Week 07 Lectures

Signature-based Selection

Indexing with Signatures

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Signature-based indexing:

- designed for *pmr* queries (conjunction of equalities)
- does not try to achieve better than O(n) performance
- attempts to provide an "efficient" linear scan

Each tuple is associated with a signature

- a compact (lossy) descriptor for the tuple
- formed by combining information from multiple attributes
- stored in a signature file, parallel to data file

Instead of scanning/testing tuples, do pre-filtering via signatures.

... Indexing with Signatures

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File organisation for signature indexing (two files)

Signature File		Data File
[0]	[0]	
[1]	[1]	
[2]	[2]	
[3]	[3]	
[4]	[4]	
[5]	[5]	
[6]	[6]	
[7]	[7]	
[8]	[8]	

One signature slot per tuple slot; unused signature slots are zeroed.

Signatures do not determine record placement ⇒ can use with other indexing.

Signatures

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A signature "summarises" the data from one tuple

A tuple consists of *n* attribute values $A_1 ... A_n$

A codeword $cw(A_i)$ is

- a bit-string, m bits long, where k bits are set to 1 $(k \ll m)$
- derived from the value of a single attribute A_i

A tuple descriptor (signature) is built by combining $cw(A_i)$, i=1..n

- combine by *overlaying* codewords (bitwise-OR)
- aim to have roughly half of the bits set to 1

Generating Codewords

```
bits codeword(char *attr_value, int m, int k)
   int nbits = 0;
                     // count of set bits
   bits cword = 0;
                     // assuming m <= 32 bits</pre>
   srandom(hash(attr_value));
   while (nbits < k) {
      int i = random() % m;
      if (((1 << i) \& cword) == 0) {
         cword = (1 << i);
         nbits++;
      }
   }
                 // m-bits with k 1-bits and m-k 0-bits
   return cword;
}
```

Superimposed Codewords (SIMC)

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In a superimposed codewords (simc) indexing scheme

a tuple descriptor is formed by overlaying attribute codewords

A tuple descriptor desc(r) is

- a bit-string, m bits long, where $j \le nk$ bits are set to 1
- $desc(r) = cw(A_1)$ OR $cw(A_2)$ OR ... OR $cw(A_n)$

Method (assuming all *n* attributes are used in descriptor):

```
bits desc = 0
for (i = 1; i <= n; i++) {
   bits cw = codeword(A[i])
   desc = desc | cw
}</pre>
```

SIMC Example

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Consider the following tuple (from bank deposit database)

Branch	AcctNo	Name	Amount
Perryridge	102	Hayes	400

It has the following codewords/descriptor (for m = 12, k = 2)

```
      Ai
      cw(Ai)

      Perryridge
      010000000001

      102
      00000000001

      Hayes
      00001000100

      400
      000010000100

      desc(r)
      010011000111
```

SIMC Queries

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To answer query q in SIMC

• first generate a query descriptor desc(q)

then use the query descriptor to search the signature file

desc(q) is formed by OR of codewords for known attributes.

```
E.g. consider the query (Perryridge, ?, ?, ?).
```

... SIMC Queries

Once we have a query descriptor, we search the signature file:

Example SIMC Query

Consider the query and the example database:

Signature	Deposit Record
010000000001	(Perryridge,?,?,?)
100101001001	(Brighton,217,Green,750)
010011000111	(Perryridge,102,Hayes,400)
101001001001	(Downtown,101,Johnshon,512)
101100000011	(Mianus,215,Smith,700)
010101010101	(Clearview,117,Throggs,295)
100101010011	(Redwood,222,Lindsay,695)

Gives two matches: one true match, one false match.

SIMC Parameters

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False match probablity p_F = likelihood of a false match

How to reduce likelihood of false matches?

- use different hash function for each attribute $(h_i \text{ for } A_i)$
- increase descriptor size (m)
- choose k so that \approx half of bits are set

Larger *m* means reading more descriptor data.

Having k too high \Rightarrow increased overlapping. Having k too low \Rightarrow increased hash collisions.

