Trajectory Data Warehouses: Design and Implementation Issues

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In this paper we investigate some issues and solutions related to the design of a Data Warehouse (DW), storing several aggregate measures about trajectories of moving objects. First we discuss the loading phase of our DW which has to deal with overwhelming streams of trajectory observations, possibly produced at different rates, and arriving in an unpredictable and unbounded way. Then, we focus on the measure presence, the most complex measure stored in our DW. Such a measure returns the number of distinct trajectories that lie in a spatial region during a given temporal interval. We devise a novel way to compute an approximate, but very accurate, presence aggregate function, which algebraically combines a bounded amount of measures stored in the base cells of the data cube.

We conducted many experiments to show the effectiveness of our method to compute such an aggregate function. In addition, the feasibility of our innovative trajectory DW was validated with an implementation based on Oracle. We investigated the most challenging issues in realizing our trajectory DW using standard DW technologies: namely, the preprocessing and loading phase, and the aggregation functions to support OLAP operations.

Categories and Subject Descriptors: E.2 [DATA STORAGE REPRESENTATIONS];

General Terms: Design

Additional Key Words and Phrases: Data Warehouse, Aggregate Function

1. INTRODUCTION

Modern location-aware devices and applications deliver huge quantities of spatio-temporal data concerning moving objects, which must be either quickly processed for real-time applications, like traffic control management, or carefully mined for complex, knowledge discovering tasks. Even if such data usually originate as timed,
located observations of well identified objects, they must often be stored in aggregate form, without identification of the corresponding moving objects, either for privacy reasons, or simply for the sheer amount of data which should be kept online to perform analytical operations. Such an aggregation is usually a complex task, and prone to the introduction of errors which are amplified by subsequent aggregation operations.

For these reasons, we propose an approach to the problem which is based on classical Data Warehouse (DW) concepts, so that we can adopt a well established and studied data model, as well as the efficient tools and systems already developed for such a model. Our DW is aimed at defining, as basic elements of interest, not the observations of the moving objects, but rather their trajectories, so that we can study properties such as average speed, travelled distance, maximum acceleration, presence of distinct trajectories. We assume the granularity of the fact table given by a regular three-dimensional grid on the spatial and temporal dimensions, where the facts are the set of trajectories which intersect each cell of the grid, and the measures are properties related to that set.

One of the main issues to face is the efficient population of the Trajectory DW (TDW) from streams of trajectory observations, arriving in an unpredictable and unbounded way, with different rates. The challenge is to exploit a limited amount of buffer memory to store a few incoming past observations, in order to correctly reconstruct the various trajectories, and compute the needed measures for the base cells, reducing as much as possible the approximations.

The model of our TDW and the corresponding loading issues have been introduced in [Braz et al. 2007]. In this paper we discuss in detail the loading and computation of a complex aggregate measure, the presence, which can be defined as the number of distinct trajectories lying in a cell. Such a measure poses non trivial computational problems. On one hand, in the loading phase, only a bounded amount of memory can be used for analysing the input streams. This suffices for trajectory reconstruction, but in some cases we may still count an object, with multiple observations in the same cell, more than once. Hence the measure presence computed for base cells is in general an approximation of the exact value. A second approximation is introduced in the roll-up phase. In fact, since the roll-up function for the measure presence is, as distinct count, a holistic function, it cannot be computed using a finite number of auxiliary measures starting from sub-aggregates. Our proposal is based on an approximate, although very accurate, presence aggregate function, which algebraically combines a bounded amount of other sub-aggregate measures, stored in the base cells of the grid.

The paper will provide a thorough analysis of the above mentioned technique, focusing, in particular, on the errors introduced by the approximations. From our tests, our method turned out to yield an error which is sensibly smaller than the one of a recently proposed algorithm [Tao et al. 2004], based on sketches, which are a well known probabilistic counting method in database applications [Flajolet and Martin 1985].

We also realized a prototype of our TDW, in order to investigate its feasibility when using commodity warehousing tools, like the Oracle one. The first challenge is the ETL process of the DW: by exploiting a few memory, we have to transform
a stream of trajectory observations on-the-fly and aggregate them to update the base cells of our spatio-temporal cube. Using a Java application, performing the data transformation, and a specific loading method available in the Oracle suite, we were able to elaborate and load about 7000 observations per second. The other challenge is the efficient computation of our approximate aggregate function, used to roll-up the presence measure.

The rest of the paper is organised as follows. In Section 2 we present some related work whereas in Section 3 we discuss the issues related to the representation of trajectories. In Section 4 we recall our TDW model and in Section 5 we describe the loading phase of the measure presence and the aggregate functions supporting the roll-up operation. Then in Section 6 the approximation errors of the loading and roll-up phases are studied, both analytically and with the help of suitable tests. Section 7 focuses on our experience in designing a prototype of our TDW using Oracle. Finally, Section 8 draws some conclusion.

2. RELATED WORK

Research on spatial multidimensional data models is relatively recent. The pioneering work of Han et al. [Han et al. 1998] introduces a spatial data cube model which consists of both spatial and non-spatial dimensions and measures. Additionally, it analyses how OLAP operations can be performed in such a spatial data cube. Then, several authors have proposed data models for spatial data warehouses at a conceptual level (e.g., [Pedersen and Tryfona 2001; Damiani and Spaccapietra 2000]). Unfortunately, none of these approaches deal with objects moving continuously in time.

Research on moving objects has focused on modelling [Sistla et al. 1997; Güting et al. 2000] and on indexing such spatio-temporal data [Kollios et al. 1999; Pfoser et al. 2000], whereas the definition of TDWs is still an open problem. An issue that is gaining increasingly interest in the last years and that is relevant for the development of a TDW concerns the specification and efficient implementation of the operators for spatio-temporal aggregation. Nevertheless a standard set of operators (e.g., operators like Avg, Min, Max in SQL) has not yet been defined. A first comprehensive classification and formalisation of spatio-temporal aggregate functions is presented in [Lopez et al. 2005]. The aggregate operations are defined as functions that, applied to a collection of tuples, return a single value. To generate the collection of tuples to which these operations are applied, the authors distinguish three kinds of methods: group composition, partition composition and sliding window composition. On the other hand, in [Papadias et al. 2002; Tao and Papadias 2005] techniques for the computation of aggregate queries are developed based on the combined use of specialised indexes and materialisation of aggregate measures. For instance [Papadias et al. 2002] introduces the aRB-tree, which consists of two types of indexes: a host index and some measure indexes. The host index is an R-tree which manages the region extents, and associates with these regions an aggregate information over all the timestamps of interest. Additionally, for each entry of the host index, a B-tree containing the time-varying aggregate data is defined.

