Efficient Computation of Quantiles over Joins

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ABSTRACT

We present efficient algorithms for Quantile Join Queries, abbreviated as %JQ. A %JQ asks for the answer at a specified relative position (e.g., 50% for the median) under some ordering over the answers to a Join Query (JQ). Our goal is to avoid materializing the set of all join answers, and to achieve quasilinear time in the size of the database, regardless of the total number of answers. A recent dichotomy result rules out the existence of such an algorithm for a general family of queries and orders. Specifically, for acyclic JQs without self-joins, the problem becomes intractable for ordering by sum whenever we join more than two relations (and these joins are not trivial intersections). Moreover, even for basic ranking functions beyond sum, such as min or max over different attributes, so far it is not known whether there is any nontrivial tractable %JQ.

In this work, we develop a new approach to solving %JQ and show how this approach allows not just to recover known results, but also generalize them and resolve open cases. Our solution uses two subroutines: The first one needs to select what we call a "pivot answer". The second subroutine partitions the space of query answers according to this pivot, and continues searching in one partition that is represented as new %JQ over a new database. For pivot selection, we develop an algorithm that works for a large class of ranking functions that are appropriately monotone. The second subroutine requires a customized construction for the specific ranking function at hand.

We show the benefit and generality of our approach by using it to establish several new complexity results. First, we prove the tractability of min and max for all acyclic JQs, thereby resolving the above question. Second, we extend the previous %JQ dichotomy for sum to all partial sums (over all subsets of the attributes). Third, we handle the intractable cases of sum by devising a *deterministic* approximation scheme that applies to *every acyclic JQ*.

CCS CONCEPTS

• Theory of computation \rightarrow Database query processing and optimization (theory); *Database theory*.

KEYWORDS

join queries, quantiles, median, ranking function, answer order, pivot, approximation, inequality predicates

ACM Reference Format:

Nikolaos Tziavelis, Nofar Carmeli, Wolfgang Gatterbauer, Benny Kimelfeld, and Mirek Riedewald. 2023. Efficient Computation of Quantiles over Joins. In Proceedings of the 42nd ACM SIGMOD-SIGACT-SIGAI Symposium on Principles of Database Systems (PODS '23), June 18–23, 2023, Seattle, WA, USA. ACM, New York, NY, USA, 14 pages. https://doi.org/10.1145/3584372.3588670

1 INTRODUCTION

Quantile queries ask for the element at a given relative position $\phi \in [0, 1]$ in a given list *L* of items [21]. For example, the lower quartile, median, and upper quartile are the elements at positions $\phi = 0.25$, $\phi = 0.5$, and $\phi = 0.75$, respectively. We investigate quantile queries where *L* is the result of a Join Query (JQ) *Q* over a database, with a ranking function that determines the order between the answers. Importantly, the list *L* can be much larger than the input database *D* (specifically, *L* can be $\Omega(|D|^k)$ for some degree *k* determined by *Q*), and so, *Q* and *D* form a compact representation for *L*. Our main research question is *when we can find the quantile in quasilinear time*. In other words, the time suffices for reading *D*, but we are generally prevented from materializing *Q*(*D*).

For illustration, consider a social network where users organize events, share event announcements, and declare their plans to attend events. It has the three relations Admin(user, event), Share(user, event, #likes), and Attend(user, event, #likes). We wish to extract statistics about triples of users involved in events, beginning by joining the three relations using the JQ

Admin (u_1, e) , Share (u_2, e, l_2) , Attend (u_3, e, l_3) .

Now, suppose that all we do is apply a quantile query to the result of Q, say the 0.1-quantile ordered by $l_2 + l_3$ (the sum of likes of the share and the participation). The direct way of finding the quantile is to materialize the join, sort the resulting tuples, and take the element at position $\phi = 0.1$. Yet, this result might be considerably larger than the database, and prohibitively expensive to compute, even though in the end we care only about one value. Can we do

PODS '23, June 18-23, 2023, Seattle, WA, USA

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better? This is the research question that we study in this paper. In general, the answer depends on the JQ and order, and in this specific example we can, actually, do considerably better.

To be more precise, we study the fine-grained data complexity of query evaluation, where the query seeks a quantile over a JQ. We refer to such a query as a *Quantile Join Query* and abbreviate it as %JQ. So, the JQ Q is fixed, and so is the ranking function (e.g., weighted sum over a fixed subset of the variables). The input consists of D and ϕ . In terms of the execution cost, we allow for poly-logarithmic factors, therefore our goal is to devise evaluation algorithms that run in time quasilinear in D, that is, O(|D| polylog(|D|)).

To the best of our knowledge, little is known about the finegrained complexity of %JQ. We have previously studied this problem [7] for Conjunctive Queries (which are more general than Join Queries since they also allow for projection) under the name "*selection problem*".¹ Two types of orders were covered: sum of all attributes and lexicographic orders. On the face of it, the conclusion from our previous results is that we are extremely limited in what we can do: *The problem is intractable for every JQ with more than two atoms* (each having a set of variables that is not contained in that of another atom, and assuming no self-joins), under conventional conjectures in fine-grained complexity. Nevertheless, we argue that our previous results tell only part of the story and miss quite general opportunities for tractability:

- What if the sum involves just a *subset* of the variables, like in the above social-network example? Then the lower bounds for full sum do not apply. As it turns out, in this case we often can achieve tractability for JQs of more than 2 atoms.
- What if we allow for some small error and not insist on the precise φ-quantile? As we argue later, this relaxation makes the picture dramatically more positive.

In addition, there are ranking functions that have not been considered at all, notably minimum and maximum over attributes, such as MIN(rate1, rate2) and MAX(width, height, depth). We do not see any conclusion from past results on these, so the state of affairs (prior to this paper) is that their complexity is an open problem.

In this work, we devise a new framework for evaluating %JQ queries. We view the problem as a search problem in the space of query answers, and the framework adopts a divide-and-conquer "pivoting" approach. For a JQ Q, we reduce the problem to two subroutines, given D and ϕ :

- **PIVOT**: Find a *pivot* answer *p* such that the set of answers that precede *p* and the set of answers that follow *p* both contain at least a constant fraction of the answers.
- **TRIM**: Partition the answers into three splits: less than, equal to, and greater than *p*. Determine which one contains the sought answer and, if it is not *p*, produce a new %JQ within the relevant split using new Q', D' and ϕ' . We view this operation as *trimming* the split condition by updating the database so that the remaining answers satisfy the condition.

We begin by showing that we can select a pivot in linear time for every acyclic join query and every ranking function that satisfies a monotonicity assumption (also used in the problem of ranked enumeration [15, 23]), which all functions considered here satisfy. Note that the assumption of acyclicity is required since, otherwise, it is impossible to even determine whether the join query has any answer in quasilinear time, under conventional conjectures in finegrained complexity [6]. Hence, the challenge really lies in trimming.

Contributions. Using our approach, we establish efficient algorithms for several classes of queries and ranking functions, where we show how to solve the trimming problem.

- (1) We establish tractability for all acyclic JQs under the ranking functions MIN and MAX.
- (2) We recover (up to logarithmic factors) all past tractable cases [7] for lexicographic orders and SUM.
- (3) For self-join-free JQs and SUM, we complete the picture by extending the previous dichotomy [7] (restricted to JQs) to all *partial sums*.

We then turn our attention to approximate answers. Precisely, we find an answer at a position within $(1 \pm \epsilon)\phi$ for an allowed error ϵ . (This is a standard notion of approximation for quantiles [9, 17].) To obtain an efficient *randomized* approximation, it suffices to be able to construct in quasilinear time a direct-access structure for the underlying JQ, regardless of the answer ordering; if so, then one can use a standard median-of-samples approach (with Hoeffding's inequality to guarantee the error bounds). Such algorithms for direct-access structures have been established in the past for arbitrary acyclic JQs [6, 8]. Instead, we take on the challenge of *deterministic* approximation. Our final contribution is that:

(4) We show that with an adjustment of our pivoting framework, we can establish a deterministic approximation scheme in time quadratic in 1/ε and quasilinear in database size.

In contrast to the randomized case, we found the task of deterministic approximation challenging, and our algorithm is indeed quite involved. Again, the main challenge is in the trimming phase.

The remainder of the paper is organized as follows: We give preliminary definitions in Section 2. We describe the general pivoting framework in Section 3. In Section 4, we describe the pivot-selection algorithm. The main results are in Sections 5 and 6 where we devise exact and approximate trimmings, respectively, and establish the corresponding tractability results. We conclude in Section 7.

2 PRELIMINARIES

2.1 Basic Notions

Sets. We use [r] to denote the set of integers $\{1, \ldots, r\}$. A multiset *L* is described by a 2-tuple (Z, β) , where *Z* is the set of its distinct elements and $\beta : Z \to \mathbb{N}$ is a multiplicity function. The set of all possible multisets with elements *Z* is denoted by \mathbb{N}^Z .

Relational databases. A schema S is a set of relational symbols $\{R_1, \ldots, R_m\}$. A database D contains a finite relation $R^D \subseteq \text{dom}^{a_R}$ for each $R \in S$, where dom is a set of constants called the *domain*, and a_R is the arity of symbol R. If D is clear, we simply use R instead of R^D . The size of D is the total number of tuples, denoted by n.

Join Queries. A *Join Query* (JQ) *Q* over schema *S* is an expression of the form $R_1(\mathbf{X}_1), \ldots, R_\ell(\mathbf{X}_\ell)$, where $\{R_1, \ldots, R_\ell\} \subseteq S$ and the variables of *Q* are $\operatorname{var}(Q) = \bigcup_{i \in [\ell]} \mathbf{X}_i$, sometimes interpreted as a tuple instead of a set. Each $R_i(\mathbf{X}_i), i \in [\ell]$ is called an *atom* of

¹To be precise, in the selection problem, the position of this answer is given as an absolute index *i* rather than a relative position ϕ . (This problem is sometimes referred to as *unranking* [18].) This difference is nonessential as far as this work concerns: all our previous lower and upper bounds for selection on JQs apply to %JQs.

Q. A repeated occurrence of a relational symbol is a *self-join* and a JQ without self-joins is *self-join-free*. A *query answer* is a homomorphism from *Q* to the database *D*, i.e. a mapping from var(Q) to dom constants, such that every atom $R_i(\mathbf{X}_i)$, $i \in [\ell]$ maps to a tuple of R_i^D . The set of query answers to *Q* over *D* is denoted by Q(D) and we often represent a query answer $q \in Q(D)$ as a tuple of values assigned to var(Q). For an atom $R_i(\mathbf{X}_i)$ of a JQ and database *D*, we say that tuple $t \in R_i^D$ assigns value *a* to variable *x*, and write it as t[x] = a, if for every index *j* such that $\mathbf{X}_i[j] = x$ we have t[j] = a. For a predicate $P(\mathbf{X}_P)$ over variables $\mathbf{X}_P \subseteq var(Q)$, we denote by $(Q \land P)(D)$ the subset of query answers Q(D) that satisfy $P(\mathbf{X}_P)$.