... SIMC Parameters

How to determine "optimal" *m* and *k*?

- 1. start by choosing acceptable p_F (e.g. $p_F \le 10^{-5}$ i.e. one false match in 10,000)
- 2. then choose m and k to achieve no more than this p_F .

Formulae to derive m and k given p_F and n:

$$k = 1/\log_e 2 \cdot \log_e (1/p_F)$$

 $m = (1/\log_e 2)^2 \cdot n \cdot \log_e (1/p_F)$

Query Cost for SIMC

Cost to answer *pmr* query: $Cost_{pmr} = b_D + b_q$

- read r descriptors on b_D descriptor pages
- then read b_q data pages and check for matches

 $b_D = ceil(r/c_D)$ and $c_D = floor(B/ceil(m/8))$

E.g.
$$m=64$$
, $B=8192$, $r=10^4 \Rightarrow c_D = 1024$, $b_D=10$

 b_q includes pages with r_q matching tuples and r_F false matches

Expected false matches = $r_F = (r - r_q).p_F \approx r.p_F$ if $r_q \ll r$

E.g. Worst $b_q = r_q + r_F$, Best $b_q = 1$, Avg $b_q = ceil(b(r_q + r_F)/r)$

Exercise 1: SIMC Query Cost

Consider a SIMC-indexed database with the following properties

- all pages are B = 8192 bytes
- tuple descriptors have m = 64 bits (= 8 bytes)
- total records r = 102,400, records/page c = 100
- false match probability p_F = 1/1000
- answer set has 1000 tuples from 100 pages
- 90% of false matches occur on data pages with true match
- 10% of false matches are distributed 1 per page

Calculate the total number of pages read in answering the guery.

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SIMC has one descriptor per tuple ... potentially inefficient.

Alternative approach: one descriptor for each data page.

Every attribute of every tuple in page contributes to descriptor.

Size of page descriptor (PD) (clearly larger than tuple descriptor):

• use above formulae but with *c.n* "attributes"

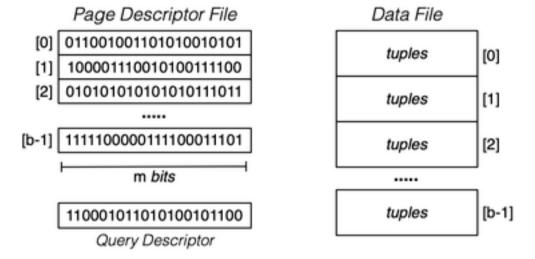
E.g.
$$n = 4$$
, $c = 64$, $p_F = 10^{-3} \implies m \approx 3680$ bits ≈ 460 bytes

Typically, pages are 1..8KB \Rightarrow 8..64 PD/page (c_{PD}).

E.g.
$$m \approx 460$$
, $B = 8192$, $c_{PD} \approx 17$

... Page-level SIMC

File organisation for page-level superimposed codeword index



... Page-level SIMC

Algorithm for evaluating *pmr* query using page descriptors

```
pagesToCheck = {}
for each descriptor D[i] in signature file {
    if (matches(D[i],desc(q))) {
        pid = i
            pagesToCheck = pagesToCheck U pid
    }
}
for each pid in pagesToCheck {
    Buf = getPage(dataFile,pid)
        check tuples in Buf for answers
}
```

Exercise 2: Page-level SIMC Query Cost

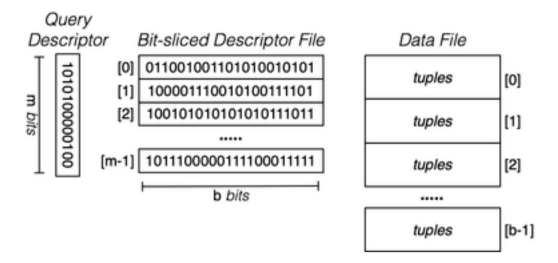
Consider a SIMC-indexed database with the following properties

- all pages are B = 8192 bytes
- page descriptors have m = 4096 bits (= 512 bytes)
- total records r = 102,400, records/page c = 100
- false match probability $p_F = 1/1000$
- answer set has 1000 tuples from 100 pages
- 90% of false matches occur on data pages with true match
- 10% of false matches are distributed 1 per page

Calculate the total number of pages read in answering the query.