As shown in [Tao et al. 2004], most of these indexes, like aRB-trees, can suffer from the distinct counting problem, i.e., if an object remains in the query region for
several timestamps during the query interval, it will be counted multiple times in the result. In fact such indexes store, for each region, an aggregate measure and they forget the identifiers of the objects.

To cope with this issue, [Tao et al. 2004] proposes an approach which combines spatio-temporal indexes with sketches, a traditional approximate counting technique based on probabilistic counting [Flajolet and Martin 1985]. The index structure is similar to aRBTrees but differs from them in the querying algorithms since one can exploit the pruning power of the sketches to define some heuristics which allow to reduce the query time.

It is worth noticing that the above cited works on aggregation methods do not take into account the fact that the spatio-temporal observations belong to moving objects. Instead they treat such observations as unrelated points. For a data warehouse of trajectories, we will show that it is important to exploit such relationships existing among observations. For example, in order to compute the approximate number of distinct trajectories intersecting a given spatio-temporal area, in this paper we exploit the knowledge about how many objects move from one region to the close ones, thus improving the result of roll-up operations.

3. TRAJECTORY REPRESENTATION

In real-world applications the movements of a spatio-temporal object, i.e. its trajectory, is often given by means of a finite set of observations. This is a finite subset of points, called sampling, taken from the actual continuous trajectory. Figure 1(a) shows a trajectory of a moving object in a 2D space and a possible sampling, where each point is annotated with the corresponding time-stamp. It is reasonable to expect that observations are taken at irregular rates for each object, and that there is no temporal alignment between the observations of different objects.

Formally, let $\mathcal{T}_S$ be a stream of samplings of 2D trajectories $\mathcal{T}_S = \{T_i\}_{i \in \{1, \ldots, n\}}$. Each $T_i$ is the sampling for an object trajectory: $T_i = (ID_i, L_i)$, where $ID_i$ is the identifier associated with the object and $L_i = \{L_i^1, L_i^2, \ldots, L_i^{M_i}\}$ is a sequence of observations. Each observation $L_i^j = (t_i^j, x_i^j, y_i^j)$ represents the presence of an object at location $(x_i^j, y_i^j)$ and time $t_i^j$. The observations are temporally ordered, i.e., $t_i^j < t_i^{j+1}$.

In many situations, e.g., when one is interested in computing the cumulative number of trajectories in a given area, an (approximate) reconstruction of each trajectory from its sampling is needed. Among the several possible solutions, we will focus on local interpolation. According to this method, although there is not a global function describing the whole trajectory, objects are assumed to move between the observed points following some rule. For instance, a linear interpolation function models a straight movement with constant speed, while other polynomial interpolations can represent smooth changes of direction. The linear (local) interpolation, in particular, seems to be a quite standard approach to the problem (see, for example, [Pfoser et al. 2000]), and yields a good trade-off between flexibility and simplicity. Hence in this paper we will adopt this kind of interpolation. However, it is straightforward to use a different interpolation, based, for example, on additional information concerning the environment traversed by the moving objects.

Starting from the stream of samplings, $\mathcal{T}_S$, we want to investigate if DW tech-
4. FACTS, DIMENSIONS, MEASURES AND AGGREGATE FUNCTIONS

We define a DW model by means of a star schema, as simple and generic as possible. The facts of our TDW are the set of trajectories which intersect each cell of the grid. Some typical properties we want to describe, i.e., the measures, are the number of trajectories, their average, maximum/minimum speed, the covered distance. The dimensions of analysis consist of the spatial dimensions $X$, $Y$ ranging over spatial intervals, and the temporal dimension $T$ ranging over temporal intervals. We assume a regular three-dimensional grid obtained by discretizing the corresponding values of the dimensions and then we associate with them a set-grouping hierarchy. A partial order can thus be defined among groups of values, as illustrated in Figure 1(c) for a temporal or a spatial dimension. Note that in the TDW of Figure 1(b), the basic information we represent concerns the set of trajectories intersecting the spatio-temporal cell having for each $X$, $Y$, or $T$ dimensions as minimum granularity 30 units, measured in the corresponding unit of measure. The hierarchy is not explicitly represented (we refer the reader to Figure 4 for a more complete picture).

It is interesting to investigate whether some pre-computation must be carried out on the trajectory observations in order to feed the TDW. Some measures require little pre-computation, and can be updated in the TDW as soon as single observations of the various trajectories arrive, others need all the observations of a trajectory to be received, before updating the TDW. In [Braz et al. 2007] we...
classified the measures according to an increasing amount of pre-calculation effort. For instance, in order to compute the average, maximum/minimum speed and the covered distance by the trajectories intersecting a cell we need two consecutive observations since we have to infer the trajectory route through interpolation. On the other hand, the measure number of observations falling in a cell can be updated directly using each single observation.

We can build the spatial data cube [Gray et al. 1997] as the lattice of cuboids, where the lowest one (base cuboid) references all the dimensions at the primitive abstraction level, while the others are obtained by summarising on different subsets of the dimensions, and at different abstraction levels along the concept hierarchy. In order to denote a component of the base cuboid we will use the term base cell, while we will simply use cell for a component of a generic cuboid.