Hypergraphs. A hypergraph $\mathcal{H} = (V, E)$ is a set V of vertices and a set $E \subseteq 2^V$ of hyperedges. A path in \mathcal{H} is a sequence of vertices such that every two consecutive vertices appear together in a hyperedge. A chordless path is a path in which no two non-consecutive ones appear in the same hyperedge (in particular, no vertex appears twice). The number of maximal hyperedges is $\mathrm{mh}(\mathcal{H}) = |\{e \in E \mid \nexists e' \in E : e \subset e'\}|$. A set of vertices $U \subseteq V$ is independent if no pair appears in a hyperedge, i.e., $|U \cap e| \leq 1, \forall e \in E$.

Join trees. A *join tree* of a hypergraph $\mathcal{H} = (V, E)$ is a tree T where its nodes² are the hyperedges of \mathcal{H} and the *running intersection* property holds, namely: for all $u \in V$ the set $\{e \in E \mid u \in e\}$ forms a (connected) subtree in T. We associate a hypergraph $\mathcal{H}(Q) = (V, E)$ to a JQ Q where the vertices are the variables of Q, and every atom of Q corresponds to a hyperedge with the same set of variables. With a slight abuse of notation, we identify atoms of Q with hyperedges of $\mathcal{H}(Q)$. A JQ Q is *acyclic* if there exists a join tree for $\mathcal{H}(Q)$, otherwise it is *cyclic*. If we root the join tree, the subtree rooted at a node U defines a subquery, i.e., a JQ that contains only the atoms of descendants of U. A partial query answer (for the subtree) rooted at U is an answer to the subquery. If we materialize a relation R_U for node U, a partial query answer (for the subtree) rooted at $t \in R_U$ must additionally agree with t.

Complexity. We measure complexity in the database size n, while query size is considered constant. The model of computation is the standard RAM model with uniform cost measure. In particular, it allows for linear-time construction of lookup tables, which can be accessed in constant time. Following our prior work [7], we only consider comparison-based sorting, which takes quasilinear time.

2.2 Orders over Query Answers

To define %JQs, we assume an ordering of the query answers by a given ranking function. The ranking function is described by a 2-tuple (w, \leq) where a weight function $w : Q(D) \rightarrow \operatorname{dom}_w$ maps the answers to a weight domain dom_w equipped with a total order \leq . We denote the strict version of the total order by \prec . Assuming consistent tie-breaking, the total order extends to query answers, i.e., for $q_1, q_2 \in Q(D), q_1 \leq q_2$ iff $w(q_1) \prec w(q_2)$ or $w(q_1) = w(q_2)$ and q_1 is (arbitrarily but consistently) chosen to break the tie.

Weight aggregation model. We focus on the case of *aggregate* ranking functions where the query answer weights are computed by aggregating weights are assigned to the input database. In particular, an input-weight function $w_x : \operatorname{dom} \to \operatorname{dom}_w$ associates each value of variable x with a weight in dom_w . An aggregate function $\operatorname{agg}_w : \mathbb{N}^{\operatorname{dom}_w} \to \operatorname{dom}_w$ takes a multiset of weights and produces a single

weight. Aggregate ranking functions are typically not sensitive to the order in which the input weights are given [11, 12], captured by the fact that their input is a multiset. Query answers map to dom_w by aggregating the weights of values assigned to a subset of the input variables $U_w \subseteq var(Q)$ with an aggregate function agg_w . Thus, the weight of a query answer $q \in Q(D)$ is w(q) = $agg_w(\{w_x(q[x]) \mid x \in U_w\})$. When we do not have a specific assignment from variables to values, we use $w(U_w)$ to refer to the expression $agg_w(\{w_x(x) \mid x \in U_w\})$. For example, if var(Q) = $\{x_1, x_2, x_3\}, U_w = \{x_1, x_3\}, w_x(x)$ is the identity function for all variables x, and agg_w is summation, then $w(U_w) = x_1 + x_3$.

Concrete ranking functions. In this paper, we discuss three types of ranking functions:

- SUM: dom_w is ℝ and agg_w is summation. We use the term full SUM when U_w = var(Q) and partial SUM otherwise.
- (2) MIN/ MAX: dom_w is \mathbb{R} and agg_w is min or max.
- (3) LEX: Lexicographic orders fit into our framework by letting the domain dom_w consist of tuples in N^{|U_w|}. Every variable x ∈ U_w is mapped to dom_w as a tuple (0,..., w'_x(x),...,0) where w'_x(x) occupies the position of x in the lexicographic order and w'_x is a function w'_x : dom → N that orders the domain of x by mapping it to natural numbers. The aggregate function agg_w is then element-wise addition, while the order ≤ compares these tuples lexicographically.

Problem definition. Let Q be a JQ and (w, \leq) a ranking function. Given a database D, a query answer $q \in Q(D)$ is a ϕ -quantile [21] of Q(D) for some $\phi \in [0, 1]$ if there exists a valid ordering of Q(D) where there are $\lceil \phi | Q(D) \rceil$ answers less-than or equal-to q and $\lfloor (1 - \phi) | Q(D) \rfloor$ answers greater than q. A %JQ asks for a ϕ -quantile given D and ϕ . Similarly, an ϵ -approximate %JQ asks for a $(\phi \pm \epsilon)$ -quantile for a given D, ϕ , and $\epsilon \in (0, 1)$.

Monotonicity. Let \forall be multiset union. An (aggregate) ranking function is *subset-monotone* [23] if $\operatorname{agg}_{w}(L_1) \leq \operatorname{agg}_{w}(L_2)$ implies that $\operatorname{agg}_{w}(L \uplus L_1) \leq \operatorname{agg}_{w}(L \uplus L_2)$ for all multisets L, L_1, L_2 . All ranking functions we consider in this work have this property. We note that subset-monotonicity has been used as an assumption in ranked enumeration [15, 23] and is a stronger requirement than the more well-known monotonicity notion of Fagin et al. [10].

Tuple weights. Our ranking function definition uses attributeweights but some of our algorithms are easier to describe when dealing with tuple weights. We can convert the former to the latter in linear time. First, we eliminate self-joins by materializing a fresh relation for every repeated symbol in the query Q. Second, to avoid giving the weight of a variable to tuples of multiple relations, we define a mapping μ that assigns each variable $x \in U_w$ to a relation R such that x occurs in the R-atom of Q. The weight of a tuple $t \in R$ is then the multiset of weights for variables assigned to R: $w_R(t) = \{w_x(t[x]) \mid x \in U_w, \mu(x) = R\}$.³ The total order \leq can be extended to sets of tuples (t_1, \ldots, t_r) (and thus query answers) by aggregating all individual weights contained in the tuple weights.

²To avoid confusion, we use the terms hypergraph vertices and tree nodes.

³The reason that we maintain the attribute weights as a set instead of aggregating them is that the aggregate ranking function can be holistic [11], in which case we lose the ability to further aggregate. If, on the other hand, the ranking function is distributive [11] like SUM, then we can aggregate to obtain a single weight for a tuple.

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2.3 Known Bounds

Certain upper and (conditional) lower bounds for %JQ follow from our previous work [7] on the *selection problem* which asks for the query answer at index k. The two problems are equivalent for acyclic JQs, since an index can be translated into a fraction ϕ , and vice-versa, by knowing |Q(D)|, which can be computed in linear time as we explain in Section 2.4.

The lower bounds are based on two hypotheses:

- HYPERCLIQUE [1, 16]: Let a (k+1, k)-hyperclique be a set of k+1 vertices such that every subset of k elements is a hyperedge. For every k ≥ 2, there is no O(m polylog m) algorithm for deciding the existence of a (k+1, k)-hyperclique in a k-uniform hypergraph with m hyperedges.
- (2) 3SUM [4, 19]: For any $\epsilon > 0$, we cannot decide in time $O(m^{2-\epsilon})$ whether there exist $a \in A, b \in B, c \in C$ from three integer sets A, B, C, each of size $\Omega(m)$, such that a + b + c = 0.

HYPERCLIQUE implies that we cannot decide in O(n polylog n) if a cyclic, self-join-free JQ has any answer [6]. For LEX, an acyclic %JQ can be answered in O(n) [7]. For full SUM, an acyclic %JQ can be answered in $O(n \log n)$ if its maximal hyperedges are at most 2, and the converse is true if it is also self-join-free, assuming 3SUM [7].

2.4 Message Passing

Message passing is a common algorithmic pattern that many algorithms for acyclic JQs follow. For example, it allows us to count the number of answers to an acyclic JQ in linear time [3, 20]. Some of the algorithms that we develop also follow this pattern, that we abstractly describe below.

Preprocessing. Choose an arbitrary root for a join tree *T* of the JQ, and materialize a distinct relation for every *T*-node. For every parent node V_p and child node V_c , group the V_c -relation by the $V_p \cap V_c$ variables. We will refer to these groups of tuples as *join groups*; a join group shares the same values for variables that appear in the parent node. The algorithm visits the relations in a bottom-up order of *T*, sending children-to-parent messages. The goal is to compute a value val(*t*) for each tuple *t* of these relations, initialized according to the specific algorithm. Sometimes, it is convenient to add an artificial root node $V_0 = \emptyset$ to the join tree, which refers to a zero-arity relation with a single tuple $t_0 = ()$. Tuple t_0 joins with all tuples of the previous root and its purpose is only to gather the final result at the end of the bottom-up pass.

Messages. As we traverse the relations in bottom-up order, every tuple t emits its val(t). These messages are aggregated as follows:

- Messages emitted by tuples t' in a join group are aggregated with an operator ⊕. The result is sent to all parent-relation tuples that agree with the join values of the group.
- (2) A tuple *t* computes val(*t*) by aggregating the messages received from all children in the join tree, together with the initial value of val(*t*), with an operator ⊗.

Example 2.1 (Count). To count the JQ answers, we initialize cnt(t) = 1 for all tuples t, set \oplus to product (\prod) , and \otimes to sum (\sum) . Figure 1 illustrates how messages are aggregated so that cnt(t) is the number of partial answers for the subtree rooted at t. To get the final count, we sum the counts in the root relation (9+4=13 in the example), e.g., by introducing the artificial root-node tuple t_0 .



(a) Example database and final (b) Messages received and aggrecounts of subtree answers. gated by an *R*-tuple.