Bit-sliced SIMC

Improvement: store *b m*-bit page descriptors as *m b*-bit "bit-slices"



... Bit-sliced SIMC 20/73

Algorithm for evaluating *pmr* query using bit-sliced descriptors

```
matches = ~0  //all ones
for each bit i set to 1 in desc(q) {
    slice = fetch bit-slice i
    matches = matches & slice
}
for each bit i set to 1 in matches {
    fetch page i
    scan page for matching records
}
```

Effective because desc(q) typically has less than half bits set to 1

Exercise 3: Bit-sliced SIMC Query Cost

Consider a SIMC-indexed database with the following properties

- all pages are B = 8192 bytes
- r = 102,400, c = 100, b = 1024
- page descriptors have m = 4096 bits (= 512 bytes)
- bit-slices have *b* = 1024 bits (= 128 bytes)
- false match probability $p_F = 1/1000$
- query descriptor has k = 10 bits set to 1
- answer set has 1000 tuples from 100 pages
- 90% of false matches occur on data pages with true match
- 10% of false matches are distributed 1 per page

Calculate the total number of pages read in answering the query.

Assignment 2

Assignment 2

Aim: implement all variants of SIMC indexing

Implement individual relations and commands to work on them.

Each relation R consists of multiple files:

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- R.info ... relation meta-data (e.g. # tuples)
- R.data ... data file containing pages of tuples
- **R.tsig** ... file containing tuple signatures
- **R.psig** ... file containing page-level signatures
- **R.bsig** ... file containing bit-sliced signatures

... Assignment 2

File structures for SIMC info + data files

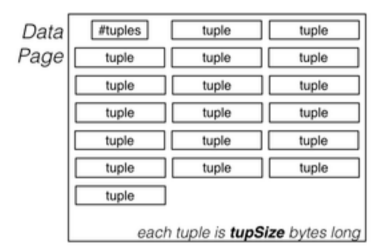


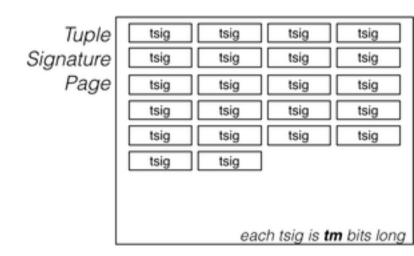
Tuples are all tupSize bytes long (based on # attributes)

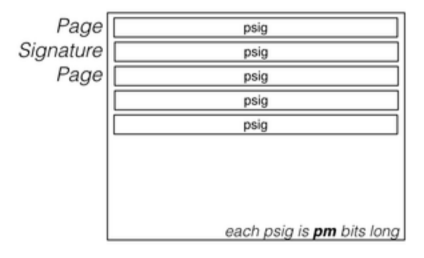
Signatures are *m* bits long, rounded to *ceil(m/8)* bytes

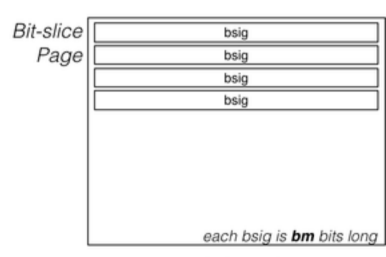
... Assignment 2

More detailed file structures for SIMC data + signature files









in theory, bm == npages

... Assignment 2

We supply:

- complete command programs to build and query relations
- partially-complete ADTs for operations needed by commands

You complete the ADTs so that the commands work properly

- create, insert, select ... build/query commands
- gendata, stats, dump ... utility commands
- x1, x2, x3 ... commands for debugging ADTs

