDV of A, and use D^{ϕ} to represent the NDV of A subject to the selection condition ϕ , which is also the query size |Q| we wish to estimate. We use \hat{D}^{ϕ} to denote an estimator of D^{ϕ} . Table 1 summarizes the notations used in this paper.

	• •
Notation	Meaning
R, R_1, \ldots, R_m	Input relation(s)
A	Projection attribute
$\mathbf{dom}(A)$	Domain of attribute A
ϕ	Selection condition
N	Number of tuples in <i>R</i> (for SP queries)
N_i	Number of tuples with $A = i$ where $i \in \mathbf{dom}(A)$
N_i^ϕ	Number of tuples with $A = i$ that pass ϕ
N_{\bowtie}	Join size $ R_1 \bowtie \cdots \bowtie R_m $
D	NDV of A
D^{ϕ}	NDV of <i>A</i> with selection condition ϕ
\hat{D}^{ϕ}	Estimator for D^{ϕ}
n	Sample size
$A_{\mathcal{S}}$	Set of sampled values from dom (<i>A</i>)
p_i	The probability that value i is sampled
$ au_i$	Sample size allocated for value <i>i</i>
n_i^{ϕ}	Number of sampled tuples with $A = i$ that pass
ı	ϕ where $i \in \mathbf{dom}(A)$
$\mu_i(\phi)$	Conditional probability of sampling a passing
	tuple for i , $\Pr[n_i^{\phi} \ge 1 \mid i \in A_s]$
ϕ°	Special filter where $N_i^{\phi^{\circ}} = 1$ for all $i \in \text{dom}(A)$
μ_i	Abbreviation of $\mu_i(\phi^{\circ}) = \tau_i/N_i$
N, p, τ, μ	Vectors of above symbols

Table 1: Notation used in the paper.

1.2 Error Metric

Most algorithms for the standard NDV problem (*i.e.*, no selection condition) provide multiplicative error guarantees. However, there

is a simple argument that such a guarantee is not possible for SP queries (hence also for SPJ queries), when the selection condition is only given at query time and can be arbitrary: Consider two selection conditions ϕ_0 and ϕ_1 . ϕ_0 blocks all tuples in R, while ϕ_1 blocks all but one that is arbitrarily chosen. So $D^{\phi_0}=0$ and $D^{\phi_1}=1$. Thus, unless the synopsis has kept more than half of the tuples, it cannot distinguish the two cases with confidence more than 1/2. Note that not being able to distinguish between 1 and 0 means that the multiplicative error is unbounded.

Therefore, we aim at designing synopses with additive errors. This helps circumvent the 0 vs 1 problem above, as an additive error of Err means that the synopsis can be insensitive to any query size smaller than Err. In fact, additive errors are more appropriate in many applications of query size estimation, in particular query optimization. In query optimization, the goal is actually not to find the optimal query plan, but to avoid bad plans [18]. A multiplicative error would require us to estimate small query sizes accurately, which is unnecessary, while incurring larger errors for large query sizes, which may correspond to plans we wish to avoid.

Specifically, we measure the accuracy of an estimator \hat{D}^{ϕ} for a particular selection condition ϕ by the standard Mean Squared Error (MSE), and:

$$\begin{aligned} \mathbf{MSE}[\hat{D}^{\phi}] &= \mathbf{E}[(\hat{D}^{\phi} - D^{\phi})^{2}] = \mathbf{E}^{2}[\hat{D}^{\phi} - D^{\phi}] + \mathbf{Var}[\hat{D}^{\phi} - D^{\phi}] \\ &= \mathbf{Bias}^{2}[\hat{D}^{\phi}] + \mathbf{Var}[\hat{D}^{\phi}]. \end{aligned}$$

Note that by Chebyshev inequality, an MSE can be translated into an additive error of $O\left(\sqrt{\text{MSE}[\hat{D}^{\phi}]}\right)$ with arbitrary constant probability.

We would like the synopsis to handle arbitrary selection conditions given at query time with a guaranteed MSE. Thus, we measure the quality of a synopsis on the worst possible ϕ , *i.e.*, $\max_{\phi} \text{MSE}[\hat{D}^{\phi}]$.

1.3 Distinct Sampling

Let $\mathbf{dom}(A)$ be the domain of the projection attribute A. All algorithms for the standard NDV problem are based on the idea of distinct sampling: For a parameter $0 , we take each <math>i \in \mathbf{dom}(A)$ into the sample with probability p, which is often implemented by checking if $h(i) \leq p$ for a random hash function $h : \mathbf{dom}(A) \to [0, 1]$. Let A_s be the set of distinct values of A sampled. Then, $|A_s|/p$ is a good estimator of D.

Now consider SP queries. As the selection condition ϕ is only given at query time and can take an arbitrary form, one natural idea is to augment A_S with additional tuples sampled from R. More precisely, for a parameter τ , we take τ tuples into the sample, randomly chosen from all tuples with A=i, for each $i\in A_S$; if there are less than τ tuples with A=i for some i, all of them are taken. The sample can be collected easily in one pass over the data by running the reservoir sampling algorithm for each i with $h(i) \leq p$.

The estimator for SP queries is then changed to

$$\hat{D}^{\phi} = \frac{1}{p} \left| \{ i \in A_s \mid n_i^{\phi} \ge 1 \} \right| = \frac{1}{p} \sum_{i \in \text{dom}(A)} I[n_i^{\phi} \ge 1], \quad (2)$$

where $I[\cdot]$ is the indicator function and n_i^{ϕ} is the number of tuples with A = i in the sample that pass the condition ϕ . Indeed, this was

the idea of the first and only work addressing size estimation for SP queries, due to Gibbons [19].

The key drawback of the simple sampling strategy above is that the same p and τ are applied for all distinct values in $\mathbf{dom}(A)$. While this makes sense for the standard NDV problem, since each distinct value in A contributes only once to the count no matter how many times it appears in R, it may lead to a sub-optimal solution for SP queries. The following example shows a scenario where it can be quadratically worse than the optimal sampling strategy in terms of MSE.

Example 1. Consider a table R with D distinct values in A. \sqrt{D} of these distinct values each have $3\sqrt{D}$ tuples, which we call *heavy* values, and the rest of the distinct values have 1 tuple each, which are called *light*. Note that the entire table R has $\sqrt{D} \cdot 3\sqrt{D} + (D - \sqrt{D}) \approx 4D$ tuples in total. Suppose the sample budget is 2D.

If $\tau > 2\sqrt{D}$, p cannot be larger than 2/3, otherwise the sample budget would be exceeded. Consider a selection condition ϕ that passes all the tuples of the light values and blocks all the rest (i.e., blocks all heavy values). We have $\operatorname{Var}[\hat{D}^{\phi}] = \frac{1}{p^2} \cdot p(1-p) \cdot (D-\sqrt{D}) = \Omega(D)$ (as $p \leq 2/3$ in this construction, $\frac{1}{p}-1$ is larger than 1/2).

If $\tau \leq 2\sqrt{D}$, then consider a series of selection conditions $\phi(x)$ that passes x tuples for each of the heavy values, for some $1 \leq x \leq 3\sqrt{D}$, and blocks the rest. We have $D^{\phi(x)} = \sqrt{D}$, and $\mathbf{E}[\hat{D}^{\phi(x)}] = \mu(x)\sqrt{D}$, where $\mu(x) = 1 - {3\sqrt{D} - x \choose \tau} / {3\sqrt{D} \choose \tau}$ is the probability that at least one of the x tuples is sampled. Observing that x is unknown to the estimator and $\mu(x)$ can range from $\frac{\tau}{3\sqrt{D}}$ to 1, an uncertainty gap of at least 1/3, so $\mathbf{Bias}[\hat{D}^{\phi(x)}]$ must be $\Omega(\sqrt{D})$ for some values of x, i.e., $\max_X \mathbf{MSE}[\hat{D}^{\phi(x)}] = \Omega(D)$. Note that this bias cannot be removed by scaling up or down the estimator (2).

On the other hand, a better sampling strategy is the following: We sample each light value with probability 1 and sample each heavy value with probability 1/3. If a value is sampled (heavy or light), we take all of its tuples. We correspondingly modify the estimator to

$$\hat{D}^{\phi} = \sum_{i \in \text{dom}(A)} \frac{1}{p_i} \mathbf{I}[n_i^{\phi} \ge 1], \tag{3}$$

where p_i is the probability that i is sampled, i.e., $p_i = 1$ for a light i and $p_i = 1/3$ for a heavy i. It can be verified that the (expected) sample size is 2D, and $\text{MSE}[\hat{D}^{\phi}] = \text{Bias}^2[\hat{D}^{\phi}] + \text{Var}[\hat{D}^{\phi}] = 0 + O(\sqrt{D}) = O(\sqrt{D})$.

1.4 Weighted Distinct Sampling

The example above suggests that, unlike the standard NDV problem where all distinct values are treated equally, SP queries call for a weighted sampling strategy according to the frequencies of the values, where the "weight" corresponds to the sample budget, *i.e.*, $p_i \tau_i$, allocated to each $i \in \mathbf{dom}(A)$.

More precisely, in this paper we study the *weighted distinct sam*pling problem: For each $i \in \mathbf{dom}(A)$, we sample it with probability p_i . This can be implemented by checking if $h(i) \le p_i$ for a random hash function². If it is sampled, then we take τ_i tuples into the sample, randomly chosen from all tuples with A = i. The goal is to minimize the worst-case MSE of the estimator (3). Denoting by N_i the number of tuples in R with A = i, then finding the optimal sampling strategy can be formulated as the following optimization problem (we use bold symbols to represent the vector forms of p_i , τ_i , N_i):

$$\begin{array}{ll} \underset{\boldsymbol{p},\tau}{\text{minimize}} & \underset{\boldsymbol{\phi}}{\text{max}} f_0(\boldsymbol{p},\tau,\phi) \\ \text{subject to} & 0 < \boldsymbol{p} \leq 1, \\ & 0 \leq \tau \leq N, \\ & \boldsymbol{p} \cdot \boldsymbol{\tau} \leq n. \end{array} \tag{4}$$

Here $f_0(\boldsymbol{p}, \boldsymbol{\tau}, \phi) = \text{MSE}[\hat{D}^{\phi}]$, < and \leq denote product (elementwise) order, and n is the given sample budget. So the last constraint ensures that the (expected) sample size does not exceed the budget. Note that we require $p_i > 0$ for all i, so that (3) is well defined. If a sampling strategy does not want to sample a particular i, it can set $\tau_i = 0$ instead.