In order to summarise the information contained in the base cells, Gray et al. [Gray et al. 1997] categorise the aggregate functions into three classes based on the space complexity needed for computing a super-aggregate starting from a set of sub-aggregates already provided, e.g., the sub-aggregates associated with the base cells of the DW. The classes are the following:

1. **Distributive**. The super-aggregates can be computed from the sub-aggregates.
2. **Algebraic**. The super-aggregates can be computed from the sub-aggregates together with a finite set of auxiliary measures.
3. **Holistic**. The super-aggregates cannot be computed from sub-aggregates, not even using any finite number of auxiliary measures.

For example, the super-aggregates for the distance and the maximum/minimum speed are simple to compute because the corresponding aggregate functions are distributive. In fact once the base cells have been loaded with the exact measure, for the distance we can accumulate such measures by using the function sum whereas for maximum/minimum speed we can apply the function max/min. The super-aggregate for the average speed is algebraic: we need the pair of auxiliary measures \((distance, time)\) where distance is the distance covered by trajectories in the cell and time is the total time spent by trajectories in the cell. For a cell \(C\) arising as the union of adjacent cells, the cumulative function performs a component-wise addition, thus producing a pair \((distance_f, time_f)\). Then the average speed in \(C\) is given by \(distance_f / time_f\).

5. **THE MEASURE PRESENCE**

In this paper we focus on the measure presence which returns the number of distinct trajectories in a cell. As already remarked, a complication in dealing with such a measure is due to the distinct count problem and this has an impact on both the loading of the base cells and on the definition of aggregate functions able to support the roll-up operation.

5.1 **Aggregate functions**

According to the classification presented in Section 4, the aggregate function to compute the presence is holistic, i.e., it needs the base data to compute the result in all levels of dimensions. Such a kind of functions represents a big issue for DW.
technology, and, in particular, in our context, where the amount of data is huge and unbounded. A common solution consists in computing holistic functions in an approximate way.

We propose two alternative and non-holistic aggregate functions that approximate the exact value of the presence. These functions only need a small and constant memory size to maintain the information to be associated with each base cell of our TDW, from which we can start computing a super-aggregate.

The first aggregate function is distributive, i.e., the super-aggregate can be computed from the sub-aggregate, and it is called PresenceDistributive. We assume that the only measure associated with each base cell is the exact (or approximate) count of all the distinct trajectories intersecting the cell. Therefore, the super-aggregate corresponding to a roll-up operation is simply obtained by summing up all the measures associated with the cells. This is a common approach (exploited, e.g., in [Papadias et al. 2002]) to aggregate spatio-temporal data. However, our experiments will show that this aggregate function may produce a very inexact approximation of the effective presence, because the same trajectory might be counted multiple times. This is due to the fact that in the base cell we do not have enough information to perform a distinct count when rolling-up.

The second aggregate function is algebraic, i.e., the super-aggregate can be computed from the sub-aggregate together with a finite set of auxiliary measures, and it is called PresenceAlgebraic. In this case each base cell stores a tuple of measures. Besides the exact (or approximate) count of all the distinct trajectories intersecting the cell, the tuple includes other measures which are used when we compute the super-aggregate. The additional measures are helpful to correct the errors, caused by the duplicates, introduced by the function PresenceDistributive.

More formally, let \( C_{x,y,t} \) be a base cell of our cuboid, where \( x, y, \) and \( t \) identify intervals of the form \([l, u]\), in which the spatial and temporal dimensions are partitioned. The tuple associated with the cell consists of \( C_{x,y,t}.presence, C_{x,y,t}.crossX, C_{x,y,t}.crossY, \) and \( C_{x,y,t}.crossT \).

\( C_{x,y,t}.presence \) is the number of distinct trajectories intersecting the cell.

\( C_{x,y,t}.crossX \) is the number of distinct trajectories crossing the spatial border between \( C_{x,y,t} \) and \( C_{x+1,y,t} \).

\( C_{x,y,t}.crossY \) is the number of distinct trajectories crossing the spatial border between \( C_{x,y,t} \) and \( C_{x,y+1,t} \).

\( C_{x,y,t}.crossT \) is the number of distinct trajectories crossing the temporal border between \( C_{x,y,t} \) and \( C_{x,y,t+1} \).

Let \( C_{x',y',t'} \) be a cell consisting of the union of two adjacent cells with respect to a given dimension, namely \( C_{x',y',t'} = C_{x,y,t} \cup C_{x+1,y,t} \). In order to compute the super-aggregate corresponding to \( C_{x',y',t'} \), we proceed as follows:

\[
\text{PresenceAlgebraic}(C_{x,y,t} \cup C_{x+1,y,t}) = C_{x',y',t'}.presence = \\
C_{x,y,t}.presence + C_{x+1,y,t}.presence - C_{x,y,t}.crossX
\]  

The other measures associated with \( C_{x',y',t'} \) can be computed in this way:

\[
C_{x',y',t'}.crossX = C_{x+1,y,t}.crossX
\]

\[
C_{x',y',t'}.crossY = C_{x,y,t}.crossY + C_{x+1,y,t}.crossY
\]

\[ C_{x',y',t'}.crossT = C_{x,y,t}.crossT + C_{x+1,y,t}.crossT \]  \hspace{1cm} (4)

Equation (1) can be thought of as an application of the well known Inclusion/Exclusion principle: \( |A \cup B| = |A| + |B| - |A \cap B| \) for all sets \( A \), \( B \). Suppose that the elements included in the sets \( A \) and \( B \) are just the distinct trajectories intersecting the cells \( C_{x,y,t} \) and \( C_{x+1,y,t} \), respectively. Hence, their cardinalities \( |A| \) and \( |B| \) exactly correspond to \( C_{x,y,t}.presence \) and \( C_{x+1,y,t}.presence \). Then \( C_{x,y,t}.crossX \) is intended to approximate \( |A \cap B| \), but, notice that, unfortunately, in some cases \( C_{x,y,t}.crossX \) is not equal to \( |A \cap B| \), and this may introduce errors in the values returned by \( Presence_{Algebraic} \). Figure 2(a) shows a trajectory that will be correctly counted, since it crosses the border between the two cells to be rolled-up. Conversely, Figure 2(b) shows a very agile and fast trajectory, which will be counted twice during the roll-up, since it is not accounted in \( C_{x,y,t}.crossX \), even if it should appear in \( |A \cap B| \). In fact, the trajectory intersects both \( C_{x,y,t} \) and \( C_{x+1,y,t} \), but does not cross the \( X \) border between the two cells.