Figure 1: Message passing for counting the answers to the JQ $R(x_1, x_2), S(x_1, x_3), T(x_2, x_4), U(x_4, x_5).$

3 DIVIDE-AND-CONQUER FRAMEWORK

We describe a general divide-and-conquer framework for acyclic %JQs that applies to different ranking functions. It follows roughly the same structure as linear-time selection [5] in a given array of elements. This classic algorithm searches for the element at a desired index k in the array by "pivoting". In every iteration, it selects a pivot element and creates three array partitions: elements that are lower, equal, and higher than the pivot. Depending on the partition sizes and the value of k, it chooses one partition and continues with that, thereby reducing the number of candidate elements. We adapt the high-level steps of this algorithm to our setting. The crucial challenge is that we do not have access to the materialized array of query answers (which can be very large), but only to the input database and JQ that produce them. In the following, we discuss the general structure of the algorithm and the subroutines that are required for it to work. In later sections, we then concretely specify these subroutines.

Pivot selection. We define what constitutes a "good" pivot. Intuitively, it is an element whose position is roughly in the middle of the ordering. With such a pivot, the partitioning step is guaranteed to eliminate a significant number of elements, resulting in quick convergence. Ideally, we would want to have the true median as our pivot because it is guaranteed to eliminate the largest fraction $(\frac{1}{2})$ of elements. However, to achieve convergence in a logarithmic number of iterations, it is sufficient to choose any pivot that eliminates any constant fraction c > 0 of elements.

Definition 3.1 (*c*-pivot). For a constant $c \in (0, 1)$ and a set *Z* equipped with a total order \leq , a *c*-pivot *p* for *Z* is an element of *Z* such that $|\{z \in Z \mid p \leq z\}| \geq c|Z|$ and $|\{z \in Z \mid p \geq z\}| \geq c|Z|$.

Our goal is to find such a *c*-pivot for the set of query answers Q(D) ordered by the given ranking function.

Partitioning. Assuming an appropriate query answer p as our pivot, we use it to partition the query answers. This means that we want to separate the answers into those whose weight is less than, equal to, and greater than the weight of the pivot. Since we do not have access to the query answers, this partitioning step must be performed on the input database and JQ. The less-than and greater-than partitions can be described by the original JQ, together with inequality predicates: (1) $w(U_w) < w(p)$ and (2) $w(U_w) > w(p)$

respectively. The equal-to partition can be assumed to contain all answers that do not fall into either of the other two.

Trimming inequalities. If we materialize as database relations the inequalities that arise from the partitioning step, their size can be very large. For example, the inequality $x_1 + x_2 + x_3 < 0$ for three variables x_1, x_2, x_3 has a listing representation of size $O(n^3)$. However, in certain cases it is possible to represent them more efficiently, e.g., in space O(n polylog n), by modifying the original JQ and database. We call this process "*trimming*."

Definition 3.2 (Predicate Trimming). Given a JQ Q and a predicate P(U) with variables $U \subseteq var(Q)$, a trimming of P(U) from Q receives a database D and returns a JQ Q' of size O(|Q|) and with $var(Q) \subseteq var(Q')$, and a database D' for which there exists an O(1)-computable bijection from Q'(D') to $(Q \land P)(D)$. Trimming time is the time required to construct Q' and D'.

Efficient trimmings of predicates are for instance known for additive inequalities when the sum variables are found in adjacent JQ atoms [22] and for not-all-equals predicates [14], which are a generalization of non-equality (\neq). Ultimately, our ability to partition and the success of our approach relies on the existence of efficient trimmings of inequalities that involve the aggregate function.

Choosing a partition. After we obtain three new JQs and corresponding databases by trimming, we count their query answers to determine where the desired index (calculated from the given percentage) falls into. To ensure that this can be done in linear time, we want all JQs to be acyclic, and so we restrict ourselves to trimmings that do not alter the acyclicity of the JQs. To keep track of the candidate query answers, we maintain two weights low and high as bounds, which define a contiguous region in the sorted array of query answers. Every iteration then applies trimming for two additional inequalities $w(U_w) > 1$ ow and $w(U_w) <$ high in order to restrict the search to the current candidate set.

Termination. The algorithm terminates when the desired index falls into the equal partition since any of its answers, including our pivot, is a ϕ -quantile.⁴ It also terminates when the number of candidate answers is sufficiently small, by calling the Yannakakis algorithm [24] to materialize them and then applying linear-time selection [5]. With *c*-pivots, we eliminate at least c|Q(D)| answers in every iteration; hence, the candidate query answers will be O(n) after a logarithmic number of iterations. Notice that our trimming definition allows the new database to be larger than the starting one, so the database size may increase across iterations. However, the number of JQ answers decreases, ensuring termination.

Summary. Our algorithm repeats the above steps (selecting pivot, partitioning, trimming) iteratively. It requires the implementation of two subroutines: (1) selecting a *c*-pivot among the JQ answers, which we call "PIVOT", and (2) trimming of inequalities, which we call "TRIM". The pseudocode is in Appendix B.

LEMMA 3.3 (EXACT QUANTILES). Let Q be a class of acyclic JQsand (w, \leq) a ranking function. If for all $Q' \in Q$

 there exists a constant c such that for any database D, a cpivot for Q'(D) can be computed in time O(g_p(n)) for some function q_p, and (2) for all $\lambda \in \text{dom}_w$, there exist trimmings of $w(U_w) \prec \lambda$ and $w(U_w) \succ \lambda$ from Q' that return $Q'' \in Q$ in time $O(g_t(n))$ for some function g_t ,

then a %JQ can be answered for all $Q \in Q$ in time $O(\max\{g_p(n), g_t(n)\} \log n)$.

Notice that trimming can result in a *different* query than the one we start with. This is why pivot-selection and trimming need to applicable not just to the input query *Q*, but to all queries that we may obtain from trimming (referred to as a class in Lemma 3.3).

Example 3.4. Suppose that Q is $R_1(x_1, x_2), R_2(x_2, x_3)$ over a database D and we want to compute the median by SUM with attribute weights equal to their values. First, we call PIVOT to obtain a pivot answer p, which we use to create two partitions: one with $x_1 + x_2 + x_3 < w(p)$, and one with $x_1 + x_2 + x_3 > w(p)$. Second, we call TRIM on these inequalities. By a known construction [22], these inequalities can be trimmed in $O(n \log n)$. This construction adds a new column and variable v to both relations. We now have two JQs Q_{1t} and Q_{gt} , over databases D_{1t} , D_{gt} . For example, Q_{1t} is $R_{1_{1t}}(x_1, x_2, v)$, $R_{2_{1t}}(v, x_2, x_3)$. Suppose that |Q(D)| = 1001, hence the desired index is k = 500 (with zeroindexing). If $|Q_{1t}(D_{1t})| = 400$ and $|Q_{gt}(D_{gt})| = 600$, we can infer that the middle partition contains 1 one answer with weight w(p). Thus, we have to continue searching in the index range from 401 to 1000 with k' = 500 - 400 - 1 = 99. To create less-than and greaterthan partitions in the next iteration, we will start with the original *Q* and *D* and apply inequalities $w(p) \le x_1 + x_2 + x_3 < w(p')$ and $w(p') < x_1 + x_2 + x_3 < \infty$ with some new pivot p'.

In Section 4, we will show that an efficient algorithm for PIVOT exists for *any subset-monotone ranking function*. For TRIM, the situation is more tricky and no generic solution is known. For each ranking function, we design a trimming algorithm tailored to it. This is precisely where we encounter the known conditional hardness of SUM [7]. For example, a quasilinear trimming for $Q(x_1, x_2, x_3) := R_1(x_1), R_2(x_2), R_3(x_3)$ and $x_1 + x_2 + x_3 < 0$ would violate our 3SUM hypothesis (see Section 2.3) because it would allow us to count the number of answers in the less-than and greater-than partitions. In Section 5, we will show that efficient trimmings exist for MIN/MAX and LEX, as well as (partial) SUM in certain cases.

3.1 Adaptation for Approximate Quantiles

Since %JQ can be intractable (under our efficiency yardstick) for some ranking functions such as SUM [7], we aim for ϵ -approximate quantiles. We can obtain a *randomized* approximation by the standard technique of sampling answers uniformly and taking as estimate the ϕ -quantile of the sample (e.g., as done by Doleschal et al. [9] for quantile queries in a different model). Concentration theorems such as Hoeffding Inequality imply that it suffices to collect $O(1/\epsilon^2)$ samples and repeat the process $O(\log(1/\delta))$ times (and select the median of the estimates) to get an ϵ -approximation with probability $1 - \delta$. So, it suffices to be able to efficiently sample uniformly a random answer of an acyclic JQ; we can do so using linear-time algorithms for constructing a logarithmic-time random-access structure for the answers [6, 8].

We will show that with our pivoting approach, we can obtain a *deterministic* approximation, which we found to be much more

⁴If we want to enforce the same tie-breaking scheme across different calls to our algorithm, we could continue searching within the equal partition with a LEX order, but this requires also trimming for equality-type predicates.

challenging than the randomized one. Pivot selection remains the same as in the exact algorithm, while for trimming (which as we explained is the missing piece for SUM), we introduce an approximate solution based on the notion of a *lossy trimming*. Intuitively, a lossy trimming does not retain all the JQ answers that satisfy a given predicate. Such a trimming results in some valid query answers being lost in each iteration and causes inaccuracy in the final index of the returned query answer. However, if the number of lost query answers is bounded, then we can also bound the error on the index.

Definition 3.5 (Lossy Predicate Trimming). Given a JQ Q, a predicate P(U) with variables $U \subseteq var(Q)$, and a constant $\epsilon \in [0, 1)$, an ϵ -lossy trimming of P(U) from Q receives a database D and returns a JQ Q' of size O(|Q|) and with $var(Q) \subseteq var(Q')$, and a database D' for which there exists an O(1)-computable injection from Q'(D') to $(Q \land P)(D)$, and also $|Q'(D')| \ge (1 - \epsilon)|(Q \land P)(D)|$. Trimming time is the time required to construct Q' and D'.

The injection in the definition above implies that some query answers that satisfy the given predicate do not correspond to any answers in the new instance we construct, but we also ask their ratio to be bounded by ϵ . For $\epsilon = 0$, we obtain an exact predicate trimming (Definition 3.2) as a special case.⁵

LEMMA 3.6 (APPROXIMATE QUANTILES). Let Q be a class of acyclic JQs and (w, \leq) a ranking function. If for all $Q' \in Q$

- there exists a constant c such that for any database D, a cpivot for Q'(D) can be computed in time O(g_p(n)) for some function g_p, and
- (2) for all λ ∈ dom_w and ε' > 0, there exist ε'-lossy trimmings of w(U_w) < λ and w(U_w) > λ from Q' that return Q'' ∈ Q in time O(g_t(n, ε')) for some function g_t,

then an ϵ -approximate %JQ can be answered for all $Q \in Q$ in time $O\left(\max\left\{g_p(n), g_t\left(n, \frac{\epsilon}{2\lceil \ell \log_{1/(1-c)} n\rceil}\right)\right\} \log n\right)$, where ℓ is the number of atoms of Q.