Intuitively, higher sample budgets should be allocated to values i's with higher N_i 's; indeed, determining this relationship quantitatively is a key technical problem we solve in this paper. To contrast, we call the sampling strategy with the same p and τ for all i uniform distinct sampling, which simply allocates the same sample budget to all i.

1.5 Our Contributions

In addition to proposing the idea of weighted distinct sampling, we make the following technical contributions in this paper:

- We design an O(D)-time algorithm that, given the frequency vector N (in sorted order), finds a near-optimal solution to the optimization problem (4). The solution found by the algorithm is larger than the optimum by only a lower order term. Note that after we have solved (4) for p and τ , the sample can be collected in one pass over the data using a hash function h and reservoir sampling.
- As illustrated by Example 1, weighted distinct sampling works better on more skewed data. We substantiate this intuition by analyzing its MSE against the Zipfian distribution, which quantitatively shows that the MSE reduces as data skewness increases.
- We also show that our sampling strategy has a worst-case MSE of O(min{DN/n, D²}) for SP queries on any database instance, while Gibbons' uniform distinct sampling strategy as described in [19] may have an MSE of Ω(D²) on certain instances and queries, even with a sample size of n = Θ(N). In this case, our MSE is quadratically better than that of uniform distinct sampling.
- The straightforward way to dealing with SPJ queries is to first compute the join $R_1 \bowtie \cdots \bowtie R_m$, and then build the SP-query synopsis on the join results. This can be expensive, especially for multi-way cyclic joins. We extend our sampling strategy to a random walk based algorithm that can collect the sample much more efficiently, while incurring a small loss to the accuracy.
- Finally, we conducted an extensive experimental study to evaluate these algorithms on synthetic, benchmark, and real data, giving recommendations to their practical use.

 $^{^2\}mathrm{Our}$ analysis only requires h to be a pairwise-independent hash function.

Value	i = 1, 2, 3		i = 4, 5, 6		i = 7		i = 8		i = 9		i = 10								
Frequency	$N_1 =$	$= N_2 = N_3 = 1$		$N_4 = N_5 = N_6 = 2$		$N_7 = 3$		$N_8 = 5$		$N_9 = 8$		$N_{10} = 20$							
	p_i	τ_i	$p_i \tau_i$	p_i	τ_i	$p_i \tau_i$	p ₇	τ7	$p_7\tau_7$	p_8	$ au_8$	$p_8\tau_8$	p 9	τ9	<i>p</i> 9 <i>τ</i> 9	p_{10}	τ_{10}	$p_{10}\tau_{10}$	M
n = 10	0.89	1	0.89	0.63	2	1.26	0.51	3	1.54	0.40	5	1.99	0.32	0	0	0.20	0	0	8
n = 15	1	1	1	1	2	2	0.87	3	2.62	0.68	5	3.38	0.53	0	0	0.34	0	0	8
n = 20	1	1	1	1	2	2	0.93	3	2.80	0.72	5	3.62	0.57	8	4.58	0.36	0	0	9

Table 2: Example of near-optimal solutions

2 RELATED WORK

Cardinality estimation is a fundamental problem that has been studied extensively. However, most previous work only studied each relational operator in isolation.

As mentioned, the projection operator acted on its own is the standard NDV problem. This problem has a long history and is well understood by now [5, 6, 11, 13–15, 30, 41]. None of them works in conjunction with selection and/or joins, though, except the distinct sampling algorithm [19], which was reviewed in Section 1.3.

Estimating the result size of a selection operator is the classical selectivity estimation problem, and has also been widely studied. When the selection condition ϕ can be arbitrary, however, the only way to estimate its selectivity is by random sampling, which is also the approach we take in this paper. On the other hand, if ϕ is restricted to a particular form, most commonly a range condition, a host of techniques have been proposed, including sampling [7, 26, 39], histograms [20, 21, 24, 28, 34, 36, 40], and quantiles [16, 22, 23, 33].

Join size estimation is another well studied problem, and there are two main categories: sketch-based [4, 9, 10, 37] and sample-based [2, 8, 12, 17, 42]. The former usually offers better accuracy, but cannot support selection conditions. The latter may incur larger errors compared with sketch-based synopses, but can often incorporate selection conditions easily. However, none of these techniques can handle SPJ queries.

Query size estimation falls under the general umbrella of *approximate query processing (AQP)* [3, 27, 29, 32, 38, 43]. AQP techniques generally fall into two categories: query-time sampling and precomputed samples. The former takes samples according to the query, so can often return more accurate estimates, but the downside is that it needs to access the entire database at query time, which may not be desirable for use in query optimizers for large-scale distributed databases. Our sampling method falls into the category of pre-computed samples, and aims at producing query size estimates by only examining the pre-computed sample at query time.

3 OPTIMAL SAMPLING STRATEGY

In this section, we provide a near-optimal solution to problem (4). Without loss of generality, we assume $\mathbf{dom}(A) = \{1, \dots, D\}$ and their frequencies N_i are given in ascending order. Denote the optimal solution to problem (4) by $(\mathbf{p}^*, \mathbf{r}^*)$ and the optimal worst-case MSE by OPT, namely,

$$\mathrm{OPT} = \max_{\phi} f_0(\boldsymbol{p}^*, \boldsymbol{\tau}^*, \phi) = \min_{\boldsymbol{p}, \boldsymbol{\tau}} \max_{\phi} f_0(\boldsymbol{p}, \boldsymbol{\tau}, \phi) \,.$$

The main result of this section is the following characterization of a near-optimal solution to problem (4):

Theorem 3.1. There exists $0 \le M \le D$, $\kappa > 0$ and

$$p_i = \min\left\{1, \frac{\kappa}{\sqrt{N_i}}\right\}, \quad \tau_i = \begin{cases} N_i, & if \ i \leq M, \\ 0, & if \ i > M, \end{cases}$$

such that $p \cdot \tau \leq n$ and

$$\max_{\phi} f_0(\boldsymbol{p}, \boldsymbol{\tau}, \phi) \leq \text{OPT} + \sqrt{\text{OPT}} + 1.$$

The key message in this result is that the sample budget $p_i\tau_i$ should be proportional to $\sqrt{N_i}$, except for a few very large N_i 's, which we simply ignore (by setting $\tau_i=0$). Another way of looking at this result is that each tuple with A=i has a per-tuple sample budget of $p_i\tau_i/N_i\sim 1/\sqrt{N_i}$, i.e., inversely proportional to $\sqrt{N_i}$. Fundamentally, this is the balance point between two forces in a tug of war: On the one hand, a value $i\in \operatorname{dom}(A)$ will contribute to the SP query size as long as at least one tuple with A=i passes the filter ϕ , so values with higher N_i are more likely to contribute. On the other hand, all N_i tuples with the same A=i can only contribute one to the query size, no matter how many of them pass ϕ , so the per-tuple importance is actually less than those tuples with a smaller N_i . Below, we illustrate this idea with a more concrete example:

Example 2. In Table 2 we give a concrete example with D=10 distinct values and show the near-optimal solutions under different sample sizes n. The values of κ and M are found by our algorithm to be discussed in Section 4. This example illustrates that for most values i in the middle, the sample budget $p_i\tau_i$ is proportional to $\sqrt{N_i}$. For very small N_i 's, the p_i hits the ceiling of 1 and we take all their tuples; for very large N_i 's, we decide not to sample them at all, the intuition being that the sample budget they would consume is so high that it actually makes more sense to ignore them and accept the error thus caused.

Now consider applying a filter ϕ° that passes one tuple for each value i (so $D^{\phi^{\circ}} = 10$) on the sample of size n = 20. Suppose $A_s = \{1, 2, 3, 4, 5, 6, 7, 9\}$, our estimator will be

$$\hat{D}^{\phi^{\circ}} = 6 + 0.93^{-1} + 0.57^{-1} = 8.82.$$

In the rest of this section we prove Theorem 3.1. The proof involves a series of transformations starting from problem (4), each of which either preserves optimality or introduces a quantifiable small error.

3.1 Rewriting the Optimization Problem

Problem (4) is a minimax problem. To convert it to a regular optimization problem, we need to write $\max_{\phi} f_0(\pmb{p}, \pmb{\tau}, \phi)$ into an explicit form of \pmb{p} and $\pmb{\tau}$, *i.e.*, to identify the worst ϕ that maximizes f_0 for any given \pmb{p} and $\pmb{\tau}$. However, this is another optimization problem and it has no closed-form solutions. To get around this

difficulty, we first minimize $f_0(\boldsymbol{p}, \boldsymbol{\tau}, \phi^\circ)$ for a particular ϕ° , *i.e.*, find $(\boldsymbol{p}^\circ, \boldsymbol{\tau}^\circ) = \arg\min_{\boldsymbol{p}, \boldsymbol{\tau}} f_0(\boldsymbol{p}, \boldsymbol{\tau}, \phi^\circ)$. This is a regular optimization problem. Then we show that $(\boldsymbol{p}^\circ, \boldsymbol{\tau}^\circ)$ must also be an optimal solution to problem (4).