In order to understand Equations 2–4, notice that the right \( X \) border of the new cell \( C_{x',y',t'} \) is the corresponding \( X \) border of the rightmost cell \( C_{x+1,y,t} \), hence the number of distinct trajectories crossing the \( X \) border of \( C_{x',y',t'} \) is \( C_{x+1,y,t}.crossX \) (Equation 2). Instead, the top \( Y \) border of \( C_{x',y',t'} \) is the union of the corresponding borders of the component cells \( C_{x,y,t} \) and \( C_{x+1,y,t} \). This is why in the Equation 3 we take the sum of \( C_{x,y,t}.crossY \) and \( C_{x+1,y,t}.crossY \). Similar considerations apply to Equation 4.

5.2 Loading phase

In this section we deal with the problem of feeding the DW, i.e., the base cells of our base cuboid, with suitable sub-aggregate measures, from which the aggregate functions compute the super-aggregates.

We recall that trajectory observations arrive in streams at different rates, in an unpredictable and unbounded way. In order to limit the amount of buffer memory needed, it is essential to store information only about active, i.e., not ended trajectories. In our simple model of trajectory sampling, since we do not have an end-mark associated with the last observation of a given trajectory, the system module that is responsible for feeding data decides to consider a trajectory as ended when for a long time interval no further observation for the object has been received.

In the following we present two options to compute the sub-aggregates stored in each base cell \( C_{x,y,t} \), namely \( C_{x,y,t}.presence \), \( C_{x,y,t}.crossX \), \( C_{x,y,t}.crossY \), and
$C_{x,y,t} \cdot \text{cross} T$, which are used by the aggregate functions $\text{Presence}_{Distributive}$ and $\text{Presence}_{Algebraic}$.

1. **Single observations.** Considering a single observation at a time, without buffering any past observation, we can only update/increment the measure $C_{x,y,t} \cdot \text{presence}$. As a consequence, we cannot compute $\text{Presence}_{Algebraic}$, since $C_{x,y,t} \cdot \text{cross} X$, $C_{x,y,t} \cdot \text{cross} Y$ and $C_{x,y,t} \cdot \text{cross} T$ are not available.

2. **Pairs of observations.** We consider a pair of observations consisting of the currently received observation $L^i_t$ of trajectory $T_i$, with the previously buffered $L^{i-1}_t$. Using this pair of points, we can linearly interpolate the trajectory, thus registering further presences for the cells only traversed by $T_i$. Moreover, if we store in the buffer not only $L^{i-1}_t$, but also the last base cell $C_{x,y,t}$ that was modified on the basis of trajectory $T_i$, we can reduce the number of duplicates by simply avoiding updating the measure when $L^i_t$ falls into the same $C_{x,y,t}$. It is worth noting that since this method only exploits a very small buffer, it is not able to remove all the possible duplicates from the stored presence measures. Consider, for example, three observations of the same trajectory, all occurring within the same base time interval. If the first and the third points fall into the same cell $C_{x,y,t}$, but the second one falls outside (e.g., into $C_{x,y+1,t}$), this method will store a duplicate count in $C_{x,y,t} \cdot \text{presence}$ when the third observation is encountered.

Finally, by exploiting linear interpolation, we can also identify the cross points of each base cell, and accordingly update the various sub-aggregates $C_{x,y,t} \cdot \text{cross} X$, $C_{x,y,t} \cdot \text{cross} Y$, and $C_{x,y,t} \cdot \text{cross} T$.

5.3 Accuracy of the approximate aggregate function

Let us give an intuitive idea of the errors introduced by the aggregate function $\text{Presence}_{Algebraic}$, starting from the measures loaded in the base cells by the method **Pairs of observations.** Notice that the overall error can be obtained as the sum of the errors computed for each single trajectory in isolation.

Let us first consider the simplest case: a trajectory is a line segment $l$. In this case, no roll-up errors are introduced in the computation of presence. In fact, let $C_{x',y',t'}$ be the union of two adjacent cells, $C_{x,y,t}$ and $C_{x+1,y,t}$, i.e., $C_{x',y',t'} = C_{x,y,t} \cup C_{x+1,y,t}$, and $l$ intersect $C_{x',y',t'}$. Then by definition:

$$\text{Presence}_{Algebraic}(C_{x,y,t} \cup C_{x+1,y,t}) = C_{x,y,t} \cdot \text{presence} + C_{x+1,y,t} \cdot \text{presence} - C_{x,y,t} \cdot \text{cross} X$$

If $l$ intersects the $X$ border, then correctly $\text{Presence}_{Algebraic}(C_{x',y',t'}) = 1 + 1 - 1$. Otherwise, $C_{x,y,t} \cdot \text{cross} X = 0$, but $l$ intersects only one of the two cells. Hence $\text{Presence}_{Algebraic}(C_{x',y',t'}) = 1 + 0 - 0$ or $\text{Presence}_{Algebraic}(C_{x',y',t'}) = 0 + 1 - 0$.

This result can be extended to a trajectory which is composed of a set of line segments whose slopes are in the same octant of the three-dimensional coordinate system. Let us call uni-octant sequence a maximal sequence of line segments whose slopes are in the same octant. Clearly, every trajectory can be uniquely decomposed in uni-octant sequences and, in the worst case, the error introduced by the aggregate function $\text{Presence}_{Algebraic}$ will be the number of uni-octant sequences composing a trajectory.
6. EVALUATING APPROXIMATE SPATIO-TEMPORAL AGGREGATES

In this section we evaluate our method to approximate the measure presence and we are also interested in comparing it with a distributive approximation recently proposed [Tao et al. 2004]. The method is based on sketches, a traditional technique based on probabilistic counting, and in particular on the FM algorithm devised by Flajolet and Martin [Flajolet and Martin 1985].