In Section 6 we will give an ϵ -lossy trimming for additive inequalities, which, combined with the pivot algorithm of Section 4, will give us an ϵ -approximate quantile algorithm for SUM.

4 GENERIC PIVOT SELECTION

We describe a PIVOT algorithm for choosing a pivot element among the answers to an acyclic JQ. This is one of the two main subroutines of our quantile algorithm. We show that a *c*-pivot can be computed in linear time for a large class of ranking functions.

LEMMA 4.1 (PIVOT SELECTION). Given an acyclic $\mathcal{J}Q Q$ over a database D of size n and a subset-monotone ranking function, a c-pivot of Q(D) together with $c \in (0, 1)$ can be computed in time O(n).

4.1 Algorithm

The key idea of our algorithm is the "median-of-medians", in similar spirit to classic linear-time selection [5] or selection for the X + Y problem [13]. The main difference is that we apply the median-of-medians idea iteratively using message passing. The detailed pseudocode is given in Appendix C.

Weighted median. An important operation for our algorithm is the weighted median, which selects the median of a set, assuming that every element appears a number of times equal to an assigned weight.⁶ More formally, for a totally-ordered (\leq) set *Z* and a multiplicity function $\beta : Z \to \mathbb{N}^+$, the weighted median WMED $\leq (Z, \beta)$ is the element at position $\lfloor \frac{|B|}{2} \rfloor$ in the multiset $B = (Z, \beta)$ ordered by \leq . The weighted median can be computed in time linear in |Z| [13].

Message passing. Our algorithm employs the message-passing framework as outlined in Section 2.4 to compute pivot(t) for each tuple t bottom-up. The computed pivot(t) is a partial query answer for the subtree rooted at t and serves as a c'-pivot for these partial answers, for some $c' \ge c$. Messages are aggregated as follows: (1) The \oplus operator that combines pivots within a join group is the weighted median. The multiplicity function is given by the count of subtree answers and the order by the ranking function. The counts are also computed using message passing (see Section 2.4). (2) The \otimes operator that combines pivots from different children is simply the union of (partial) assignments to variables.

Example 4.2. Consider the binary-join $R_1(x_1, x_2)$, $R_2(x_2, x_3)$ under full SUM. Assume R_1 is the parent in the join tree with tuple weights $x_1 + x_2$, while R_2 is the child with tuple weights x_3 . First, PIVOT groups the R_2 tuples by x_2 and, for each group, it finds the median x_3 value. The message from R_2 to R_1 is a mapping from x_2 values to (1) the count of R_2 tuples that contain the x_2 value and (2) the median x_3 value over these tuples. Then, every tuple $r_1 \in R_1$ unions its x_1, x_2 values with the incoming x_3 , obtaining a pivot $(r_1) = (x_1, x_2, x_3)$ tuple. To compute the final pivot, PIVOT takes the median of these pivot (r_1) tuples, ranked by $x_1 + x_2 + x_3$, and weighted by the count of R_2 tuples that contain the x_2 value.

Example 4.3. Figure 2 shows how an *R*-tuple computes its pivot under full SUM for the example of Figure 1. Green values in brackets [.] represent the counts, while the orange assignments are the computed pivots for each tuple or join group. From a leaf node like *S* or *U*, messages are simply the relation tuples, each with multiplicity 1. To see how a pivot is computed within a join group, consider the *T*-node group. The pivot of tuple (1, 6) is smaller than the pivot of tuple (1, 7) according to the ranking function because 1+6+8 < 1+7+9. The weighted median selects the latter because it has multiplicity 2 (that is the group size for $x_4 = 6$ in the child *U*).

Pivot accuracy. As we will show, applying our two operations (weighted median, union) results in a loss of accuracy for the pivots, captured by the *c* parameter. The pivots computed for the leaf relations are the true medians (thus $\frac{1}{2}$ -pivots), but the *c* parameter decreases as we go up the join tree. Fortunately, this can be bounded by a function of the query size, making our final result a *c*-pivot with a *c* value that is independent of the data size *n*. The algorithm keeps track of the *c* value for every node and upon termination, the *c* value of the root is returned.

Running time. The time is linear in the database size. The weighted median and count operations are only performed once for every join group and both take linear time. Each tuple is visited only once, and all operations per tuple (e.g., number of child relations, finding the joining group, union) depend only on the query size.

⁵An *imprecise* trimming, which retains *more* JQ answers than it should, would also work for our quantile algorithm.

 $^{^6{\}rm This}$ weight is not the same as the weight assigned by the ranking function, thus we simply call it multiplicity.



Figure 2: Message passing for computing pivots on the example JQ and database of Figure 1 under SUM with weights equal to attribute values.

4.2 Correctness

First, we show that PIVOT returns a valid query answer. The concern is that a variable *x* may be assigned to different values in the pivots that are unioned from different branches of the tree. As we show next, this cannot happen because of the join tree properties.

LEMMA 4.4. Let V be a join-tree node and R_V the corresponding relation. For all $t \in R_V$, the variable assignment pivot(t) computed by PIVOT is a partial query answer for the subtree rooted at V.

Next, we show how the accuracy of the pivot is affected by repeated weighted median and union operations.

LEMMA 4.5. Given r disjoint sets Z_1, \ldots, Z_r equipped with a total order \leq and corresponding c-pivots p_1, \ldots, p_r , then WMED($\{p_1, \ldots, p_r\}, \beta$) with $\beta(p_i) = |Z_i|$, for all $i \in [r]$ is a $\frac{c}{2}$ -pivot for $Z_1 \cup \cdots \cup Z_r$.

LEMMA 4.6. Assume a join-tree node V, its corresponding relation R_V , its children V_1, \ldots, V_r , a subset-monotone ranking function, and c-pivots $p_i, i \in r$ for the partial answers which are rooted at V_i and restricted to those that agree with t. Then, $t \cup p_1 \cup \ldots \cup p_r$ is a c^r -pivot for the partial answers rooted at t.

With the three above lemmas, we can complete the proof of Lemma 4.1 by induction on the join tree.

5 EXACT TRIMMINGS

We now look into trimmings for different types of inequality predicates that arise in the partitioning step of our quantile algorithm (i.e., the TRIM subroutine). Our construction essentially removes these predicates from the query, while ensuring that the modified query can only produce answers that satisfy them.

5.1 MIN/MAX

When the ranking function is MIN or MAX, then we need to trim predicates of the type min{ U_w } < $\lambda, \lambda \in \mathbb{R}$.

Example 5.1. Suppose the ranking function is MAX, the weighted variables are $U_w = \{x_1, x_2, x_3\}$, attribute weights are equal to database values, and our pivot has weight 10. To create the appropriate partitions, we trim predicates $\max\{x_1, x_2, x_3\} < 10$ and $\max\{x_1, x_2, x_3\} > 10$. Enforcing $\max\{x_1, x_2, x_3\} < 10$ is straightforward by removing from the database all tuples with values greater than or equal to 10 for either of the three variables. For

 $\max\{x_1, x_2, x_3\} > 10$, there are three ways to satisfy the predicate: (1) $x_1 > 10$, (2) $x_1 \le 10 \land x_2 > 10$, or (3) $x_1 \le 10 \land x_2 \le 10 \land x_3 > 10$. These three cases are disjoint and cover all possibilities. For each case, we create a fresh copy of the database and then enforce the predicates in linear time by filtering the tuples. Our JQ over one of these three databases produces a partition of the answers that satisfy the original inequality. To return a single database and JQ, we union the corresponding relations and distinguish between the three partitions by appending a partition identifier to every relation.

Generalizing our example in a straightforward way, we show that trimmings of such inequalities exist for all acyclic JQs.

LEMMA 5.2. Given an acyclic $\mathcal{J}Q \ Q$, variables $U_w \subseteq \operatorname{var}(Q)$, weight functions $w_x : \operatorname{dom} \to \mathbb{R}$ for $x \in U_w$, and $\lambda \in \mathbb{R}$, a trimming of $\min_{x \in U_w} w_x(x) < \lambda$, $\min_{x \in U_w} w_x(x) > \lambda$, $\max_{x \in U_w} w_x(x) < \lambda$, or $\max_{x \in U_w} w_x(x) > \lambda$ takes time O(n) and returns an acyclic $\mathcal{J}Q$.

Combining Lemma 5.2 together with Lemma 4.1 and Lemma 3.3 gives us the following result:

THEOREM 5.3. Given an acyclic JQ over a database D of size n, MIN or MAX ranking function, and $\phi \in [0, 1]$, the %JQ can be answered in time $O(n \log n)$.

5.2 LEX

For lexicographic orders, we provided [7] an O(n) selection algorithm that can also be used for %JQs. Our divide-and-conquer approach can recover this result up to a logarithmic factor, i.e., our %JQ algorithm runs in time $O(n \log n)$. To achieve that, we trim lexicographic inequalities, similarly to the case of MIN and MAX.

LEMMA 5.4. Given an acyclic $\mathcal{J}QQ$, variables $U_w = \{x_1, \ldots, x_r\} \subseteq$ var(Q), weight functions $w'_x : \text{dom} \to \mathbb{R}$ for $x \in U_w$, and $\lambda \in \mathbb{R}^r$, a trimming of $(w'_{x_1}(x_1), \ldots, w'_{x_r}(x_r)) <_{\text{LEX}} \lambda$ or $(w'_{x_1}(x_1), \ldots, w'_{x_r}(x_r)) >_{\text{LEX}} \lambda$ takes time O(n) and returns an acyclic $\mathcal{J}Q$.

5.3 Partial SUM

We now consider the case of SUM. While we previously gave a dichotomy [7] for all self-join-free JQs, this result is limited to full SUM. We now provide a more fine-grained dichotomy where certain variables may not participate in the ranking. For example, the 3-path JQ $R_1(x_1, x_2), R_2(x_2, x_3), R_3(x_3, x_4)$ would be classified as intractable by the prior dichotomy, yet with weighted variables $U_w = \{x_1, x_2, x_3\}$, we show that it is in fact tractable.

The positive side of our dichotomy requires a trimming of additive inequalities. We rely on a known algorithm that can be applied whenever the SUM variables appear in adjacent join-tree nodes.

LEMMA 5.5 ([22]). Given a set of variables U_w , let Q be the class of acyclic JQs Q for which there exists a join tree where $U_w \subseteq var(Q)$ belong to adjacent join-tree nodes. Then, for all $Q \in Q$, weight functions $w_x : \text{dom} \to \mathbb{R}$ for $x \in U_w$, and $\lambda \in \mathbb{R}$, a trimming of $\sum_{x \in U_w} w_x(x) < \lambda$ or $\sum_{x \in U_w} w_x(x) > \lambda$ takes time $O(n \log n)$ and returns a JQ $Q' \in Q$.