We can write \hat{D}^{ϕ} in Equation (3) as

$$\hat{D}^{\phi} = \sum_{i=1}^{D} X_i, \quad \text{where } X_i = \frac{1}{p_i} \cdot \mathbf{I}[n_i^{\phi} \ge 1].$$

Denote the expectation of X_i as a function of ϕ by

$$\mu_{i}(\phi) = \mathbf{E}[X_{i}] = \frac{1}{p_{i}} \cdot p_{i} \cdot \Pr[n_{i}^{\phi} \geq 1 \mid i \in A_{s}]$$
$$= 1 - \binom{N_{i} - N_{i}^{\phi}}{\tau_{i}} / \binom{N_{i}}{\tau_{i}},$$

where N_i^{ϕ} is the number of tuples of value i that pass ϕ . This is equivalent to the conditional probability of sampling at least one passing tuple for i given i is sampled ($i \in A_s$). Note that $\mu_i(\phi)$ is an increasing function of N_i^{ϕ} (for any fixed τ_i), and $0 \le \mu_i(\phi) \le 1$.

Since the values in dom(A) are sampled pairwise independently, we have

$$\begin{split} \mathbf{E}[\hat{D}^{\phi}] &= \sum_{i=1}^{D} \mu_{i}(\phi) \,, \quad \mathbf{Var}[\hat{D}^{\phi}] = \sum_{i=1}^{D} \left[\frac{\mu_{i}(\phi)}{p_{i}} - \mu_{i}^{2}(\phi) \right] \,, \\ \mathbf{MSE}[\hat{D}^{\phi}] &= \left[D^{\phi} - \sum_{i=1}^{D} \mu_{i}(\phi) \right]^{2} + \sum_{i=1}^{D} \left[\frac{\mu_{i}(\phi)}{p_{i}} - \mu_{i}^{2}(\phi) \right] \,. \end{split}$$

Note that for an i with $N_i^\phi=0$, we have $\mu_i(\phi)=0$, so it has no contribution to the $\mathrm{Var}[\hat{D}^\phi]$. The bias can be written as $\mathrm{Bias}[\hat{D}^\phi]=\sum_{i=1}^D (\mathrm{I}[N_i^\phi\geq 1]-\mu_i(\phi))$, so such an i has no contribution to the bias either. Therefore, the ϕ that maximizes $f_0(\pmb{p},\pmb{\tau},\phi)$ must have $N_i^\phi\geq 1$ for all i.

Consider the special ϕ° where $N_i^{\phi^{\circ}} = 1$ for all i, then $\mu_i(\phi^{\circ}) = \tau_i/N_i$, which we abbreviate to μ_i . μ_i is also the fraction of tuples we take from i given $i \in A_s$. With ϕ fixed to ϕ° , we rewrite problem (4) to the following optimization problem, where τ_i is replaced by $\mu_i N_i$:

minimize
$$p, \mu$$
 $f_1(p, \mu)$ subject to $0 , $0 \le \mu \le 1$, $\sum_{i=1}^{D} p_i \mu_i N_i \le n$, (5)$

where

$$f_1(\boldsymbol{p},\boldsymbol{\mu}) = f_0(\boldsymbol{p},\boldsymbol{\tau},\phi^\circ) = \left(D - \sum_{i=1}^D \mu_i\right)^2 + \sum_{i=1}^D \left(\frac{\mu_i}{p_i} - \mu_i^2\right).$$

3.2 Relating the Two Optimization Problems

Now we prove that an optimal solution to problem (5) is also an optimal solution to problem (4). To do this, we first identify some constraints that an optimal solution to (5) must satisfy.

First, since the constraint 0 is not a compact set, we need to argue that an optimal solution to problem (5) exists,*i.e.*, an

optimal solution cannot set some p_i infinitely close to 0. Note that $f_1(\mathbf{1}, \mathbf{0}) = D^2$, it suffices to consider solutions $(\mathbf{p}, \boldsymbol{\mu})$ with objective value at most D^2 . This means

$$D^2 \ge f_1(\boldsymbol{p}, \boldsymbol{\mu}) \ge \frac{\mu_i}{p_i} - \mu_i^2$$

for any i, which implies $p_i \ge \mu_i/(D^2 + \mu_i^2)$. The constraint on \boldsymbol{p} can therefore be replaced by

$$\frac{\mu_i}{D^2 + \mu_i^2} \le p_i \le 1.$$

Now that all the constraints are compact and f_1 is continuous, the optimal solution must exist.

Next, we present two lemmas, each of which reveals some properties of optimal solutions to problem (5).

LEMMA 3.2. Suppose $(\mathbf{p}, \boldsymbol{\mu})$ is an optimal solution to problem (5), then for any i with $\mu_i > 0$, $p_i \ge (D - \sum_{j \ne i} \mu_j)^{-1}$. Besides, if $0 < \mu_i < 1$, then $p_i = (D - \sum_{j \ne i} \mu_j)^{-1}$.

PROOF. For any i with $\mu_i > 0$, consider a new assignment (p', μ') that sets $\mu'_i = x, p'_i = p_i \mu_i / x$, and keeps all other entries unchanged. As long as $p_i \mu_i \le x \le 1$, this remains a feasible solution. Now the objective value in problem (5) is

$$\begin{split} f(x) &= \left(D - \sum_{j \neq i} \mu_i - x \right)^2 + \frac{x^2}{p_i \mu_i} - x^2 + \sum_{j \neq i} \left(\frac{\mu_j}{p_j} - \mu_j^2 \right) \\ &= \frac{x^2}{p_i \mu_i} - 2 \left(D - \sum_{i \neq i} \mu_j \right) x + C \,, \end{split}$$

where *C* is irrelevant to *x*. Since $D - \sum_{j \neq i} \mu_j \geq 1$, *f* takes the minimal value in $[p_i \mu_i, 1]$ if and only if

$$x = \min \left\{ p_i \mu_i \left(D - \sum_{i \neq i} \mu_j \right), 1 \right\}.$$

Since the optimal solution (p, μ) takes $x = \mu_i$, it must be

$$\mu_{i} = \min \left\{ p_{i} \mu_{i} \left(D - \sum_{j \neq i} \mu_{j} \right), 1 \right\}$$

$$\leq p_{i} \mu_{i} \left(D - \sum_{j \neq i} \mu_{j} \right),$$

equivalently $p_i \ge (D - \sum_{j \ne i} \mu_j)^{-1}$. Besides, if $\mu_i < 1$ then

$$\mu_i = p_i \mu_i \Big(D - \sum_{i \neq i} \mu_i \Big) < 1,$$

equivalently $p_i = (D - \sum_{j \neq i} \mu_j)^{-1}$.

LEMMA 3.3. Suppose (p, μ) is an optimal solution to problem (5), then $\mu_i = 0$ or $\mu_i = 1$ for all but one i.

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PROOF. Proof by contradiction. Suppose for some i < j, $0 < \mu_i < 1$ and $0 < \mu_j < 1$. Denote $C = D - \sum_k \mu_k > 0$. By Lemma 3.2, $p_i = (C + \mu_i)^{-1}$ and $p_j = (C + \mu_j)^{-1}$. Consider a new solution (p', μ') where

$$\mu'_{i} = \min\{\mu_{i} + \mu_{j}, 1\}, \ p'_{i} = (C + \mu'_{i})^{-1};$$

$$\mu'_{j} = \max\{\mu_{i} + \mu_{j} - 1, 0\}, \ p'_{j} = (C + \mu'_{j})^{-1};$$

$$\mu'_{k} = \mu_{k}, \ p'_{k} = p_{k} \text{ for all } k \neq i, j.$$

It is easy to verify $\mu_j' < \mu_i, \mu_j < \mu_i'$ and $\mu_i' + \mu_j' = \mu_i + \mu_j$.

We prove (p', μ') is a feasible solution. Both μ'_i and μ'_j are obviously within [0,1]. To verify p'_i and p'_j , note that

$$p_i' = (C + \mu_i')^{-1} = (D - \sum_{k \neq i,j} \mu_k - \mu_j')^{-1} \le (2 - \mu_j')^{-1} < 1 \,,$$

and similarly $p_j' \le 1$. The expected sample size strictly decreases as its change

$$\begin{split} &p_i'\mu_i'N_i + p_j'\mu_j'N_j - p_i\mu_iN_i - p_j\mu_jN_j \\ &= \frac{\mu_i'}{C + \mu_i'}N_i + \frac{\mu_j'}{C + \mu_j'}N_j - \frac{\mu_i}{C + \mu_i}N_i - \frac{\mu_j}{C + \mu_j}N_j \\ &= \frac{C(\mu_i' - \mu_i)}{(C + \mu_i)(C + \mu_i')}N_i + \frac{C(\mu_j' - \mu_j)}{(C + \mu_j)(C + \mu_j')}N_j \\ &= C(\mu_i' - \mu_i) \left[\frac{N_i}{(C + \mu_i)(C + \mu_i')} - \frac{N_j}{(C + \mu_j)(C + \mu_j')} \right] \\ &< C(\mu_i' - \mu_i) \cdot \frac{N_i - N_j}{(C + \mu_i)(C + \mu_j)} < 0 \; . \end{split}$$

Now we argue that (p', μ') is as good as (p, μ) since their objective values satisfy

$$\begin{split} f_1(\boldsymbol{p}', \boldsymbol{\mu}') - f_1(\boldsymbol{p}, \boldsymbol{\mu}) \\ &= \left[\mu_i'(C + \mu_i') + \mu_j'(C + \mu_j') - \mu_i'^2 - \mu_j'^2 \right] \\ &- \left[\mu_i(C + \mu_i) + \mu_j(C + \mu_j) - \mu_i^2 - \mu_j^2 \right] \\ &= C(\mu_i' + \mu_j' - \mu_i - \mu_j) = 0 \; . \end{split}$$

This is a contradiction since we can slightly increase p_i without violating any constraint. The objective value strictly decreases from the term μ_i/p_i , contradicting that (p, μ) is optimal.