6.1 FM sketches

FM is a simple, bitmap-based algorithm to estimate the number of distinct items. In particular, each entry in the sketch is a bitmap of length \( r = \log UB \), where \( UB \) is an upper bound on the number of distinct items. FM requires a hash function \( h \) which takes as input an object ID \( i \) (in our case a trajectory identifier), and outputs a pseudo-random integer \( h(i) \) with a geometric distribution, that is, \( \text{Prob}[h(i) = v] = 2^{-v} \) for \( v \geq 1 \). Indeed, \( h \) is obtained by combining a uniformly distributed hash function \( h' \), and a function \( \rho \) that selects the least significant 1-bit in the binary representation of \( h'(i) \). The hash function is used to update the \( r \)-bit sketch, initially set to 0. For every object \( i \) (e.g., for every trajectory observation in our stream), FM thus sets the \( h(i) \)-th bit. In the most naive and simple formulation, after processing all objects, FM finds the position of the leftmost bit of the sketch that is still equal to 0. If \( k \) is this position, then it can be shown that the overall object count \( n \) can be approximated with \( 1.29 \times 2^k \).

Unfortunately, this estimation may entail large errors in the count approximation. The workaround proposed in [Flajolet and Martin 1985] is the adoption of \( m \) sketches, which are all updated by exploiting a different and independent hash function. Let \( k_1, k_2, \ldots, k_m \) be the positions of the leftmost 0-bit in the \( m \) sketches. The new estimate of \( n \) is \( 1.29 \times 2^{k_a} \), where \( k_a \) is the mean of the various \( k_i \) values, i.e. \( k_a = (1/m) \sum_{i=1}^{m} k_i \). This method reduces the standard error of the estimation, even if it increases the expected processing cost for object insertion. The final method that reduces the expected insertion cost back to \( O(1) \) is Probabilistic Counting with Stochastic Averaging (PCSA). The trick is to randomly select, for each object to insert, one of the \( m \) sketches. Each sketch thus becomes responsible for approximately \( n/m \) (distinct) objects.

An important feature of FM sketches is that they can be merged in a distributive way. Suppose that each sketch is updated on the basis of a different set of objects. We can merge a pair of sketches together, in order to get a summary of the number of distinct items seen over the union of both sets of items, by simply taking the bitwise-OR of the corresponding bitmaps. We can find a simple application of these FM sketches in our TDW. First, we can exploit a small logarithmic space for each base cell, used to store a sketch that approximates the distinct count of trajectories seen in the corresponding spatio-temporal region. The distributive nature of the sketches can then be used to derive the counts at upper levels of granularities, by simply OR-ing the sketches at the lowest granularities.

6.2 Experimental Setup

**Datasets.** In our experiments we have used several different datasets. Most of them are synthetic ones, generated by the traffic simulator described in [Brinkhoff
2002], but also some real-world sets of trajectories, presented in [Frentzos et al.
2005], were used.

Due to space limitations, we present only the results produced for one of the
synthetic datasets and for one real dataset. The synthetic dataset was generated
using a map of the San Joaquin road network (US Census code 06077), and contains
the trajectories of 10000 distinct objects monitored for 50 time units, with an
average trajectory length of 170k space units. The real-world dataset contains the
trajectories of 145 school buses. The average number of sample points per trajectory
is 455, and, differently from the case of synthetic data, not all points are equally
distant in time. Since the setting does not suggest a specific granularity, we define as
a standard base granularity for each dimension the average difference of coordinate
values between two consecutive points in a trajectory. We will specify granularities
as multiples of this base granularity \( g \). For the synthetic dataset \( g = (1, 2000, 2000) \)
and for the school buses one \( g = (2, 100, 100) \), where the granularities of the three
dimensions are in the order: \( t, x, y \).

**Accuracy assessment.** Vitter et al. [Vitter et al. 1998] proposed several error
measures. We selected the one based on absolute error, which appears to be the
most suited to represent the error on data that will be summed. Instead of the
\( \infty - \text{norm} \), however, we preferred to use the \( 1-\text{norm} \) as a normalisation term, since
it is more restrictive and less sensible to outliers.

The formula we used to compute the normalised absolute error for all the OLAP
queries \( q \) in a set \( Q \) is thus the following:

\[
\text{Error} = \frac{\| \tilde{M} - M \|_1}{\| M \|_1} = \frac{\sum_{q \in Q} |\tilde{M}_q - M_q|}{\sum_{q \in Q} M_q}
\]  

(5)

where \( \tilde{M}_q \) is the approximate measure computed for query \( q \), while \( M_q \) is its exact
value. The various queries \( q \) may either refer to base cells, or to cells at a coarser
level of granularity. In particular, we used this error measure for evaluating the
*loading errors* at the level of base cells, and the *roll-up errors* at the level of coarser
cells. In all these cases, we always applied Equation (5) to evaluate the errors
tained by a set \( Q \) of queries at uniform granularity.

6.3 Experimental Evaluation

First we evaluate the errors in the loading phase of our TDW. During this phase,
the reconstruction of the trajectories takes place. Figures 3.(a) and 3.(b) show the
accuracy of the loaded (sub-aggregate) presence, for different granularities of the
base cells. In particular, the two figures illustrate the normalised absolute error as
a function of the size of base cells for the two datasets.

Let us focus on the curves labelled *single observations*. In this case we simply fed
the TDW with each single incoming trajectory observation, without introducing
any additional interpolated point. For small granularities, errors are mainly due to
the fact that some cells are traversed by trajectories, but no observation falls into
them. On the other hand, for very coarse granularities, the main source of errors
is the duplicate observations of the same trajectory within the same cell. There
exists an intermediate granularity (around 2 \( g \) for these particular datasets) which
represents the best trade-off between the two types of errors, thus producing a small error.

Then consider the curves labelled as *observation pairs*. In this case the presence (sub-aggregate) measure of the base cells of the TDW is loaded by exploiting a buffer, which stores information about active trajectories. For each of these trajectories, the buffer only stores the last received observation. This information is used to reduce the errors with respect to the previous feeding method, by adding interpolated points, and also avoiding to record duplicates in the base cells. The resulting normalised absolute error is close to 0 for small granularities, and it remains considerably small for larger values. For very large granularities, a buffer that only stores information about the last previously encountered observation does not suffice to avoid recording duplicates in the base cells.