We are now in a position to state our dichotomy:

THEOREM 5.6. Let Q be a self-join-free $\mathcal{J}Q, \mathcal{H}(Q)$ its hypergraph, and $U_w \subseteq \operatorname{var}(Q)$ the variables of a SUM ranking function.

- If H(Q) is acyclic, any set of independent variables of U_w is of size at most 2, and any chordless path between two U_w variables is of length at most 3, then %JQ can be answered in O(n log² n).
- Otherwise, %JQ cannot be answered in O(n polylog n), assuming 3SUM and HYPERCLIQUE.

We note that the positive side also applies to JQs with self-joins.

6 APPROXIMATE TRIMMING FOR SUM

We now move on to devise an ϵ -lossy trimming for additive inequalities in order to get a deterministic approximation for SUM and JQs beyond those covered by Theorem 5.6.

LEMMA 6.1. Given an acyclic JQ Q, variables $U_w \subseteq var(Q)$, weight functions $w_x : dom \to \mathbb{R}$ for $x \in U_w$, $\lambda \in \mathbb{R}$, and $\epsilon \in (0, 1)$, an ϵ -lossy trimming of $\sum_{x \in U_w} w_x(x) < \lambda$ or $\sum_{x \in U_w} w_x(x) > \lambda$ takes time $O(\frac{1}{\epsilon^2} n \log^2 n \log \frac{n}{\epsilon})$ and returns an acyclic JQ.

This, together with Lemmas 3.6 and 4.1 gives us the following:

THEOREM 6.2. Given an acyclic $\mathcal{J}QQ$ over a database D of size n, SUM ranking function, $\phi \in [0, 1]$, and $\epsilon \in (0, 1)$, the ϵ -approximate $\mathcal{J}Q$ can be answered in time $O(\frac{1}{\epsilon^2}n\log^5 n\log \frac{n}{\epsilon})$.

To achieve the trimming, we adapt an algorithm of Abo-Khamis et al. [2], which we refer to as APXCNT. It computes an approximate count (or more generally, a semiring aggregate) over the answers to acyclic JQs with additive inequalities. In contrast, we need not only the count of answers, but an efficient *relational representation* of them as JQ answers over a new database. We only discuss the case of less-than (<), since the case of greater-than (>) is symmetric. The detailed pseudocode is given in Appendix E.

Message passing. APXCNT uses message passing (see Section 2.4). We first describe the exact, but costly, version of the algorithm. The message sent by a tuple is a multiset containing the (partial) sums of partial query answers in its subtree. Messages are aggregated as follows: (1) The \oplus operator that combines multisets within a join group is multiset union (\uplus). (2) The \otimes operator that combines multisets from different children is pairwise summation (applied as a binary operator). The messages emitted by the rootnode tuples contain all query-answer sums, which can be leveraged to count the number of answers that satisfy the inequality.

Sketching. Sending all possible sums up the join tree is intractable since, in the worst case, their number is equal to the number of JQ answers. For this reason, APXCNT applies *sketching* to compress the messages. The basic idea is to replace different elements in a multiset with the same element; the efficiency gain is due to the fact that an element *s* that appears *r* times can be represented as $s \times r$. In more detail, the multiset elements are split into buckets, and subsequently, each element within a bucket is replaced by the maximum element of the bucket. A sketched multiset *L* is denoted by $\mathbb{S}_{\epsilon}(L)$, where ϵ is a parameter that determines the number of buckets. Let $\downarrow_{\lambda}(L)$ be the number of elements of *L* that are less than $\lambda \in \mathbb{R}$. By choosing buckets appropriately, we can guarantee that $\downarrow_{\lambda}(\mathbb{S}_{\epsilon}(L))$ is close to $\downarrow_{\lambda}(L)$ for all possible λ .

LEMMA 6.3 (ϵ -SKETCH [2]). For a multiset $L \in \mathbb{N}^{\mathbb{R}}$ and $\epsilon \in (0, 1)$, we can construct a sketch $\mathbb{S}_{\epsilon}(L)$ with $O(\log_{1+\epsilon} |L|)$ distinct elements such that for all $\lambda \in \mathbb{R}$, we have $(1 - \epsilon) \downarrow_{\lambda}(L) \leq \downarrow_{\lambda}(\mathbb{S}_{\epsilon}(L)) \leq \downarrow_{\lambda}(L)$.



Figure 3: Example of how we use the message passing framework [2] to create a relational representation of the query answers that satisfy an inequality $x + y + z < \lambda$.

ApxCNT sketches all messages and bounds the error incurred by the two message-passing operations (\oplus, \otimes) .

Relational representation. Our goal is to construct a relational representation of the JQ answers which satisfy the inequality that we want to trim. The key idea is to *embed the sums* contained in the messages of APXCNT into the database relations so that each tuple stores a *unique sum* and all answers in its subtree approximately have that sum. The reasoning behind this is that we can then remove from the database the root tuples whose associated sum does not satisfy the inequality. However, in APXCNT a message is a multiset of sums, and hence the main technical challenge we address below is how to achieve a unique sum per tuple (and its subtree).

Separating sums. Let $\sigma(t')$ be the message sent by a tuple t' in a child relation S. Then, according to APXCNT, a tuple t in the parent relation R receives a message $\sigma(b) = \mathbb{S}_{\epsilon'}(\uplus_{t' \in b} \sigma(t'))$ for some ϵ' and join group b. We separate the sums in this multiset by creating a number of copies of t, equal to the number of distinct sums in $\sigma(b)$. Each t-copy is associated with a unique bucket e, described by a sum value e_s and a multiplicity e_m . To avoid duplicating query answers, we restrict each t-copy to join only with the *source tuples* of its associated bucket e, i.e., the child tuples $t' \in S$ whose messages were assigned to bucket e during sketching.

Example 6.4. Figure 3 illustrates how we embed messages into the database relations for a leaf relation *S* and a parent relation *R* with no other children, and assuming weights equal to attribute values. The messages from S(x, y) to R(y, z) are the sums x + y (because *y*-weights are assigned to *S*). After sketching their union using two buckets, sums 4 and 5 are both mapped to 5; we keep track of this with a multiplicity counter (shown as ×2), reflecting the number of answers in the subtree. Upon reaching relation *R*, the weight of the *R*-tuple (which is the *z*-value) is added to all sums. For our relational representation, we duplicate the *R*-tuple and associate each copy with a unique sum. A copy corresponds to a bucket in the sketch, so we can trace its "source" *S*-tuples, i.e., those that belong to that bucket. Instead of joining with all *S*-tuples like before, a copy now only joins with the source *S*-tuples of the bucket via a new variable v_{RS} that stores the sum and the multiplicity.

Adjusting the sketch buckets. An issue we run into with the approach above is that the sum sent by a single tuple may be *assigned to more than one bucket* during sketching. To see why this is problematic, consider a tuple t' that sends $\sigma(t') = 5 \times 10$. For simplicity, assume these are the only values to be sketched and that the two buckets contain 5×3 and 5×7 . With these buckets, we will

create two copies of a tuple t in the parent and both will join with t', because t' is the source tuple for both buckets. By doing so, we have effectively doubled the number of (partial) query answers since there are now 10 answers in the subtree of each copy. To resolve this issue, we need to guarantee that all elements in $\sigma(t')$ are assigned to the same bucket. We adjust the sketching $\mathbb{S}_{\epsilon}(L)$ of a multiset L as follows. The *r* buckets are determined by an increasing sequence of r + 1 indexes on an array that contains L sorted. The first index is 0 and the last index is equal to |L| - 1 (where |L| takes into account the multiplicities). Consider three consecutive indexes i, j, k which define two consecutive buckets where the values at the borders are the same, i.e., L[j-1] = L[j]. Let j' and j'' be the smallest and largest indexes that contain L[j] in the buckets i - j and j - k, respectively. We replace the indexes *i*, *j*, *k* with *i*, j', j'' + 1, *k* (and if 2 consecutive indexes are the same, then we remove that bucket). As a result, all values *L*[*j*] from these two buckets now fall into the same bucket. We repeat this process for every two consecutive buckets. This can at most double the number of buckets, which, as we show, does not affect the guarantee of Lemma 6.3.

Binary join tree. The running time of our algorithm (in particular the logarithmic-factor exponent) depends on the maximum number of children of a join-tree node. This is because we handle each parent-child node pair separately, and each child results in the parent relation growing by the size of the messages, which is logarithmic. To achieve the time bound stated in Lemma 6.1, we impose a binary join tree, i.e., every node has at most two children. Such a join tree can be constructed by creating copies of a node that has multiple children, connecting these copies in a chain, and distributing the original children among them. In the worst case, this doubles the number of nodes in the join tree (hence the number of relations that we materialize), but it does not affect the data complexity.

7 CONCLUSIONS

We can often answer quantile queries over joins of multiple relations much more efficiently than it takes to materialize the result of the join. Here, we adopted *quasilinear time* as our yardstick of efficiency. With our divide-and-conquer technique, we recovered known results (for lexicographic orders) and established new ones (for partial sums, minimum, and maximum). We also showed how the approach can be adapted for deterministic approximations.

We restricted the discussion to JQs, that is, full Conjunctive Queries (CQs), and left open the treatment of non-full CQs (i.e., joins with projection). Most of our algorithms apply to CQs that are acyclic and free-connex, but it is not yet clear to us whether our results cover all tractable cases (under complexity assumptions). More general open directions are the generalization of the challenge to *unions* of CQs, and the establishment of nontrivial algorithms for general CQs beyond the acyclic ones.

ACKNOWLEDGMENTS

This work was supported in part by NSF under award numbers IIS-1762268 and IIS-1956096. Benny Kimelfeld was supported by the German Research Foundation (DFG) Project 412400621. Nikolaos Tziavelis was supported by a Google PhD fellowship.