We can now argue that solving problem (5) on the particular ϕ° is enough for solving the original problem (4).

Theorem 3.4. Let (p^*, μ^*) be an optimal solution to problem (5), then (p^*, τ^*) is an optimal solution to problem (4) where $\tau_i^* = \mu_i^* N_i$ for all i.

PROOF. Recall that ϕ° is the selection condition such that $N_i^{\phi^{\circ}} = 1$ for all i. We show that with (p^*, τ^*) fixed,

$$f_0(\boldsymbol{p}^*, \boldsymbol{\tau}^*, \phi^\circ) = \max_{\phi} f_0(\boldsymbol{p}^*, \boldsymbol{\tau}^*, \phi).$$

Let $\phi'=\arg\max_{\phi}f_0(\boldsymbol{p}^*,\tau^*,\phi)$. We show that we can modify ϕ' into ϕ° without changing the value of the target function. As observed previously, $N_i^{\phi'}\geq 1$ for all i. For those i's with $\tau_i^*=0$, setting $N_i^{\phi'}$ to 1 does not affect the objective value as f_0 only depends on $\mu_i(\phi')$, which is always 0. Similarly, if $\tau_i^*=N_i$, any $N_i^{\phi'}\geq 1$ has no effect on f_0 since $\mu_i(\phi')$ is always 1. Therefore for these values we can let $N_i^{\phi'}=1$.

these values we can let $N_i^{\phi'}=1$. Recall from Lemma 3.3 that $\mu_i^*=\tau_i^*/N_i=0$ or 1 for all but one i, so we are left with at most one i with $0<\tau_i^*< N_i$. By Lemma 3.2, this value satisfies

$$p_i^* = \left(D - \sum_{j \neq i} \frac{\tau_i^*}{N_i}\right)^{-1} = \left(1 + \sum_{j \neq i} \mathbf{I}[\tau_j^* = 0]\right)^{-1} \,.$$

We compute the partial derivative of $f_0(\mathbf{p}, \tau, \phi)$ with respect to $\mu_i(\phi)$:

$$\begin{split} &\frac{\partial f_0}{\partial \mu_i(\phi)} = -2 \Big(D - \sum_{j=1}^D \mu_j(\phi) \Big) + \frac{1}{p_i} - 2\mu_i(\phi) \\ &= -2 \Big(D - \sum_{j \neq i} \mu_j(\phi) \Big) + \sum_{j \neq i} \mathbf{I} [\tau_j^* = 0] + 1 \\ &= \sum_{j \neq i} \Big(2\mu_j(\phi) + \mathbf{I} [\tau_j^* = 0] - 2 \Big) - 1 \\ &= \sum_{j \neq i} \Big(2\mu_j(\phi) \cdot \mathbf{I} [\tau_j^* > 0] + \mathbf{I} [\tau_j^* = 0] - 2 \Big) - 1 < 0 \; . \end{split}$$

This means $f_0(\boldsymbol{p}^*, \boldsymbol{\tau}^*, \phi)$ is maximized when $\mu_i(\phi)$ takes the smallest possible value. Since $\mu_i(\phi)$ is an increasing function of N_i^{ϕ} , we must have $N_i^{\phi'}=1$. Thus, we conclude that $\phi'=\phi^{\circ}$ maximizes $f_0(\boldsymbol{p}^*, \boldsymbol{\tau}^*, \phi)$.

Now, for any valid solution (p', τ') to problem (4), we have

$$\max_{\phi} f_0(\boldsymbol{p}', \boldsymbol{\tau}', \phi) \ge f_0(\boldsymbol{p}', \boldsymbol{\tau}', \phi^{\circ}) = f_1(\boldsymbol{p}', \boldsymbol{\mu}')$$

$$\ge f_1(\boldsymbol{p}^*, \boldsymbol{\mu}^*) = f_0(\boldsymbol{p}^*, \boldsymbol{\tau}^*, \phi^{\circ}) = \max_{\phi} f_0(\boldsymbol{p}^*, \boldsymbol{\tau}^*, \phi),$$

so (p^*, τ^*) is also an optimal solution to problem (4).

3.3 A Near-Optimal Solution

Problem (5) has linear constraints and a closed-form objective function, but the objective function is non-convex. General solvers for such optimization problems only return a local-optimal solution without any guarantee on its global optimality. Furthermore, they usually use iterative algorithms that can be slow on large instances. In this section, we exploit the special structure of problem (5) and transform it to an easier one that admits a near-optimal solution.

By Lemma 3.3, for any optimal solution, there is at most one i with $0 < \mu_i < 1$. For simplicity, we also sets $\mu_i = 0$ for this i, i.e., we only consider solutions such that $\mu_i \in \{0,1\}$ for all i. The problem now becomes the following:

minimize
$$f_1(\boldsymbol{p}, \boldsymbol{\mu})$$

subject to $0 < \boldsymbol{p} \le 1$,
 $\boldsymbol{\mu} \in \{0, 1\}^D$, (6)

$$\sum_{i=1}^D p_i \mu_i N_i \le n$$
.

We quantify its near-optimality by the following lemma.

Lemma 3.5. Let (p, μ) be the optimal solution to problem (6), and OPT = $f_1(p^*, \mu^*)$ be the optimal objective value of problem (5), then

$$f_1(\boldsymbol{p}, \boldsymbol{\mu}) \leq \text{OPT} + \sqrt{\text{OPT}} + 1$$
.

PROOF. First note that the objective value for any feasible solution (p', μ') is lower bounded by

$$f_1(\mathbf{p}', \boldsymbol{\mu}') \ge (D - \|\boldsymbol{\mu}'\|_1)^2$$
.

Let (p^*, μ^*) be an optimal solution to problem (5). According to Lemma 3.3, $\mu_i^* = 1$ or $\mu_i^* = 0$ for all but possibly one *i*. If there is

no exception then we are done. Otherwise for this 0 < $\mu_i^* <$ 1, by Lemma 3.2 we have

$$p_i^* = (D - \sum_{i \neq i} \mu_i^*)^{-1} = (D - \|\boldsymbol{\mu}^*\|_1 + \mu_i^*)^{-1} \,.$$

Now consider a new solution (p^*, μ') that sets $\mu'_i = 0$ and keeps $\mu'_j = \mu^*_j$ for all $j \neq i$. Plug in the objective function f_1 , we have

$$f_{1}(\boldsymbol{p}^{*}, \boldsymbol{\mu}') - f_{1}(\boldsymbol{p}^{*}, \boldsymbol{\mu}^{*})$$

$$= \left[(D - \|\boldsymbol{\mu}^{*}\|_{1} + \mu_{i}^{*})^{2} - (D - \|\boldsymbol{\mu}^{*}\|_{1})^{2} \right] - \left[\frac{\mu_{i}^{*}}{p_{i}^{*}} - (\mu_{i}^{*})^{2} \right]$$

$$= \mu_{i}^{*}(D - \|\boldsymbol{\mu}^{*}\|_{1} + \mu_{i}^{*}) < \sqrt{f_{1}(\boldsymbol{p}^{*}, \boldsymbol{\mu}^{*})} + 1.$$

But now (p^*, μ') is in the feasible region of problem (6), whose optimal solution is (p, μ) , we must have

$$f_1(\mathbf{p}, \boldsymbol{\mu}) \leq f_1(\mathbf{p}^*, \boldsymbol{\mu}') < f_1(\mathbf{p}^*, \boldsymbol{\mu}^*) + \sqrt{f_1(\mathbf{p}^*, \boldsymbol{\mu}^*)} + 1.$$

Now we focus on solving problem (6). We first give a characterization of its optimal solutions, similar to Lemma 3.3.

LEMMA 3.6. There exists an optimal solution $(\mathbf{p}, \boldsymbol{\mu})$ to problem (6) such that $\mu_i = 1$ for $i \leq M$ and $\mu_i = 0$ for i > M, for some $M \in \{0, 1, ..., D\}$.

PROOF. For any solution (p, μ) , if $\mu_i = 0$ and $\mu_j = 1$ for some i < j, we can swap (p_i, μ_i) with (p_j, μ_j) so that the new solution is still feasible and the MSE does not change. By repeating the process at most D times, we can put all $\{i \mid \mu_i = 1\}$ in front of $\{i \mid \mu_i = 0\}$.

Thus the problem of finding μ is reduced to finding $M = \|\mu\|_1$, whose possible values are $0, \ldots, D$. Suppose we have obtained an optimal M. We still need to find p. For any given M, this is another optimization problem:

minimize
$$f_2(\mathbf{p}; M) = \sum_{i=1}^{M} \frac{1}{p_i}$$

subject to $\mathbf{0} < \mathbf{p} \le \mathbf{1}$, (7)
 $\sum_{i=1}^{M} p_i N_i \le n$.

Note that f_2 and f_1 differ only in terms independent of p, so it is sufficient to minimize f_2 . Without the constraint $p \le 1$, problem (7) can be solved easily by the Cauchy-Schwartz inequality, and the optimal solution is $p_i \sim 1/\sqrt{N_i}$. The constraint $p \le 1$ has introduced some technical complications, but we still manage to get the following closed-form solution.

Lemma 3.7. An optimal solution to problem (7) is

$$p_i = \min\left\{1, \frac{\kappa}{\sqrt{N_i}}\right\},\,$$

where $\kappa = \frac{n - \sum_{j=1}^{K} N_j}{\sum_{i=K+1}^{M} \sqrt{N_j}}$, where K is the largest integer such that

$$n - \sum_{i=1}^{K} N_i > \sqrt{N_K} \sum_{i=K+1}^{M} \sqrt{N_i}.$$
 (8)

If
$$n \leq \sqrt{N_1} \sum_{i=1}^{M} \sqrt{N_i}$$
, we set $K = 0$.