The other curves in Figure 3(a) and 3(b) represent the error introduced in the base cell by approximating the presence (sub-aggregate) measure by using sketches. It is worth noting that also to load the sketch-base cells, we added further interpolated observations, by using the buffer during the loading phase. Although the error slightly decreases when a larger number of sketches, \( m \), is used for each cell, it remains considerably larger than the curve labelled as *observation pairs*, except for coarse granularities and a large number of sketches \( (m = 40) \).

The second set of tests, whose results are reported in Figures 3(c) and 3(d), regards the errors introduced by rolling-up our cube, starting from the base cells at granularity \( g \). The base granularity \( g = 1 \) used for these tests was the same as the one in Figures 3(a-b), while the method used to load the base cells was *observation pairs*. Therefore the normalised absolute error at the various base cells was close to 0. In all the roll-up tests, the error is averaged by submitting many queries at a given granularity \( g' > g, g' = n \cdot g \). Each query computes the presence measure associated with a distinct coarse-grain cell, obtained by aggregating different adjacent base cells. As an example of query, eight adjacent base cells are aggregated to make one cell of granularity \( 2 \cdot g \).

As shown by the corresponding curves, the *distributive aggregate* function *(sum)* quickly reaches very large errors as the roll-up granularity increases. This is due to the duplicated counting of the same trajectories, when they are already counted by multiple sub-aggregates. Conversely, we obtained very accurate results with our *algebraic* method, with an error always smaller than 10%, for most granularities (except for very coarse ones and 40 sketches). Finally, the *distributive* function *(or)* used to aggregate sketches produce much larger errors than our method, even because such large errors are already present in the sketches stored in the base cells. However, a nice property of sketches is that the cumulative errors do not increase, and, are actually often reduced when rolling-up, but sketches also exploit much more memory per cell. In case of \( m = 40 \), the memory usage of the sketch-based method is one order of magnitude larger than our algorithm. Thus it is remarkable that our method achieves a better accuracy in most cases, in spite of this unfavourable bias.
Figure 3. Errors on different datasets during the load and roll-up phases as a function of the granularity. Granularity is expressed as a multiple of the minimal granularity $g$ for load, and of the base granularity $g$ for roll-up (note logarithmic scale for both axes).

7. PROTOTYPE OF OUR TDW USING ORACLE

In this section we will focus on the implementation of our TDW. In particular we will discuss the most challenging issues with respect to a traditional DW: namely, the preprocessing phase to transform data to be loaded, and the aggregation functions, necessary to support OLAP operations. As described in Section 5.2, the set of streaming observations are not enough to correctly compute the measures, e.g., we have to generate new points through interpolations. Moreover, the common aggregation functions provided by DWs are sum, max/min, average. Instead, in our case, we have to implement a more complex algebraic function, in order to approximate the holistic function needed to compute the measure presence in a roll-up query.

The TDW has been implemented and tested by using the DW tools of the Oracle DBMS suite. We experimented different solutions to tackle the mentioned problems: the most efficient ones use ad-hoc constructs of Oracle, but, in order to
have a portable software, we also tested alternative implementations which employ standard SQL.

Let us first describe the data model for our TDW, illustrated in Figure 4. As presented in Section 4, we define three dimension tables, two for the spatial dimension, DimX and DimY, ranging over equally sized spatial intervals, and one for the temporal dimension, DimT, ranging over equally sized temporal intervals. For each base interval \( I \) in each dimension, the corresponding table contains its identifier and some attributes \texttt{level1}, \texttt{level2}, \ldots, modelling a set-grouping hierarchy, defined by the user. In particular, attribute \texttt{levelj} assumes as value the identifier of the interval at the \( j \)-th level of the hierarchy including \( I \). Referring to the simple hierarchy shown in Figure 1(c), if \( I = "[60,90)" \), the corresponding attribute \texttt{level1} contains the father of \( I \) in the hierarchy, i.e. "[60,120)". The \texttt{Fact_Table} has the conjunction of the foreign keys of the dimension tables, \((\texttt{dimX}._\texttt{id}, \texttt{dimY}._\texttt{id}, \texttt{dimT}._\texttt{id})\) as primary key, and \texttt{presence}, \texttt{crossX}, \texttt{crossY} and \texttt{crossT} as measures. Since in this paper we focus on the measure \texttt{Presence}, the other measures, such as \texttt{distance} and \texttt{average speed}, are not reported in the figure.

As already remarked in Section 4, the fact table contains only aggregate information about trajectories (i.e., the number of distinct trajectories in a cell and the number of distinct trajectories crossing the \( X, Y \) and \( T \) border of the cell). The identifiers of the trajectories and their observations are not kept in the TDW. These data can be stored into a Moving Object Database (MOD) [Güting and Schneider 2005], which can be useful also to extract higher level knowledge that in turn will feed the TDW. In this context, the features of a spatial Database, like Spatial Oracle, could be of great help for storing, querying and indexing trajectories.
7.1 The ETL process

In our prototype we assume that data arrive in streams of observations of the form $(id, t, x, y)$, ordered by time, and we extract an observation at a time from this ordered stream.

In the transformation phase we prepare data for loading. In particular, we transform the various observations into single contributions to the measures to be stored in the various cells. Then such contributions must be grouped-by, in order to load the actual final measures into the DW cells.

The main operations of our transformation phase consist of:

(i) transforming each observation $(id, t, x, y)$ into a cell tuple, containing the key of the corresponding base cell into which the observation falls into, and its contribution to the cell measures;

(ii) determining, through linear interpolation, the base cells traversed by a trajectory where no observation falls, and creating the relative cell tuples;

(iii) avoiding the production of duplicate contributions to the same cell for the same trajectory. Therefore, if the current observation of a trajectory falls into the same base cell of the last received point for the same trajectory, we avoid producing a cell tuple.