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Symbol	Definition
Ζ	generic set
L	generic multiset
R, S, T, R_1, R_2	relation
V, V_1, V_2	atom/hyperedge/node of join tree
S	schema
D	database (instance)
n	size of D (number of tuples)
dom	database domain
t	tuple
x, y, z, u, v	variable
Q	Join Query (JQ)
l	number of atoms in a JQ
var(Q)	variables contained in Q
Q(D)	set of answers of Q over D
$(Q \land P)(D)$	subset of $Q(D)$ answers that satisfy a predicate P
$q \in Q(D)$	query answer
$\mathcal{H}(Q) = (V, E)$	hypergraph associated with query Q
Т	join tree
ϕ	fraction in [0, 1] used to ask for a quantile
w	weight function for query answers
dom_w	domain of weights
U_w	subset of variables that participate in the ranking
w_{χ}	input weight function for variable $x: \operatorname{dom} \to \operatorname{dom}_w$
w _R	input weight function for tuples of relation $R: \operatorname{dom}^{\operatorname{ar}(R)} \to$
	dom_w where $ar(R)$ is the arity of R
agg_w	aggregate function that combines input weights to derive
	weights for query answers
$w(U_w)$	aggregate function applied on the weighted variables, i.e.,
	$\arg_w(\{w_x(x) x \in U_w\})$
λ	a weight from dom_w

A NOMENCLATURE

B DETAILS OF DIVIDE-AND-CONQUER FRAMEWORK

Algorithm 1 returns the desired (approximate) quantile for a given JQ, database, and ranking function, as presented in Section 3. The exact version is obtained by simply setting $\epsilon = 0$.

B.1 Proof of Lemma 3.6

Let Q_i and D_i be the JQ and database at the start of iteration i, with $i \ge 0$ (these are the variables Q' and D' in Algorithm 1). Even though trimmings may increase the size of our queries by a constant factor (by the definition of trimming), all queries Q_i have constant size. This is because we start every iteration with the original query Q and every other query we construct is the result of applying at most two consecutive trimmings.

First, we bound the number of iterations. Iteration *i* is guaranteed to eliminate at least $c|Q_i(D_i)|$ query answers because (i) we select a *c*-pivot to partition and (ii) the lossy trimmings may result in more query answers being eliminated than they should, but never less. Consequently, at the beginning of iteration *i*, we have at most $(1 - c)^i |Q(D)|$ query answers remaining. The number of query answers is bounded by n^ℓ where ℓ is the number of atoms in *Q*. If *I* is the total number of iterations, then $I \leq \lceil \log_{1/(1-c)} n \rceil = O(\log n)$ since *c* and ℓ are constants.



Figure 4: Proof of Lemma 3.6: Query answers that are "lost" due to lossy trimmings are implicitly moved to the equal-to partition (middle). Consequently, index k_i in the less-than partition contains an element that was previously at a higher index k'_i , but $k'_i - k_i$ is bounded by the number of lost answers.

Second, we show that the returned answer is an ϵ -approximate quantile. The less-than partition $Q_{1t}(D_{1t})$ is constructed by trimming the inequalities $w(U_w) \prec w(p)$ and $w(U_w) > 1$ ow with some error ϵ' , where low lower-bounds the weights of the candidate query answers. Because these two trimmings are lossy, we "lose" a number of query answers which are at most $2\epsilon' |Q(D)|$. These are the answers that satisfy the inequalities, but do not appear in $Q_{1t}(D_{1t})$. As Figure 4 illustrates, all answers not contained in the less-than or greater-than partition, including these lost query answers, are assumed to be contained in the equal-to partition which we do not explicitly count. We now bound the distance between the desired index and the index of the answer that our algorithm returns. Each iteration *i* starts with an index k_i and results in a new index k'_i , which the following iteration is asked to retrieve (or, in case this is the last iteration, the index that is returned). Note that in Algorithm 1 the variable k indexes the subarray of query answers that are currently candidates; thus it is offset by the index of the answer with weight low. Here, the indexes k_i and k'_i refer to the original array that contains all the query answers. Suppose that k_i falls into the less-than partition at the beginning of the iteration. Then, if k'_i is different than k_i , it has to be a higher index because of lost query answers that precede it and which are moved to the middle equal-to partition (see Figure 4). Thus, $|k_i - k'_i| \le 2\epsilon' |Q(D)|$. If k_i falls into the equal-to partition, then we still choose that partition and return the pivot because the size of the partition can only increase from the lossy trimmings. For the greater-than partition, the analysis is symmetric to lower-than since the lossy trimmings of the latter do not affect the indexes of the former. To conclude, the accumulated absolute error is $I \cdot 2\epsilon' |Q(D)| \le 2\lceil \ell \log_{1/(1-c)} n \rceil \epsilon' |Q(D)|$. To obtain an ϵ -approximate quantile of Q(D), we set $\epsilon' = \frac{\epsilon}{2\lceil \ell \log_{1/(1-\epsilon)} n \rceil}$.

Finally, we prove the running time. Since our trimmings return acyclic JQs, the answers of all queries we construct can be counted in linear time. Thus, the running time per iteration is $O(g_p(n) + 4g_t(n, \epsilon') + n)$ which is $O(\max\{g_p(n), g_t(n, \epsilon')\}$ since $g_p(n)$ and $g_t(n, \epsilon')$ are necessarily $\Omega(n)$.

We note that this proof also covers Lemma 3.3 since Lemma 3.6 is a stronger version of it.

C DETAILS OF CHOOSING A PIVOT

Algorithm 2 shows the algorithm that returns a *c*-pivot for a given JQ, database, and ranking function, as presented in Section 4.

Efficient Computation of Quantiles over Joins

Algorithm 1: Pivoting Algorithm

¹ Input : acyclic JQ Q , database D , ranking function (w , ≤), quantile	φ,
approximation bound ϵ	
² Output : the ϕ -th quantile of $Q(D)$	
3 //Calculate desired index	
⁴ Determine $ Q(D) $ and set $k = \lfloor \phi \cdot Q(D) \rfloor$ (with zero-indexing)	
5 //Calculate parameter for trimming ($\epsilon = \epsilon' = 0$ for exact)	
$\epsilon \epsilon' = \frac{\epsilon}{2\lceil \ell \log_{1/(1-\epsilon)} n \rceil}$	
7 //Each iteration modifies $Q'(D')$ by bringing low and high closer	
$(Q', D', \text{low}, \text{high}) = (Q, D, \bot, \top)$	
9 while $ Q'(D') > D $ do	
10 //Select a c -pivot p	
11 $(p,c) = \operatorname{PIVOT}(Q',D',(w,\leq))$	
12 //Partition	
13 $(Q_{1t}, D_{1t}) = \operatorname{trim}(Q, D, w(U_w) \prec w(p), \epsilon')$	
14 $(Q_{lt}, D_{lt}) = \text{trim}(Q_{lt}, D_{lt}, w(U_w) > \text{low}, \epsilon')$	
15 $(Q_{gt}, D_{gt}) = \operatorname{trim}(Q, D, w(U_w) \succ w(p), \epsilon')$	
$(Q_{\text{gt}}, D_{\text{gt}}) = \text{TRIM}(Q_{\text{gt}}, D_{\text{gt}}, w(U_w) \prec \text{high}, \epsilon')$	
17 //Choose partition	
18 Set $ Q_{eq}(D_{eq}) $ to $ Q'(D') - Q_{lt}(D_{lt}) - Q_{gt}(D_{gt}) $	
19 if $k < Q_{lt}(D_{lt}) $ then	
$(Q', D', high) = (Q_{1t}, D_{1t}, w(p))$	
else if $k < Q_{lt}(D_{lt}) + Q_{eq}(D_{eq}) $ then	
²² return <i>p</i>	
23 else	
24 $(Q', D', 1t) = (Q_{gt}, D_{gt}, w(p))$	
25 $k = k - Q_{1t}(D_{1t}) - Q_{eq}(D_{eq}) $	
Materialize and sort $Q'(D')$	
²⁷ return answer at index k in $Q'(D')$	

C.1 Proof of Lemma 4.4

Let b_1, \ldots, b_r be the joining groups from the children V_1, \ldots, V_r of V, We show by induction on the join tree that $pivot(b_1), \ldots, pivot(b_r)$, and t all agree on their common variables. For the leaf relations, we have no children and pivot(t) is initialized to t. For the inductive step, let x be a common variable between two children V_i and V_j of $V, i, j \in [r]$. Because of the running intersection property of the join tree, x also needs to appear in the parent V. Since the groups b_i, b_j join with t, all their tuples necessarily assign value t[x] to variable x. We show that $pivot(b_i)$ also assigns t[x] to x and the case of $pivot(b_j)$ is similar. We have that $pivot(b_i) = pivot(t_i)$ for some tuple $t_i \in b_i$ where $pivot(t_i)$ is picked as the weighted median of the group. From the inductive hypothesis, $pivot(t_i)$ needs to agree with t_i on the value of x which we argued is equal to t[x].

C.2 Proof of Lemma 4.5

Without loss of generality, let the indexing of the *r* sets be consistent with the ordering of their *c*-pivots, i.e., $p_i \leq p_j$ for $Z_i, Z_j, 1 \leq i \leq j \leq r$. Let p_m be the weighted median, selected from set Z_m for some $m \in [r]$. We prove that p_m is greater than or equal to (according to \leq) at least $\frac{c}{2}|Z_1 \cup Z_2 \cup \ldots \cup Z_r|$ elements, and the case of less than or equal to is symmetric. Because of the indexing we enforced, we know that $p_i \leq p_m$ for all $i \in [m]$. Combining that with the definition of a *c*-pivot (for p_i), we obtain that p_m is greater than or equal to at least $c|Z_i|$ elements of $|Z_i|$, or $c \sum_{i \in [1,m]} |Z_i|$ in total. Now, because the median is weighted by the set sizes and there is no overlap between their elements, $\sum_{i \in [1,m]} |Z_i| \geq |Z_1 \cup Z_2 \cup \ldots \cup Z_r|/2$. Thus, p_m is greater than or equal to at least $c|Z_1 \cup Z_2 \cup \ldots \cup Z_r|/2$ elements of $Z_1 \cup Z_2 \cup \ldots \cup Z_r$.

Algorithm 2: PIVOT

1	Input : acyclic JQ <i>Q</i> , database <i>D</i> , ranking function (w, \leq)		
2	Output : a <i>c</i> -pivot of $Q(D)$ ordered by \leq , and the value of <i>c</i>		
3	Convert attribute weights to tuple weights		
4	Construct a join tree T of Q with artificial root $V_0 = \{t_0\}$		
5	Materialize a relation for every T -node and group it by the variables it has in		
	common with its parent node		
6	Initialize $pivot(t) = t$, $cnt(t) = 1$ for all tuples t of all relations		
7	Initialize $c(R) = 1$ for leaf relations R		
8	for relation R in bottom-up order of T do		
9	if R is not leaf then		
10	S_1, \ldots, S_r = children of R		
11	$S_1, \dots, S_r = \text{children of } R$ $c(R) = \frac{c(S_1)}{2} \times \dots \times \frac{c(S_r)}{2}$		
12	for tuple $t \in R$, child S of R do		
13	b = join group of S that agrees with the values of t		
14	//Compute the weighted median (and the count of subtree answers)		
	the first time we visit this group		
15	if <i>pivot</i> (<i>b</i>) <i>not already computed</i> then		
16	pivot(b) = WMED \leq ({pivot(t') t' \in b}, β) with		
	$\beta(\operatorname{pivot}(t')) = \operatorname{cnt}(t')$		
17	$\operatorname{cnt}(b) = \sum_{t' \in b} \operatorname{cnt}(t')$		
18	//Combine results from different branches of the join tree		
19	$pivot(t) = pivot(t) \cup pivot(b)$		
20	$\operatorname{cnt}(t) = \operatorname{cnt}(t) \times \operatorname{cnt}(b)$		
21	return pivot $(t_0), c(V_0)$		

C.3 Proof of Lemma 4.6

Let *M* be the partial answers rooted at *t*, and let M_i be the partial answers rooted at V_i and restricted to those that agree with *t* for all $i \in [r]$. We only show that $t \cup p_1 \cup \ldots \cup p_r$ is greater than or equal to at least $c^r |M|$ partial answers, since the case of less than or equal to is symmetric. For $i \in [r]$, let L_i be the subset of M_i answers that are less than or equal to p_i .