PROOF. We focus on $p_1 \dots p_M$ as the remaining are irrelevant. First note that there exists an optimal solution to problem (7) where $\{p_i\}$ is descending, otherwise we may swap a reverse pair so that the solution is still feasible and the objective value $\sum p_i^{-1}$ does not change. Let p be an optimal solution where $\{p_i\}$ is descending, and K be the largest integer such that

$$n - \sum_{i=1}^{K} p_i N_i > \sqrt{N_K} \sum_{i=K+1}^{M} \sqrt{N_i},$$

which implies

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$$n - \sum_{i=1}^{K} p_{i} N_{i} \leq p_{K+1} N_{K+1} + \sqrt{N_{K+1}} \sum_{i=K+2}^{M} \sqrt{N_{i}}$$

$$\leq \sqrt{N_{K+1}} \sum_{i=K+1}^{M} \sqrt{N_{i}}.$$
(9)

We now show that $p_K = 1$.

Note the values p_{K+1}, \ldots, p_M satisfy

$$\sum_{i=K+1}^M p_i N_i = n - \sum_{j=1}^K p_j N_j \,.$$

By Cauchy-Schwarz inequality, we know

$$\sum_{i=K+1}^{M} \frac{1}{p_i} \ge \frac{(\sum_{j=K+1}^{M} \sqrt{N_j})^2}{n - \sum_{j=1}^{K} p_j N_j}$$

where minimum is taken exactly when

$$p_{i} = \frac{n - \sum_{j=1}^{K} p_{j} N_{j}}{\sqrt{N_{i}} \sum_{j=K+1}^{M} \sqrt{N_{j}}}, \quad i > K.$$
 (10)

By Inequality 9 $p_i \le 1$, so they are feasible. Therefore these are exactly the values for p_{K+1}, \ldots, p_M due to optimality. The minimum objective value is thus

$$\sum_{i=1}^{K} \frac{1}{p_i} + \frac{(\sum_{j=K+1}^{M} \sqrt{N_j})^2}{n - \sum_{i=1}^{K} p_j N_j}.$$

Now we increase p_K and decrease p_{K+1}, \ldots, p_M correspondingly according to Equation 10. The solution will still be feasible. We view the objective value as a function of p_K , and calculate

$$\frac{\partial}{\partial p_K} \left(\sum_{i=1}^K \frac{1}{p_i} + \frac{(\sum_{j=K+1}^M \sqrt{Nj})^2}{n - \sum_{j=1}^K p_j N_j} \right) < 0 \; .$$

Increasing p_K will reduce the objective value, which contradicts optimality. Thus we conclude $p_K = 1$. Since $\{p_i\}$ is descending, we have

$$p_i = \begin{cases} 1, & i \leq K, \\ \frac{n - \sum_{j=1}^K N_j}{\sqrt{N_i} \sum_{j=K+1}^M \sqrt{N_j}}, & i > K. \end{cases}$$

By setting $\kappa = \frac{n - \sum_{j=1}^{K} N_j}{\sum_{i=K+1}^{M} \sqrt{N_i}}$, we prove that $p_i = \min\{1, \kappa/\sqrt{N_i}\}$. \square

Combining Theorem 3.4, Lemma 3.5, 3.6, and 3.7, we obtain Theorem 3.1. In addition, these lemmas yield a method to find M and κ , although not very efficient: For each $M = 0, 1, \ldots, D$, we use Lemma 3.7 to find the optimal p. Then we choose the best M.

4 EFFICIENT ALGORITHMS

Recall that we assume $N_1 \leq N_2 \leq \cdots \leq N_D$ are given in sorted order. The straightforward algorithm described above has a running time of $O(D^2)$. For each M, we spend O(D) time to find the corresponding K (which we denote as K_M to stress that it depends on M) and p, as specified in Lemma 3.7. Then we calculate the objective function

$$f_3(M) = (D - M)^2 + \frac{(\sum_{i=K_M+1}^M \sqrt{N_i})^2}{n - \sum_{i=1}^{K_M} N_i} + K_M - M,$$

and return the M with the minimum $f_3(M)$.

In this section, we first give an improved algorithm with running time O(D). Then we also show how the algorithm can be made to work if only a histogram is given in lieu of all value frequencies.

4.1 An O(D)-time Algorithm

We see that the bottleneck of the above algorithm is in finding all the K_M 's. To improve it, we make the key observation that K_M is non-increasing with respect to M.

Let M^* be the optimal M minimizing $f_3(M)$, and let M_0 be the largest integer such that $\sum_{i=1}^{M_0} N_i \le n$. Note that it must be the case $M^* \ge M_0$, otherwise we could have added more tuples into the sample. By the definition of K_M , $n - \sum_{i=1}^{K_M} N_i > 0$, so $K_M \le M_0$ for all M. Also, when $M > M_0$,

$$n - \sum_{i=1}^{K_M} N_i > \sqrt{N_{K_M}} \sum_{i=K_M+1}^M \sqrt{N_i} \ge \sqrt{N_{K_M}} \sum_{i=K_M+1}^{M-1} \sqrt{N_i},$$

so $K_M \leq K_{M-1}$.

This means that we can find K_M incrementally for $M = M_0, \ldots, D_n$, starting from $K = M_0$. This suggests the O(D)-time algorithm as described in Algorithm 1. After M^* and K_{M^*} are found, we can calculate κ , p and τ in O(D) time following Lemma 3.7.

Example 3. We illustrate how Algorithm 1 works using the example in Table 2 with n = 20. The algorithm first finds $M_0 = 8$, as the first 8 N_i 's add up to 17, while adding N_9 would exceed n.

- When M = 8, we find $K_M = 8$, and $f_3(M) = (10 8)^2 + 0 = 4$.
- When M = 9, we reduce K_M to 6, and update $f_3(M)$ as

$$f_3(9) = (10 - 9)^2 + \frac{(\sqrt{3} + \sqrt{5} + \sqrt{8})^2}{20 - 9} + 6 - 9 = 2.20$$
.

The algorithm maintains the prefix sum $\sum N_i$ and the suffix sum $\sum \sqrt{N_i}$ so that both K_M and $f_3(M)$ can be updated incrementally.

• When M = 10, we reduce K_M to 3, while $f_3(M)$ is updated to

$$f_3(10) = 0^2 + \frac{(3\sqrt{2} + \sqrt{3} + \sqrt{5} + \sqrt{20})^2}{20 - 3} + 3 - 10 = 2.46$$
.

The optimal value is therefore taken at M = 9.

```
Algorithm 1: Find M^* and K^* = K_{M^*}

Input : D, \{N_i\}_{i=1}^D and n

Output: M^* and K^* = K_{M^*}
```

 $M_0 \leftarrow 0$, $sum \leftarrow 0$;

while $M_0 \le D$ and $n \ge sum$ do

$$M_0 \leftarrow M_0 + 1;$$

 $sum \leftarrow sum + N_{M_0};$

 $sum \leftarrow sum - N_{M_0}, M_0 \leftarrow M_0 - 1;$

 $M^* \leftarrow 0, K^* \leftarrow 0, min_mse \leftarrow +\infty;$

 $K \leftarrow M_0$, $sum_sqrt \leftarrow 0$;

for $M \leftarrow M_0$ to D do

return M^* and K^* ;

THEOREM 4.1. There is an O(D)-time algorithm that, given $\{N_i\}$ in ascending order, finds a near-optimal solution to problem (4).

4.2 Histogram Approximation

 $sum_sqrt \leftarrow sum_sqrt + \sqrt{N_M}$

If not all frequencies N_i are available or expensive to obtain, our algorithm can also work with a heavy hitters histogram, which is commonly maintained by database systems. A heavy hitters histogram only maintains the H values in $\mathbf{dom}(A)$ with the largest frequencies, namely, N_{D-H+1}, \ldots, N_D . Then, for the low-frequency values, we simply assume that they have equal frequency, i.e., we use

$$\hat{N_L} = \frac{N - \sum_{j=D-H+1}^D N_j}{D-H}$$

as an approximation of N_i for i = 1, ..., D - H.

Naively running Algorithm 1 after replacing N_i with \hat{N}_L for $i=1,\ldots,D-H$ would still take O(D) time. Below, we show how to improve it to O(H). Consider two cases. In the first case $n \le (D-H)\hat{N}_L$, we must have $M_0 = n/\hat{N}_L$, therefore

$$n = M_0 \hat{N}_L \le M \hat{N}_L \le \sqrt{N_1} \sum_{i=1}^M \sqrt{N_i},$$

which implies K=0. When $M \leq D-H$, $f_3(M)$ becomes a quadratic function whose minimal value can be computed in O(1) time, and when M>D-H, we only need to search M from D-H to D with the knowledge $K_M=0$, which requires O(H) time. In the second case $n>(D-H)\hat{N}_L$, we have $M_0\geq D-H$. Note that when $K\leq D-H$, inequality (9) simplifies to

$$n > (D-H) \cdot \hat{N}_L + \sqrt{\hat{N}_L} \sum_{i=D-H+1}^{M} \sqrt{N_i},$$

which is irrelevant to K, meaning $K_M = 0$ or $K_M \ge D - H$. Therefore we only need to search M from M_0 to D and K_M in $\{M_0, M_0 - 1, \dots, D - H, 0\}$, which requires O(H) time as $\{K_M\}$ is non-increasing.

5 MSE ANALYSIS

Our algorithm finds a near-optimal weighted distinct sampling strategy. However, we still need to answer the question: how good is its MSE, in particular, in comparison with uniform distinct sampling [19]? In this section, we give a theoretical analysis; results from empirical studies will be presented in Section 7.

Worst-case data distribution. We first show that the MSE of our optimal weighted distinct sampling strategy is $O(\min\{DN/n, D^2\})$ in the worst case.

THEOREM 5.1. For any input data and SP query, the MSE of the near-optimal weighted distinct sampling strategy found by our algorithm is bounded by $O(\min\{DN/n, D^2\})$.