Finally, during the loading phase of the cell tuples obtained by the previous procedure, another transformation operation is necessary to obtain the measures to store in a cell:

(iv) aggregating the cell tuples according to the cell key, and summing up the corresponding measures, thus resulting into a single tuple for each cell traversed by at least one trajectory. This tuple corresponds to the base cell to insert into the TDW.

It is worth remarking that, in order to obtain a correct update of our TDW, this loading procedure must occur at fixed times. In particular, if $\Delta T$ is the temporal granularity of the base cells of the TDW, we can load the tuple cells that have been so far produced when all the observations that fall in a given $\Delta T$ interval (or a multiple of $\Delta T$) have been received and transformed. All the tuple cells grouped-by and loaded in the TDW have to refer to this $\Delta T$ interval (or a multiple of $\Delta T$).

Let us give an example of the result of the transformation process. Consider the set of observations depicted in Figure 1(a). Then, we reconstruct the whole trajectory by using linear interpolation, as discussed in Section 3. Figure 5 shows the reconstructed trajectory together with the discretized 2D space and the points where the interpolated trajectory intersects the spatio-temporal cells. The transformation phase generates the cell tuples described in Table I where $(dimX\_id, dimY\_id, dimT\_id)$ – i.e., the conjunction of the foreign keys for the $X$, $Y$, and $T$ dimensions – is the primary key for the fact table of our TDW\(^1\). The observation with

\(^1\)For the sake of clarity, we use the corresponding intervals instead of the numeric values of such
Figure 5. Linear interpolation of a 2D trajectory. The white points correspond to intersections between the interpolated trajectory and spatial/temporal borders.

Table I. The cell tuples obtained by the transformation phase. For the sake of clarity, we also report the timestamps (labels) of the original (or interpolated) points shown in Figure 5 that originated each tuple. Note that some points, like the one labelled by 23, may not correspond to any tuple: this avoid counting duplicates in the DW.

<table>
<thead>
<tr>
<th>Tstamp</th>
<th>dimX_id</th>
<th>dimY_id</th>
<th>dimT_id</th>
<th>presence</th>
<th>crossX</th>
<th>crossY</th>
<th>crossT</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>(30,60)</td>
<td>(30,60)</td>
<td>(0,30)</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>30</td>
<td>(30,60)</td>
<td>(30,60)</td>
<td>(0,30)</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>32</td>
<td>(30,60)</td>
<td>(30,60)</td>
<td>(30,60)</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>60</td>
<td>(60,90)</td>
<td>(30,60)</td>
<td>(30,60)</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>65</td>
<td>(60,90)</td>
<td>(30,60)</td>
<td>(60,90)</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>67</td>
<td>(60,90)</td>
<td>(30,60)</td>
<td>(60,90)</td>
<td>0</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>70</td>
<td>(60,90)</td>
<td>(60,90)</td>
<td>(60,90)</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>73</td>
<td>(90,120)</td>
<td>(60,90)</td>
<td>(60,90)</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>75</td>
<td>(90,120)</td>
<td>(90,120)</td>
<td>(60,90)</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>81</td>
<td>(90,120)</td>
<td>(90,120)</td>
<td>(60,90)</td>
<td>0</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>87</td>
<td>(120,150)</td>
<td>(90,120)</td>
<td>(60,90)</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

timestamp 10 generates the first tuple, whereas no tuple is created by the second observation (timestamp 23) since it is inside the same cell of the first point. The second, the third and the fourth tuples are inserted because the interpolated trajectory between the observations with timestamp 23 and 65 crosses a T border, a X border and a T border, respectively, of spatio-temporal base cells. Note that, by construction, in each tuple cell the measures can assume only the values 0 or 1, since we process each single trajectory in isolation. Therefore we can have different cell tuples having the same value for the primary key. Only when we aggregate

foreign keys.

Table II. Performances of the loading techniques

<table>
<thead>
<tr>
<th>Method</th>
<th>rows per second</th>
</tr>
</thead>
<tbody>
<tr>
<td>SQL Loader</td>
<td>26695</td>
</tr>
<tr>
<td>INSERT</td>
<td>1446</td>
</tr>
<tr>
<td>INSERT Batch</td>
<td>4135</td>
</tr>
<tr>
<td>EXTERNAL Table</td>
<td>30884</td>
</tr>
<tr>
<td>MERGE INTO</td>
<td>4834</td>
</tr>
</tbody>
</table>

these cells according to \((dimX_id, dimY_id, dimT_id)\), we obtain a unique tuple containing the actual values to be loaded into the fact table.

7.1.0.1 Transformation/Loading Tools. In order to feed the DW, we analysed different techniques:

—SQL Loader: this is the standard loading tool for Oracle. It is a command line tool that can refresh the data in the DBMS. A configuration file describes the format of data in the input file and how these data are loaded into a DBMS table.

—SQL INSERT: this is a common way to process and load data by using an external ad-hoc application, which can exploit an API for Database Connectivity (e.g., JDBC for Java applications) to load data. It can perform whichever operations, and it has complete control on the operations to perform on importing data.

—batch SQL INSERT: as the above, but optimised for large number of insertions (the insertions are executed in blocks for optimising the performance).

—EXTERNAL Table: this is a way for accessing external data from within the DBMS, which can operate on external text files in the same way as DBMS tables. To use EXTERNAL Table as a transformation/loading tool, consider that we can perform any SQL transformation on external tables, and copy the results of these SQL queries into the DBMS internal tables.

—MERGE INTO: this is the Oracle tool for updating a DBMS table from other tables, or even EXTERNAL ones. Each single row of the DBMS table can be updated or inserted, depending on a Boolean condition.

We compared the above solutions to load the TDW, using an enlarged version (151k trajectories) of the Tiger dataset described in Section 6.2. The results obtained, expressed in terms of maximum throughput, are reported in Table II. It turns out that the fastest way of importing data is through EXTERNAL Tables.

7.1.0.2 Implementing the ETL process. Even if EXTERNAL Tables allow us also to make whichever SQL manipulations on the data during the import, for performance reasons, we decided to implement the most complex transformations, such as interpolation and the computation of the contribution of each observation to the measures, by using a Java application. This Java application is also in charge of extracting the observations from the incoming stream.