We first show that $w(t \cup q_1 \cup \ldots \cup q_r) \leq w(t \cup p_1 \cup \ldots \cup p_r)$ whenever $q_i \in L_i, i \in [r]$. We know that $w(q_i) \leq w(p_i)$ for all $i \in [r]$. We proceed inductively in *i*, showing that $w(q_1 \cup \ldots \cup q_i) \leq i$ $w(p_1 \cup \ldots \cup p_i)$. The inductive hypothesis is that $w(q_1 \cup \ldots \cup q_{i-1}) \leq d_i$ $w(p_1 \cup \ldots \cup p_{i-1})$. Each of these two terms is an aggregate over the values of variables $U_1 \cup \ldots \cup U_{i-1}$ mapped to their weights, where U_i is the subset of weighted variables U_w that appear in the subtree rooteed at node V_i . For example, $w(q_1 \cup \ldots \cup q_{i-1})$ is $\arg_{w}(\{w_{x}((q_{1} \cup \ldots \cup q_{i-1})[x]) \mid x \in U_{1} \cup \ldots \cup U_{i-1}\})$. By subset-monotonicity, we can add to both aggregates the weighted values of p_i without changing the inequality, i.e., we obtain $w(q_1 \cup$ $\dots \cup q_{i-1} \cup p_i) \leq w(p_1 \cup \dots \cup p_{i-1} \cup p_i)$ (A). With a similar argument of subset-monotonicity, we start from $w(q_i) \leq w(p_i)$ and add to both sides the weighted values of $q_1 \cup \ldots \cup q_{i-1}$ to obtain $w(q_1 \cup \ldots \cup q_{i-1} \cup q_i) \leq w(q_1 \cup \ldots \cup q_{i-1} \cup p_i)$ (B). (A) and (B) together prove the inductive step. To complete the first part of the proof, we add to both aggregates the weighted values of t (that do not appear in any child) to obtain $w(t \cup q_1 \cup \ldots \cup q_r) \leq dt$ $w(t \cup p_1 \cup \ldots \cup p_r)$, again by subset-monotonicity.

Next, notice that there are $|L_1 \times \ldots \times L_r|$ partial answers of the form $t \cup q_1 \cup \ldots \cup q_r$ with $q_i \in L_i, i \in [r]$. Since every L_i comprises of elements that are less than or equal to a *c*-pivot, we have $|L_i| \ge c|M_i|$. Also notice that $|M| = \prod_{i \in [r]} |M_i|$. Overall, we have that $|L_1 \times \ldots \times L_r| \ge \prod_{i \in [r]} c|M_i| = c^r|M|$, and so $t \cup p_1 \cup \ldots \cup p_r$ is greater than or equal to at least $c^r|M|$ partial answers.

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Algorithm 3: TRIM for MAX

1	Input : acyclic JQ Q , database D , predicate $\max(U_w) > \lambda$
2	Output : acyclic JQ Q' , database D'
3	$x_1,\ldots,x_r=U_w$
4	$(Q',D')=(Q,\emptyset)$
5	//Construct the new JQ
6	Eliminate self-joins from Q' by materializing new relations in D
7	Add the same variable x_p to all the atoms and the head of Q'
8	//Create <i>r</i> databases
9	for i from 1 to r do
10	//Each P_i is a conjuntion of unary predicates
11	$P_i = \{ w_{x_1}(x_1) \le \lambda, \dots, w_{x_{i-1}}(x_{i-1}) \le \lambda, w_{x_i}(x_i) > \lambda \}$
12	$D_i = \text{copy of } D$ with conditions P_i applied
13	//An identifier separates the answers from different D_i after the union
14	Add the column x_p with value <i>i</i> to all relations of D_i
15	//Union the databases into one
16	for relation \mathbb{R}^D in D do
17	Add to D' relation $R^{D'} = \bigcup_{i \in [r]} R^{D_i}$ of database D_i
18	return (Q', D')

D DETAILS OF EXACT TRIMMINGS

D.1 Proof of Lemma 5.2

We always start by creating fresh copies of relations to eliminate selfjoins from Q. This ensures that every column in the database corresponds to a unique variable, avoiding situations like R(x, y), R(y, x).

First, consider $\max_{x \in U_w} w_x(x) < \lambda$. We scan the given database D once and if a tuple t contains a value t[x] with $w_x(t[x]) \ge \lambda$ for a variable $x \in U_w$, then we remove t from the database. This process removes precisely the answers $q \in Q(D)$ that do not satisfy the predicate, since, for the maximum to be greater than or equal to λ , at least one variable needs to map to such a weight. The JQ we return is Q itself. The case of $\min_{x \in U} w_x(x) > \lambda$ is symmetric.

Second, consider $\max_{x \in U_w} w_x(x) > \lambda$. Algorithm 3 shows the pseudocode of TRIM for this case. If there are r variables in U_w , then we create r databases, each enforcing condition P_i , which is a conjunction of unary predicates. The conditions P_i partition the space of possible U_w values that satisfy $\max_{x \in U_w} w_x(x) > \lambda$. To return a single database D', we union together the copies of each relation and separate the different databases with a partition identifier $i \in [r]$. This identifier is added as a variable x_p to all atoms of the returned JQ Q'. As a consequence, each query answer of the returned Q' can only draw values from database tuples that belong to the same partition. The bijection from Q'(D') to Q(D) simply removes the variable x_p . Since r does not depend on D, the entire process can be done in linear time. Furthermore, Q' remains acyclic because every join tree of Q is also a join tree of Q' by adding x_p to all nodes. The case of $\min_{x \in U_w} w_x(x) < \lambda$ is symmetric.

D.2 Proof of Lemma 5.4

Let $\lambda = (\lambda_1, ..., \lambda_r)$. The proof is the same as in the case of MIN/MAX (Appendix D.1), except that the conditions we enforce in the *i*th of the *r* copies of the database *D* are $P_i = \{w'_{x_1}(x_1) = \lambda_1, ..., w'_{x_{i-1}}(x_{i-1}) = \lambda_{i-1}, w'_{x_i}(x_i) < \lambda_i\}$ for \leq_{LEX} and $P_i = \{w'_{x_1}(x_1) = \lambda_1, ..., w'_{x_{i-1}}(x_{i-1}) = \lambda_{i-1}, w'_{x_i}(x_i) > \lambda_i\}$ for \geq_{LEX} .

D.3 Proof of Theorem 5.6

First, we prove that the condition in our dichotomy is equivalent to having the SUM variables on one or two adjacent join tree nodes.

LEMMA D.1. Consider the hypergraph $\mathcal{H}(Q)$ of a JQQ and a set of variables U_w . If $\mathcal{H}(Q)$ is acyclic, any set of independent variables of U_w is of size at most 2, and any chordless path between two U_w variables is of length at most 3, then there exists a join tree for Qwhere U appears on one or two adjacent nodes.

PROOF. If there is one query atom that contains all U_w variables, then we are done. Otherwise, since any set of independent variables of U_w is of size at most 2, then there are 2 atoms that together contain all U_w variables. Indeed, consider any 3 atoms. If each of them has a U_w variable that does not appear in the other two, then these three variables are an independent set of size 3, which contradicts our condition. Thus, 2 of these atoms contain all U_w variables that appear in the 3 atoms. By applying this repeatedly to the selected 2 atoms and an untreated atom until all atoms are treated, we get 2 atoms that contain all of U_w variables.

Since Q is acyclic, it has a join tree. Let R' and S' be two join-tree nodes that together contain all of U_w . Consider the path P' from R' to S' in the join tree. Let R be the last node on P' that contains all U_w variables that are in R', and let S be the first node on P' that contains all U_w variables that are in S'. If R and S are neighbors, we are done. Otherwise, we show we can find an alternative join tree where they are neighbors. Consider the path P from R to Sin the join tree. Let V be all the variables that appear on the path between *R* and *S* (not including *R* and *S*), such that each variable in V appears in either R or S (or both). We consider three cases. The first case is $V \subseteq R$. We directly connect R and S and remove the edge connecting S to the node preceding it on the path from R. The running intersection property is maintained as for each variable, the nodes containing this variable remain connected. The second case is $V \subseteq S$. It is handled similarly by directly connecting *R* to *S* and removing the edge from R to its succeeding node on the path to *S*. The third case is that a variable $u \in V$ appears in *R* but not in *S* and another variable $v \in V$ appears in *S* but not in *R*. Since *R* is the last in *P* to contain all U_w variables of *R'*, there is a variable $x \in U_w$ that appears in *R* but nowhere else in *P*. Similarly, there is a variable $y \in U_w$ that appears in *S* and nowhere else in *P*. If every two consecutive nodes on P share a variable, then we have a chordless path $x - u - \ldots - v - y$ of length at least 4, contradicting our condition. Otherwise, we remove the edge between the two nodes that do not share a variable, and add an edge between R and *S*, which preserves the running intersection property.

We now show the dichotomy of Theorem 5.6.

For the positive side, we apply Lemma D.1. When all U_w variables are contained in a single join-tree node, trimming can be done in linear time by filtering the corresponding relation. When they are contained in two adjacent join-tree nodes, $O(n \log n)$ trimming follows from Lemma 5.5. Combining these two cases with Lemmas 3.3 and 4.1 completes the proof of the positive side.