PROOF. It is sufficient to find one valid solution (p, μ) to problem (4) such that $f_1(p, \mu) = O(\min\{DN/n, D^2\})$. The near-optimal solution found by our algorithm can only be better, modulo a lower-order term.

Note that the min takes O(DN/n) for $n \ge N/D$ and $O(D^2)$ for n < N/D. In the former case, consider the solution (p, μ) where $p_i = n/N$ and $\mu_i = 1$ for all i. It can be verified this is a feasible solution to problem (4), and

$$f_1(\mathbf{p}, \boldsymbol{\mu}) = 0 + \sum_{i=1}^{D} \left(\frac{N}{n} - 1 \right) = O(DN/n).$$

In the latter case, we simply set $\mu_i = 0$ for all i, and $f_1(\boldsymbol{p}, \boldsymbol{\mu}) = O(D^2)$.

The bound given in Theorem 5.1 is tight in the worst case. The hardest input distribution is simply the uniform distribution: $N_i = N/D$ for all i, and the selection condition is $\phi = \phi^{\circ}$, i.e. $N_i^{\phi^{\circ}} = 1$ for all i. In the optimal solution, $K_M = 0$, and

$$f_3(M) = (D - M)^2 + \frac{N}{nD} \cdot M^2 - M$$
$$\ge \frac{(N/n - 1)D + 1/4}{1 + N/(nD)}.$$

When n > N/D, it is $\Omega(DN/n)$, and when $n \le N/D$, it is $\Omega(D^2)$. In fact, it is not surprising that uniform distribution is the worst-case input for weighted distinct sampling; on uniform data, all distinct values are equally important, losing the advantage of weighted sampling.

Analysis of Gibbons' algorithm. On the other hand, the MSE of the uniform distinct sampling strategy of Gibbons [19] is $\Omega(D^2)$ on certain datasets and queries, even with a sample size of $n = \Omega(N)$, as we demonstrate below.

In fact, Gibbons' algorithm [19] assumes that the selectivity q of the condition ϕ , *i.e.*, $q = |\sigma_{\phi}R|/N$, is given in advance, whereas our algorithm makes no assumption on ϕ . We show that even with q given, the MSE of Gibbons' algorithm is still $\Omega(D^2)$.

Gibbons' algorithm sets $\tau = \min\{2/q, n/50\}$ for all i, and adaptively chooses p such that the sample size is the given budget n.

Consider a table in which $N_i=3N/D$ for $i\leq D/4$ and $N_i=N/(3D)$ for i>D/4. The selection condition is again $\phi=\phi^\circ$, *i.e.* $N_i^{\phi^\circ}=1$ for all i. Note that the selectivity of ϕ° is q=D/N. Assume $n=\Omega(N)$. Thus algorithm sets $\tau=\min\{2/q,n/50\}=2/q=2N/D$. We have for $i\leq D/4$, the conditional probability

$$\mu_i = \Pr[n_i^{\phi^{\circ}} \ge 1 \mid i \in A_s] = \tau/N_i \le \frac{2}{3}.$$

As a result,

$$\begin{split} \mathbf{E}[\hat{D}^{\phi^{\circ}}] &= \frac{1}{p} \sum_{i=1}^{D} \mathbf{E}[\mathbf{I}[n_{i}^{\phi^{\circ}} \geq 1]] = \sum_{i=1}^{D} \mu_{i} \\ &\leq \frac{3D}{4} + \frac{D}{4} \cdot \frac{2}{3} = \frac{11}{12}D. \end{split}$$

Equivalently,

$$\mathbf{MSE}[\hat{D}^{\phi^{\circ}}] \geq \mathbf{Bias}^{2}[\hat{D}^{\phi^{\circ}}] \geq \left(\frac{11}{12}D - D\right)^{2} = \Omega(D^{2}).$$

Recall that the worst-case MSE of our weighted distinct sampling strategy is O(DN/n) when $n \ge N/D$. Therefore, for any sample size $n = \omega(N/D)$, we obtain an asymptotic improvement over Gibbons' uniform distinct sampling algorithm, and the improvement is quadratic when $n = \Theta(N)$.

Analysis on Zipfian distribution. Our earlier analysis indicates that uniform distribution is the hardest input for our algorithm. Thus, we also tried to see if the algorithm could perform better on skewed data. For this purpose, we analyzed its MSE on Zipfian distribution, a commonly used distribution for modeling skewed data. In the Zipfian distribution, the frequency of the *i*-th most popular value has frequency proportional to $i^{-\alpha}$, for a parameter α . Larger α indicates higher skewness. Note that when $\alpha = 0$, it degenerates into the uniform distribution.

Our analytical results are given in Table 3, which indeed shows that the MSE reduces as data skewness increases: When $0 \le \alpha < 1$, the MSE is the same as the uniform case. When $\alpha > 2$, the MSE becomes O(N/n). This means that a constant-fraction (say, 1%) sample can already achieve to a constant additive error.

Range of α	Range of n	Asymptotic MSE
[0, 1)	(0, N)	DN/n
{1}	$(0, N/\ln D]$	$(N/n)(D/\ln D)$
11)	$(N/\ln D, N)$	$D^{1-n/N}$
(1, 2)	$(0, N/D^{\alpha-1}]$	$(N/n)D^{2-\alpha}$
(1, 2)	$(N/D^{\alpha-1},N)$	$(N/n)^{1/(\alpha-1)}$
(3)	$(0, N \ln D/D]$	$(N/n)\ln^2 D$
{2}	$(N \ln D/D, N)$	$(N/n)\ln^2(N/n)$
(2, +∞)	(0,N)	N/n

Table 3: MSE analysis on Zipfian distribution

6 SPJ QUERIES

A straightforward way of handling SPJ queries is to first compute the join $R_1 \bowtie \cdots \bowtie R_m$, and then run our optimal weighted distinct sampling algorithm on the join results. This way, an SPJ query becomes a SP query, and all our MSE analysis from the previous section will hold. In particular, we can guarantee a maximum MSE of $O(DN_{\bowtie}/n)$ for SPJ queries over any database instance and any

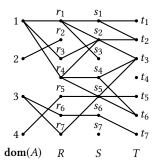


Figure 1: Example with the conceptual column.

 ϕ , where $N_{\bowtie} = |R_1 \bowtie \cdots \bowtie R_m|$ is the join size. However, computing the full join and running the algorithm on all join results can be very expensive, especially for multi-way joins. In this section, we describe a random walk based algorithm that collects the sample in just one pass over the data, without performing the join. The resulting sample is not optimal, though, but we demonstrate through our experiments that the accuracy loss is still small enough in comparison to the significant savings in the sample construction cost.

6.1 Random Walk Algorithm

Our random walk algorithm is inspired by the Wander Join algorithm of Li et al. [32]. However, Wander Join does not support projections. Thus, we make the following modifications. To make things more concrete, consider a 3-way line join $R \bowtie S \bowtie T$, where the projection attribute A belongs to R. We conceptually add one more table that consists of a single attribute and D tuples, one for each distinct value in $\operatorname{dom}(A)$ (see Figure 1). For each tuple $r \in R$ with $\pi_A(r) = i$, we add an edge between value i and tuple r. We also add an edge between any two tuples that can join. Then, the idea is to start the random walks from the conceptual table $\operatorname{dom}(A)$ and walk towards T, where each random walk yields a sample from the join results.

Another major difference is that Wander Join [32] takes samples at query time, which requires indexes to be available, and results in a lot of random accesses in the database at query time. This does not fit the scenario of query optimization, which is the main application of query size estimation. Here, we would like to have a pre-computed small synopsis from which the query size can be estimated quickly without having to rely on indexes.

Below, we describe our random walk algorithm for obtaining a weighted distinct sample from a multi-way join. It does not need indexes, either, and the sample can be collected in one pass over each relation.

We first compute the optimal sampling strategies p, τ using the frequencies of values of $\mathbf{dom}(A)$ in R as described in Section 3, 3 and collect a distinct sample on R. Suppose i=1 has been sampled, and $\tau_1=3$ tuples r from R have been sampled with $\pi_A(r)=i$. Next, we extend each of the τ_1 tuples into a join result, sampled from all join results whose projection on A is i. This corresponds to the subgraph rooted at i=1 (see Figure 2).

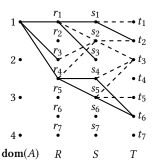


Figure 2: Distinct sampling from a multi-way join.

As we may encounter a "deadend" when extending a tuple into a join result, such as (r_1, s_3) in Figure 2, we take the following twophase approach. We use a space budget of $\tau_i' = c\tau_i$ for each distinct $i \in \mathbf{dom}(A)$ sampled from R, where c > 1 is a constant scaling parameter. We are in phase one as long as the space budget is not reached. In phase one, we try to obtain a random sample of size up to τ_1' from the partial join results $\sigma_{A=1}(R)\bowtie S$, using reservoir sampling. Consider i = 1 and suppose $\tau'_1 = 4$ in Figure 2. Our initial random walks consist of 3 tuples in R, which is smaller than $\tau'_1 = 4$, so we are in phase one. Then when scanning S, for each $s_i \in S$, we join it with the current sample from R, and feed the join results to reservoir sampling. If $|\sigma_{A=1}(R) \bowtie S| \le \tau'_1$, we take all the join results into the sample and remain in phase one; otherwise we go to phase two. In Figure 2, $|\sigma_{A=1}(R)| \leq S = 7 > 4$, so only 4 partial join results have been sampled (solid lines), and we move onto phase two when scanning T.

We will be in phase two when scanning T. In phase two, for each partial random walk (r, s), we extend it to one tuple in T, randomly chosen from all those that join with s (if such tuples exist). An example is shown in Figure 2, where 3 join results, (r_1, s_1, t_2) , (r_4, s_4, t_6) and (r_4, s_5, t_6) , have been eventually sampled. Note that (r_1, s_3) fails to extend to T, which is why we use a slightly larger space budget τ_1' . In the end, if more than τ_1 join results have been obtained, we sub-sample them down to τ_1 .