In summary, our ETL process works as follows:

—a Java application reads, one at a time, the trajectory observations from the temporally-ordered stream of observations. For each observation, it retrieves from an internal buffer the previous observation of the same trajectory, and
checks whether the current observation falls in the same base cell of the stored one. If it belongs to a different cell (or, it is the first read observation of a given trajectory), our application determines its contribution to the cell measures, and creates a new cell tuple (like the ones shown in Table I) that is appended to a temporary file. Moreover, it computes other possible cells traversed by the trajectory via interpolation, also appending the corresponding cell tuples to the temporary file (operations (i)–(ii)–(iii) of the transformation stage).

As mentioned above, to simplify the following phase and avoid errors while loading the DW, multiple temporary files are created/updated by our application. More precisely, all the cell tuples that fall into a given interval $\Delta T$ of the time dimension of our DW, are appended to the same file.

—Each completed temporary file can be viewed by the DBMS as an External Table. The cell tuples are thus aggregated via a GROUP BY command, according to the primary key of the cell $(\text{dimX\textunderscore id}, \text{dimY\textunderscore id}, \text{dimT\textunderscore id})$, and the measures are simply summed up. In this way, we prepare a single/distinct tuple for each cell (operation (iv) of the transformation stage). The resulting tuples can be simply inserted into the fact table of the DW, and, finally, the temporary file can be thrown away.

We chose this distribution of the ETL task between the Java application and the DBMS also to keep the amount of memory used by the Java application low: we only need a buffer containing, for each trajectory, the last processed observation, whereas the cell tuples with the measures are directly appended to a temporary file. In this way, we do not need to maintain in the main memory all the contributions to the measures to store in the various base cells of our TDW.

We tested the ETL implementation we have discussed so far, by using the Tiger dataset. It took 27 minutes to load all the 151000 trajectories (10867722 observations) of the dataset. We also tested other ETL implementations, based on either SQL INSERT or MERGE INTO. They required about 6 hours to complete.

7.2 Aggregation

The challenging problem concerning aggregation is due to the measure presence. As discussed in Section 5.1, we compute the value of this measure, for the higher levels of the hierarchy, by using two approximate functions, one distributive and the other one algebraic. The implementation of the distributive function is based on the aggregate standard function $\text{SUM}$ applied to the measure presence. Then, by using the CUBE operation we can summarise between multiple dimensions. For instance, given $\text{level1}$ as the layer of the hierarchy on dimension $X$, $\text{level1}$ as the layer of the hierarchy on dimension $Y$ and $\text{level1}$ as the layer of the hierarchy on dimension $T$, we can obtain an aggregate information for all the combinations of these three levels, i.e., $2^3$ groupings. The SQL query is the following:

```sql
SELECT dt.level1, dx.level1, dy.level1, SUM(presence) FROM Fact_Table ft, DimT dt, DimX dx, DimY dy WHERE ft.dimT_id = dt.dimT_id AND ft.dimX_id = dx.dimX_id AND ft.dimY_id = dy.dimY_id GROUP BY CUBE (dt.level1, dx.level1, dy.level1);
```

The implementation of the algebraic function is more complex since it requires the combination of several aggregate functions and the use of a non-standard SQL operation. In fact, suppose that we want to roll-up \( n \) adjacent cells along the \( X \) dimension. We have to implement the following operation:

\[
\sum_{i=0}^{n-1} C_{x+i,y,t}.\text{presence} - \sum_{i=0}^{n-2} C_{x+i,y,t}.\text{crossX}
\]

The difficulty consists in avoiding to subtract the \text{crossX} of the last cell of this sequence of cells. In fact, considering the sequence of cells as a single cuboid, the presence in it is given by the sum of the presence in each cell corrected by removing the flow (i.e., the number of crossing trajectories) at the \text{internal} \( X \) borders. In Oracle, the LAST function is available: it operates on a set of values from a set of rows that rank as the last with respect to a given sorting specification. As for the above query, we use the CUBE operation to obtain all the possible combinations.

```sql
SELECT dt.levelI, dx.levelH, dy.levelJ,
SUM(presence) - (SUM(crossX) + SUM(crossY) + SUM(crossT)) + (SUM(crossT))
KEEP (DENSE_RANK LAST ORDER BY dimT_id ASC) + SUM(crossX)
KEEP (DENSE_RANK LAST ORDER BY dimX_id ASC) + SUM(crossY)
KEEP (DENSE_RANK LAST ORDER BY dimY_id ASC)) FROM Fact_Table ft, DimT dt,
DimX dx, DimY dy WHERE ft.dimT_id = dt.dimT_id AND ft.dimX_id=
dx.dimX_id AND
ft.dimY_id=dy.dimY_id
GROUP BY CUBE (dt.levelI, dx.levelH, dy.levelJ);
```

The construct KEEP...DENSE_RANK LAST indicates that the aggregation will be executed over only those rows having \text{maximum} dense rank. This construct simplifies the computation of the algebraic function for the presence. In fact, a solution without LAST would require several nested queries combined with the use of outer joins.

As far as the performance is concerned, the first query has an execution time less than the second one, as expected. However, the difference is around 30%.

8. CONCLUSIONS

In this paper we discussed some relevant issues arising in the design of a DW devised to store measures regarding trajectories. In particular, we focused on the research problems related to storing and aggregating the holistic measure presence. We contributed a novel way to compute such a measure in an approximate, but nevertheless very accurate way. We also compared our approximate function with the method proposed in [Tao et al. 2004], based on sketches, showing, with the help of various experiments, that in our case the error is in general much smaller.

We also analysed the design issues deriving from the implementation of an Oracle prototype of our TDW. We have investigated in depth its feasibility, and, in particular, the challenges deriving from the ETL process of the DW, which has to start from a stream of trajectory observations, possibly produced at different rates, and arriving in an unpredictable and unbounded way. In addition, we have analysed the issue related to the computation of our approximate aggregate functions, used to roll-up on the presence measure.
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REFERENCES


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