For the negative side, there are 3 cases. 1) If Q is cyclic, an answer to %JQ would also answer the decision problem of whether Q has any answer, which precludes time O(n polylog n) assuming HYPERCLIQUE [6]. Assume Q is acyclic. 2) If there exists a set of

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Algorithm 4: Approximate TRIM for SUM		
¹ Input : acyclic JQ Q with ℓ atoms, database D , predicate		
$\sum_{x \in U_W} w_x(x) < \lambda$, approximation bound ϵ		
2 Output : acyclic JQ Q' , database D'		
3 Convert attribute weights to tuple weights		
⁴ Construct a binary join tree T of Q , set an arbitrary root		
Materialize a relation for every T -node and group it by the variables it has in		
common with its parent node		
6 Initialize $\sigma(t) = (\sigma_s(t), \sigma_m(t)) = (w(t), 1)$ for all tuples t of all relations		
7 $\epsilon' = \frac{1}{4\ell}\epsilon$		
s for relation R in bottom-up order of T do		
9 for child S of R do		
Add variable v_{RS} to R and S in Q , and corresponding columns in D		
11 for tuple $t \in R$, child S of R do		
b = join group of S that agrees with the values of t		
13 //Sketch messages the first time we visit this group		
14 if $\sigma(b)$ not already computed then		
15 $\sigma(b) = \mathbb{S}_{\epsilon'}(\cup_{t' \in b} \sigma(t'))$ such that each value falls into a		
single bucket		
16 //A bucket e in the sketch is described by a sum e_s ,		
multiplicity e_m , and a set of source tuples from S		
17 for bucket $e \in \sigma(b)$ with source tuples $S_e \subseteq S$ do		
18 //Add the bucket values to the child column		
19 $t_e[v_{RS}] = (e_s, e_m)$ for all $t_e \in S_e$		
for bucket $e \in \sigma(b)$ do		
21 //Add the bucket values to the parent column		
22 Create a copy t_e of t in R with $t_e[v_{RS}] = (e_s, e_m)$		
23 $\sigma(t_e) = (\sigma_s(t) + e_s, \sigma_m(t) \times e_m)$		
24 Remove t from R		
25 Remove all tuples <i>t</i> from the root relation with $\sigma_s(t) \ge \lambda$		
$_{26}$ return Q, D		

independent variables of U_w of size 3, selection by SUM is not possible in $O(n^{2-\varepsilon})$ for all $\varepsilon > 0$ assuming 3SUM [7, Corollary 7.11]. Since we can count the answers to an acyclic JQ in linear time, the selection problem and %JQ are equivalent. 3) If there is a chordless path between two U_w variables of length 4 or more, we apply a known reduction [7, Lemma 7.13] to show that solving %JQ in quasilinear time can be used to detect a triangle in a graph in quasilinear time, which is not possible assuming HYPERCLIQUE. There are two ways in which the statement of that lemma differs from our needs: first, all variables there were allowed to participate in the ranking. However, the reduction only assigns non-zero weights to the first and last variables in the path, so this difference is non-essential. Second, the path there contains exactly 3 atoms (i.e., 4 variables); if our path is longer, we simply make the the remaining relations equality, and the rest of the proof is the same.

E DETAILS OF APPROXIMATE TRIMMING FOR SUM

Algorithm 4 shows the pseudocode of our lossy trimming for SUM.

E.1 Proof of Lemma 6.1

Preservation of JQ answers. Let Q' and D' be the returned JQ and database. We argue that, before removing the root tuples that violate the inequality (Line 25), the JQ answers are preserved in the sense that there exists a bijection from Q'(D') to Q(D) which simply removes the new variables. Consider the step where we introduce variable v_{RS} between parent R and child S. Let $t \in R$ be a tuple in the original database D and b the join group in S that

agrees with *t*. Then, every tuple $t' \in b$ joins with exactly one copy of *t* after the introduction of v_{RS} . This is because there is a copy of *t* for each bucket (with the bucket identifier in v_{RS}) and our bucket adjustment guarantees that the weight of *t'* is assigned to precisely one bucket.

Error from sketch adjustment. Recall that in our sketch $\mathbb{S}_{\epsilon}(L)$ of a multiset L we made the adjustment that if i, j, k are three consecutive indexes in the bucketization, L[j - 1] = L[j], and i', i'' are the smallest and largest indexes that contain L[i] in the two consecutive buckets, then we replace *i*, *j*, *k* with *i*, j', j'' + 1, *k*. We say that a multiset is an ϵ -sketch of another multiset if it satisfies the guarantee of Lemma 6.3. Also, let S be the original sketch with approximation error ϵ (see Lemma 6.3) and S' be the resulting sketch. What we will show is that S' is an ϵ -sketch of L. In particular, we claim that $\downarrow_{\lambda}(\mathbb{S}) \leq \downarrow_{\lambda}(\mathbb{S}') \leq \downarrow_{\lambda}(L)$ for all values of λ . Our adjustment can only change elements in the index ranges [i, j')and [j, j''), while all other elements stay the same since the largest element in their bucket continues to be the same. The elements that can potentially change may only decrease in value because the upper index of their bucket is now smaller (but they may not decrease beyond L[i] and L[j] respectively). Consequently, if $\lambda < 1$ L[i] or $\lambda \ge L[j'']$ then $\downarrow_{\lambda}(\mathbb{S}) = \downarrow_{\lambda}(\mathbb{S}')$. If $L[i] \le \lambda < L[j]$, then $\downarrow_{\lambda}(\mathbb{S}) \leq \downarrow_{\lambda}(\mathbb{S}')$ because all elements in this bucket were mapped to Z[i] in S but now they are mapped to a number that can only be smaller, and thus closer to their original value. If $L[j] \leq \lambda < L[j']$, all elements in that bucket are equal to L[j], thus $\downarrow_{\lambda}(\mathbb{S}') = \downarrow_{\lambda}(L)$.

Approximation guarantee. Let us introduce the notation and tools we need. Recall that each tuple *t* computes $\sigma(t)$ that represents the approximate sum of partial query answers in its subtree. Let $cp(t) = \{t_1, \ldots, t_r\}$ be the copies of t that we create in our algorithm, W_t be the partial query answers in the subtree of t mapped to their weights, $jg_{S}(t)$ be the join group of relation S that joins with a tuple *t* of the parent relation, and \otimes be the pairwise summation operator for multisets. Abo-Khamis et al. [2] have shown that if L'_1 is an ϵ_1 -sketch of L_1 and L'_2 is an ϵ_2 -sketch of L_2 , then $L'_1 \uplus L'_2$ is a $\max{\epsilon_1, \epsilon_2}$ -sketch of $L_1 \uplus L_2$ and $L'_1 \otimes L'_2$ is an $(\epsilon_1 + \epsilon_2)$ -sketch of $L_1 \otimes$ *L*₂. Additionally, an ϵ_1 -sketch of an ϵ_2 -sketch is a $(2 \max{\epsilon_1, \epsilon_2})$ sketch (using the definition and that $(1 - \epsilon)^2 \ge 1 - 2\epsilon$). With these, we will show that the removal of root-node tuples (Line 25) removes the JQ answers that fall into buckets with values greater than or equal to λ in an ϵ -sketch of the multiset $\{w(q) | q \in Q(D)\}$. Note that in the algorithm, we apply sketching with $\epsilon' \leq \epsilon$ (Line 15).

First, we prove inductively that for a tuple $t \in R$ where R is a relation at level d (i.e., the maximum-length path from R to a leaf node is d), $end time_{i_i \in \operatorname{cp}(t)} \sigma(t_i)$ is a $(4^d \epsilon')$ -sketch of W_t . Each weight in W_t is the sum of the weight of t and the weights of the joining partial answers (in the original database D) from the child relations, i.e., $W_t = \{w(t)\} \otimes (\bigotimes_S \bigcup_{t' \in jg_S(t)} W_{t'})$). If t joins with a tuple t' of a child relation S, then it needs to join with all copies $\operatorname{cp}(t')$ that were created when we handled S and its children. The algorithm computes the values $\sigma(t_i)$ as follows: $eqtildelta_i \in \operatorname{cp}(t) \sigma(t_i) = \{w(t)\} \otimes (\bigotimes_S \mathbb{S}_{\epsilon'}(\biguplus_{t' \in jg_S(t)} \oiint_{t'_j \in \operatorname{cp}(t')} \sigma(t'_j)))$. We know inductively that $eqtildelta_{t'_j \in \operatorname{cp}(t')} \sigma(t'_j)$ is a $(4^{d-1}\epsilon')$ -sketch of $W_{t'}$. The error bound of the sketch remains $4^{d-1}\epsilon'$ after the union, then becomes $2 \cdot 4^{d-1}\epsilon'$ after applying the ϵ' -sketch, and finally $2 \cdot 2 \cdot 4^{d-1}\epsilon' = 4^d\epsilon'$ after taking the pairwise sums between the two children.

Second, we claim that the height of the binary join tree we construct is no more than ℓ , where ℓ is the number of atoms of Q. To see why, note that the new nodes we introduce in order to make the tree binary cannot be leaves and will always have 2 children. Suppose that there exists a root-to-leaf path of length greater than ℓ . For every new node on the path, there must be an original node that is a descendant of it, but not on this path. This implies that the number of original nodes would be greater than ℓ , which is a contradiction. To conclude, we get ϵ -sketches of W_t for tuples t at the root level if we set $\epsilon' = \frac{1}{4t}\epsilon$. Their union is an ϵ -sketch of $\{w(q)|q \in Q(D)\}$.

Returned JQ properties. The fact that the JQ Q' that we return is acyclic is evident from the fact that every variable v_{RS} that we introduce appears in two adjacent nodes of the join tree of Q. Therefore, Q' also has a join tree.

Running time. The size of any relation is initially bounded by *n*. Consider the step where we handle a relation and increase its size by

creating copies of its tuples. The sizes of the child relations (which have already been handled) have size bounded by $n' \ge n$. The total size of the messages sent from the children is $O(\log_{1+\epsilon'} n')$ because the messages are sketched. The parent relation receives the messages of a child and creates copies of its tuples whose number is equal to the message size. Since we have at most 2 children, the size of the parent relation becomes $O(n(\log_{1+\epsilon'} n')^2)$. Applying this for every relation bottom-up, we can conclude that all relations after the algorithm terminates have size $O(n(\log_{1+\epsilon'} n)^2)$ (because the double-logarithmic terms are dominated). Changing base, this is $O(n\frac{\log^2 n}{\log^2(1+\epsilon')}) \text{ or } O(\frac{1}{\epsilon^2}n\log^2 n) \text{ since } \epsilon' = \Theta(\epsilon) \text{ and also } \log(1+\epsilon)$ is very close to ϵ for small ϵ . All other operations of the algorithm are linear in this size, except for sketching, which is only done once for each join group. A sketch of a multiset $L = (Z, \beta)$ can be computed in $O(|Z| \log |Z|)$ by sorting. Since $O(\log(\frac{1}{\epsilon^2} n \log^2 n)) =$ $O(\log \frac{n}{\epsilon})$, we get the desired time bound $O(\frac{1}{\epsilon^2} n \log^2 n \log \frac{n}{\epsilon})$.