For general multi-way joins, we reorder relations so that each relation (except the first one) shares at least one common attribute with a previous relation. We then scan the relations one by one in this order. When scanning, we only take tuples that join with the corresponding tuples sampled from all previous relations so far. For example, if there is also a join condition between R and T, then when extending a partial random walk (r,s) to a $t \in T$, we only sample from all tuples $t \in T$ such that t joins with both r and s.

Finally, we run the algorithm above for all $i \in \mathbf{dom}(A)$ sampled in the first relation in parallel. So the algorithm makes one pass over each relation (i.e., its time complexity is *only linear to the sum of all relation sizes from the join*), using space cn = O(n), where n is the total sample size.

6.2 Bias Reduction

Another technical difference between our SPJ algorithm and SP algorithm as described in Section 3 is that, in the SP algorithm, τ_i is either N_i or 0, so individual estimators have no bias. In the random walk algorithm, however, for a sampled $i \in A_s$, only up to τ_i of its join results are sampled. As such, there is a downward bias

³Essentially, we use the frequencies of values in *R* to approximate their frequencies in the full join results, which is why we cannot guarantee optimality of the sampling strategy.

in the estimator. To see this problem more clearly, for a selection condition ϕ , let $\mu_i(\phi)$ be the probability that any sampled tuples for *i* pass ϕ , given *i* is sampled. Then $\hat{D}^{\phi} = \sum_{i=1}^{D} Y_i$ would be an unbiased estimator of D^{ϕ} , where

$$Y_i = \frac{\mathbf{I}[n_i^{\phi} \ge 1]}{p_i \mu_i(\phi)} .$$

However, as illustrated in Example 1, the problem is that $\mu_i(\phi)$ is unknown⁴. Due to the uncertainty in $\mu_i(\phi)$, bias cannot be eliminated. Nevertheless, we introduce the following heuristic to reduce it to some extent.

Our idea is to reduce the bias based on the value of n_i^{ϕ} , which is known. If $n_i^{\phi} \geq 2$, then N_i^{ϕ} is also likely large, so the chance that some tuple passes ϕ is also large, so we take $\mu_i(\phi) = 1$. If $n_i^{\phi} = 1$, then $\mu_i(\phi)$ is likely small. We observe that

$$p_t \leq \mu_i(\phi) \leq 1$$
,

where p_t is the probability that the only tuple t in the sample that passes ϕ was sampled when the sample was initially collected, conditioned upon i being sampled. Then we strike a balance between the two extremes by taking geometric mean and setting $\mu_i(\phi) \approx \sqrt{p_t}$.

To summarize, our biased-reduce estimator is $\hat{D}^{\phi} = \sum_{i=1}^{D} \hat{Y}_{i}$, where

$$\hat{Y}_i = \begin{cases} 0, & \text{if } n_i^\phi = 0\,,\\ \frac{1}{p_i\sqrt{p_t}}, & \text{if } n_i^\phi = 1\,,\\ \frac{1}{p_i}, & \text{if } n_i^\phi > 1\,. \end{cases}$$

The remaining issue is how to compute p_t . Recall our two-phase sample collection algorithm. If the algorithm terminates during phase one, then p_t is the same for all tuples with A = i:

$$p_t = \frac{\tau_i'}{|\sigma_{A=i}(R_1) \bowtie \cdots \bowtie R_m|}.$$

Note that the partial join size $|\sigma_{A=i}(R_1)|\bowtie \cdots \bowtie R_m|$ can be computed when we scan R_m .

If the algorithm goes on to phase two, then p_t might be different for different t. Suppose phase one terminates after scanning R_s , for some s < m. Consider a tuple $t = r_1 \bowtie r_2 \bowtie \cdots \bowtie r_m$ sampled in the end, where $r_j \in R_j$. First, (r_1, \ldots, r_s) are sampled during phase one, so the probability that it is sampled is $\frac{\tau'_i}{|\sigma_{A=i}(R_1)\bowtie \cdots \bowtie R_s|}$ During phase two, we perform a series of random walks, where in each step, we pick a tuple uniformly at random from all tuples that join with previous tuples, so

$$p_t = \frac{\tau_i'}{|\sigma_{A=i}(R_1) \bowtie \cdots \bowtie R_s|} \cdot \prod_{j=s+1}^m \frac{1}{|r_1 \bowtie \cdots \bowtie r_{j-1} \bowtie R_j|} .$$
(11)

For example, in Figure 2, for $t = (r_4, s_5, t_6)$, we have

$$p_t = \frac{4}{7} \cdot \frac{1}{3} = \frac{4}{21}.$$

Finally, note that p_t 's for all the sampled tuples t can be computed easily during the sample collection process.

EXPERIMENTAL EVALUATION

In this section, we report the results from an extensive experimental evaluation on various cardinality estimation methods for both SP queries and SPJ queries.

Methods Evaluated

Exact. The exact method has been implemented to provide a reference. Note that computing the exact query size for SPJ queries requires the entire data set and is very slow.

Upper Bound (UB). As mentioned in Section 1, a simple and commonly used method for estimating the size of SPJ queries is to estimate the NDV of A and the size of the remaining query (without the projection) separately, and then return the smaller of the two. In our evaluation, we implement this method in its best-case scenario, i.e., we use the exact NDV and the remaining query size, and return the smaller of the two.

Uniform Distinct Sampling (UDS). Uniform Distinct Sampling (UDS) refers to the algorithm described in [19] for estimating SP query sizes. For SPJ queries, we compute the full join first, and then run this algorithm on the join results. MurmurHash3, implemented in SMHasher⁵, is used as the hash function for deciding if a value in dom(A) is sampled.

Weighted Distinct Sampling (WDS). This refers to our weighted distinct sampling algorithm using the optimal sampling strategy as described in Section 3. For SPJ queries, similar to UDS, we compute the full join and run the algorithm on the join results.

Histogram Approximation (HISTO). We have also implemented the algorithm in Section 4.2, where the algorithm is only given a heavy hitters histogram as an approximation of the value frequencies. The histogram contains the frequencies of the top 10% most frequent values.

Random Walk Algorithm (RW). The random walk algorithm in Section 6 can be considered an approximate version of WDS for SPJ queries. The sample can be collected in one pass over the data, at the expense of some loss in accuracy.

We implemented all the algorithms above in C++. Each relation is stored as a std::vector of std::tuple's. The experiments were carried out on a machine with an Intel(R) 2.10GHz Xeon(R) Silver 4116 CPU running CentOS Linux 7.7. The memory is large enough to hold all input relations and join results. For each dataset, we run the algorithm 30 times using independent random seeds, and report the MSE from these 30 runs. The sample size is set the same for all algorithms, which is 1% of the data size by default.

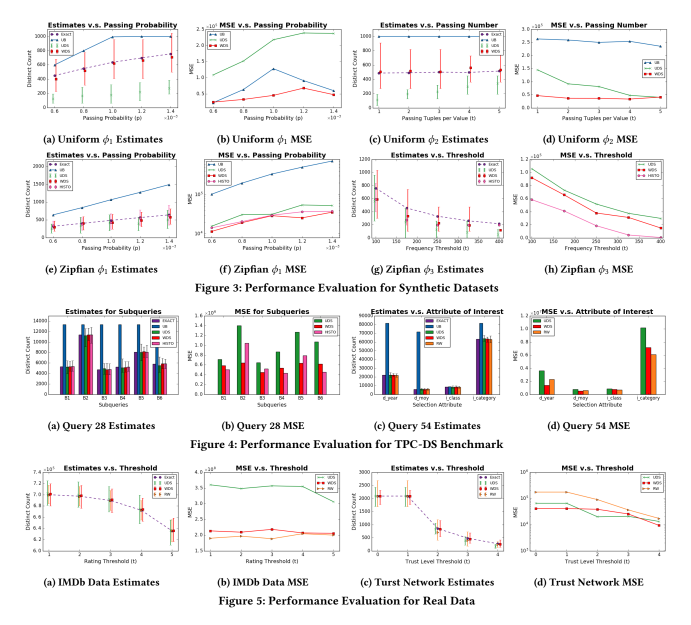
We used a wide range of datasets in our evaluation, including synthetic datasets with different skewness, the TPC-DS benchmark, and two real-world datasets. Below, we describe these datasets and queries in turn, followed by our experimental findings.

7.2 Results on Synthetic Datasets

Datasets. Our theoretical analysis shows that WDS works better on more skewed data, so we used two synthetic datasets, one uniform and the other following a Zipfian distribution, to validate this claim.

⁴More precisely, it is always 1 if all tuples with A = i are sampled; otherwise, it depends on N_i^{ϕ} , which is unknown.

⁵https://github.com/aappleby/smhasher



The table has roughly one million tuples. The uniform dataset contains D=1000 distinct A-values, each appearing in 1000 tuples. The Zipfian data have a skewness factor $\alpha=1.1$. The distinct values are integers ranging from 1 to 50000, and the frequency of value i is proportional to $1/i^{\alpha}$.

We introduced a second attribute B, which is used in the selection condition ϕ . We tested three conditions. The first one represents an average case that is considered easy, while the other two are designed to model worst-case scenarios.

- (1) For both uniform and Zipfian data, ϕ_1 passes each tuple independently with the same probability p. Essentially, p corresponds to the selectivity of the query.
- (2) ϕ_2 applies on the uniform dataset. First, we pick a random value from the D distinct values of A, and pass all its tuples. Then, we randomly take a set of D/2 values randomly from

- the remaining D-1 distinct values, and passes t tuples for each. Note that $D^{\phi_2}=D/2+1$. The smaller t is, the more difficult the query is considered, as it is more difficult to catch those passing tuples by sampling.
- (3) ϕ_3 applies on the Zipfian dataset. It passes all tuples of the most frequent value. It also passes a single tuple for each value whose frequency is at least t. As with ϕ_2 , the smaller t is, the more difficult the query will be.

Results on uniform data. Figure 3a shows the estimates of query ϕ_1 returned by various algorithms on the uniform dataset, where we vary the selectivity p. Thus, UB simply reports $\min\{D, pN\}$, which is shown as straight line turning at $p = D/N = 10^{-3}$. UDS and WDS are both sampling algorithms, so in addition to the mean, we also plot the 10% and 90% percentiles of the 30 runs. Figure 3b shows the MSE of these algorithms. From these results, we see that on

this easy query, UB actually performs quite well, while UDS is even worse than UB due to large underestimation. On the other hand, WDS has generally outperformed UB in this case. The performance of WDS is more stable across different values of p, while UB is a good estimator for either small p (when pN is a good estimate since very likely, no A-value has more than one tuple passing ϕ_1) or large p (when D is a good estimate, since almost every distinct A-value has at least one tuple passing ϕ_1).

On ϕ_2 , which is a hard query, the situation becomes quite different (see Figure 3c and 3d). Since more than D tuples in total pass ϕ_2 , UB always returns D, which is a bad estimate. On the other hand, the two sampling algorithms give more accurate estimates. Between the two, we see that WDS has stable performance across different values of t, and is always better than UDS. For larger values of t, the two have similar MSE (UDS has larger bias and smaller variance, while WDS has smaller bias but larger variance), but the accuracy of UDS drops as t gets smaller (the query gets more difficult).

Results on Zipfian data. Figure 3e to 3h show the results on the Zipfian dataset. Since Zipfian distribution produces a skewed dataset with a long tail, UB fails miserably; we cannot even plot its results on the hard query ϕ_3 . The performance of HISTO is similar to WDS, showing that our algorithm still works well even if only a histogram is given. It even outperforms WDS on ϕ_3 . This is because by computing the optimal solution based on the histogram, we set higher probabilities for more frequent values, which happens to suit the scenario of ϕ_3 . Note that this does not contradict the optimality of the solution based on full frequency information, which is assuming the *worst* filter condition ϕ .

The comparison between UDS and WDS is interesting on the Zipfian dataset. We see from Figure 3e and 3f that on ϕ_1 , which passes each tuple with equal probability, the two algorithms have very similar performance, while WDS significantly outperforms UDS on ϕ_3 across all values of t. These results are consistent with the design and analysis of WDS, which is aimed at minimizing the MSE over worst-case queries on skewed data. Meanwhile, it is also reassuring to see that, on easier, non-worst-case data and queries, WDS is no worse than its competitors.

7.3 Results on Benchmark Data

We used the TPC-DS benchmark data generator to generate a dataset of size 2 GB, and tested the following 2 queries from the benchmark.

SP queries. Query 28 finds the number of distinct list prices. It consists of 6 distinct count subqueries on a single table over different sales buckets. Thus, each subquery is an SP query with a different selection condition. The results on these 6 SP queries are shown in Figure 4a and 4b. We see that WDS achieves a smaller MSE than UDS in all cases. HISTO performs similar to WDS, and is always better than UDS as well.

SPJ queries. Query 54 finds the number of customers that purchased a given item during a given period. With the union between catalog and web sales carried out first, the query is an SPJ query involving a star join between one fact table and two dimension tables. We tested 4 queries with selection conditions on different

attributes of the dimension tables. For SPJ queries, we also included the random walk algorithm (RW) in the experimental comparison.

The results are shown in Figure 4c and 4d. We see that both WDS and RW have smaller MSE than UDS. The performances of WDS and RW are very similar. This is because the join is a foreign-key star join, *i.e.*, each tuple in the fact table joins with exactly one tuple in each of the dimension tables, and the projection attribute is in the fact table. Thus, the random walks will start from the fact table and walk towards the dimension tables. This has the same effect as sampling from all the join results, since there is a one-to-one correspondence between the tuples in the fact table and the full join results. However, we can run RW much more efficiently to collect the sample as there is no need to compute and materialize the join results. Results on the sample construction time will be given in Section 7.5.

7.4 Results on Real Data

We used two real datasets and tested the following SPJ queries:

IMDb data. We tested a query that finds all actors and actresses that ever acted in a movie whose rating is above a given threshold t on IMDb dataset⁶. This involves data stored in two tables: principals(tconst, nconst, category), the acting list, and ratings(tconst, rating), rating information of movies. We cleaned up the data by removing movies without ratings. This results in 7 million tuples in the principals table and 1 million tuples in the ratings table. Then, the query can be written as the following SPJ query with a selection condition featuring a complicated expression on multiple attributes:

```
SELECT distinct nconst FROM principals, ratings
WHERE principals.tconst = ratings.tconst
AND (principals.category = 'actor'
   OR principals.category = 'actress')
AND ratings.rating > t
```

As this is a large dataset, we used a sample size that is 0.1% of the dataset. The results are shown in Figure 5a and 5b as we vary the parameter *t*. Here, we omit UB as it is way off the chart. We see that WDS performs much better than UDS, while RW and WDS are close, because this join is also a foreign-key join.

Network data. The last dataset is the network dataset [31], which stores information about a who-trusts-whom network on a Bitcoin platform. Vertices represent users and directed edges represent trust relationships. The edges are weighted -10 to 10, denoting the degree of trust from the source user to the destination user. There are around 6k vertices and 36k edges. Our query of interest finds the users involved in at least one *trust triangle*. Specifically, three users X, Y, Z form a trust triangle if there exist edges from X to Y, Y to Z, and Z to X, and the weight of each edge is at least a given threshold t. We store the graph as a relation R(src, dst, trust), so the above query can be written as the following SPJ query involving a triangle join:

```
SELECT distinct R1.src FROM R as R1, R as R2, R as R3
WHERE R1.dst = R2.src AND R1.trust > t
AND R2.dst = R3.src AND R2.trust > t
AND R3.dst = R1.src AND R3.trust > t
```

⁶https://www.imdb.com/interfaces/

Detect	0	Saı	nple cons	truction c	ost	Query cost				
Dataset	Query	UDS	WDS	HISTO	RW	EXACT	UDS	WDS	RW	
Uniform	ϕ_1	125 ms	163 ms	156 ms	-	100 ms	< 1 ms	< 1 ms	-	
	ϕ_2					95 ms	< 1 ms	< 1 ms		
Zipfian	$\phi_1 \\ \phi_3$	300 ms	406 ms	356 ms		115 ms	< 1 ms	< 1 ms	-	
		300 1113	400 1113	JJ0 1115		182 ms	< 1 ms	< 1 ms	-	
TPC-DS	Q28	561 ms	782 ms	753 ms	-	385 ms	1 ms	1 ms	-	
	Q54	5.51 s	5.75 s	-	281 ms	6.41 s	< 1 ms	< 1 ms	< 1 ms	
IMDb		16.3 s	17.3 s	-	2.43 s	21.3 s	45 ms	36 ms	21 ms	
Network		4.33 s	4.35 s	-	61 ms	4.46 s	< 1 ms	< 1 ms	< 1 ms	

Table 4: Comparison of Time Costs

The results are shown in Figure 5c and 5d where we vary the parameter t. We see that WDS and UDS have similar performance. Recall that these two algorithms are run on the join results, which consist of all triangles in the network. This is a skewed dataset, as some vertices are involved in many triangles while the rest have few triangles. Each triangle has the same probability to pass the selection condition, so this SP query (on the join results) is an easy case, similar to what we have observed in Figure 3f. The triangle join presents a challenge for RW, since many random walks may not be able to close as a triangle. This results in a smaller effective sample size, which in turn leads to larger MSE.

7.5 Running Times

We have also measured the wall-clock running times of the algorithms, including both the time for building the sample and the time for producing an estimate for a given query, which are summarized in Table 4. First, we see that the query costs of all estimation algorithms are order-of-magnitude smaller than of computing the query size exactly. This justifies their use in a query optimizer.

The sample construction cost is larger for WDS compared to UDS, as we need to find the optimal sampling strategy first. But the costs are small anyway for SP queries. For SPJ queries, the sample construction cost is dominated by that of computing the join, which dwarfs the additional cost of finding the optimal sampling strategy. On the other hand, RW is able to build the sample without computing the join, resulting in significant savings.

7.6 Summary of Experimental Evaluation

From our experimental evaluation, we can draw the following conclusions: For SP queries, WDS performs at least as well as UDS, and outperforms it on hard queries, especially on skewed data. Thus, as long as one is willing to pay the small extra cost to find the optimal sampling strategy, WDS is always the recommended method. HISTO performs similar to WDS and is desirable when the frequency vector is not known. For SPJ queries, if one can tolerate the cost to pre-compute the join, then running WDS on the join results is the best choice. Otherwise, RW can be used, especially for foreign-key joins, where its performance is as good as that of running WDS on the full join results. On cyclic joins, however, RW may lose some accuracy compared with WDS.

8 CONCLUSIONS

In this paper, we have introduced weighted distinct sampling to tackle the problem of size estimation for SP queries, and extended it to handling SPJ queries. We have presented algorithms that can collect the sample in time linear to the size of the database, and it is also not difficult to adapt them to run in a constant number of MapReduce rounds, enabling it to run over large distributed datasets.

There are two interesting directions for further investigation: The first is dynamic maintenance. There is a lot of work on dynamic maintenance of various synopses [20–24, 33, 35, 40], and how to maintain a near-optimal weighted distinct sample under insertions and/or deletions of tuples is certainly a problem of interest. While we believe that this is doable for SP queries, the problem is more challenging for SPJ queries, as a single update could affect many join results.

The other direction is to consider some special classes of predicates, such as range conditions. For such a special class of predicates, we should be able to achieve better accuracy than weighted distinct sampling, which aims to support arbitrary selection conditions.

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