... Assignment 2

```
./create RelName #tuples #attrs 1/pF
```

- creates a new relation with prefix RelName
- uses #attrs to determine tuple size
- uses #tuples to determine b ⇒ length of bit-slices
 (#tuples suggests maximum number of tuples to be stored)
- uses pF to determine m and k for signatures
- creates files RelName.info, RelName.data, etc. etc.
- as supplied, data and signature files are empty after create
- when complete, RelName.bsig should contain all-zero bit-slices

... Assignment 2

./gendata #tuples #attributes [startID] [seed]

generates #tuples tuples in a standard format, e.g.

```
1234567, iuwhfkajewhkfjkwefbx, a3-101, a4-256, a5-013,...
```

- first attribute is unique id
- second attribute is 20-char random string (most likely unique)
- rest are aN-DDD up to #attributes
- attributes a3-DDD to aN-DDD are not unique
- if no startID given, use 1000000; if no seed given, use 0

... Assignment 2

./insert [-v] RelName

- insert tuples read from stdin into Relname
- updates all files: info, data, signature files
- tuples look like those generated from gendata
- typical usage
 - ./gendata #tuples #attrs | ./insert RelName
- -v displays each tuple and PID of page where inserted

... Assignment 2

./select RelName 'Query' SigType

- finds all matches for pmr query in relation RelName
- queries are expressed using ? for unknown attributes, e.g.

```
?,?,?,?  # matches all tuples
1234567,?,?,?,?  # matches single tuple with this ID
?,?,a3-101,?,?  # matches tuples with a3-101 as 3rd attr
?,?,a3-101,a4-200,a5-013
```

- should enclose Query in single quotes (to avoid problems with zsh)
- prints one tuple per line (note: order of tuples is not important)
- prints page read statistics at end of output

Method for ./select RelName 'Query' SigType

... Assignment 2

```
q = startQuery(r, qstr, type):
    check for valid query (e.g. #tuples)
    T = type of signature (t,p,b,x)
    Sig = build query signature of type T
    use Sig to determine list of interesting pages
    q->pages = bit-string of interesting pages
if (q == NULL) fatal error
scanAndDisplayMatchingTuples(Query q):
    foreach PID in q->pages {
        Buf = get page PID
        scan Buf for real matches and display each
}
```

Bits ADT

```
ADT to represent arbitrary-length bit-strings
Bits newBits(int n);
    • create a new bit-string of length n
Bool bitIsSet(Bits b, int i);
    • check whether bit i is set to 1 in bit-string b
void setBit(Bits b, int i);
    • set the i<sup>th</sup> bit of bit-string b to 1
void unsetBit(Bits b, int i);
    • set the i<sup>th</sup> bit of bit-string b to 0
Bit-strings of length n are indexed from 0 (least sig) .. n-1 (most sig)
                                                                                                                              33/73
Reln ADT
ADT to represent relations (where RelnRep *Reln)
Status newRelation(name, nattrs,pF,tk,tm,pm,bm)
    • make all files for relation name, based on parameters
Reln openRelation(char *name)
    • create a RelnRep, populate it and open all files
void closeRelation(Reln r)

    close all files and clean up RelnRep data structure

PageID addToRelation(Reln r, Tuple t)
    • insert tuple t into open relation r; return PID where inserted
                                                                                                                              34/73
Query ADT
ADT to represent queries (where QueryRep *Query)
Query startQuery(Reln r, char *qry, char sigType)
    • set up QueryRep for query qry for specific type of signature
void scanAndDisplayMatchingTuples(Query q)
    • evaluate the query q and display result tuples, one per line
void queryStats(Query q)
    • print statistics from the QueryRep, typically after query finishes
void closeQuery(Query)
    • clean up QueryRep data structure (i.e. free)
```

Signature ADTs

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Three different types of signature: tuple, page, bit-slice Each has its own ADT, but all ADTs are similar

Bits makeXSig(Reln r, Tuple t)

• build a signature of type X for tuple t

void findPagesUsingXSigs(Query q)

- uses Xsig to build bit-string of potentially-matching pages
- result is stored in q->pages (of type Bits)

Note: don't need makeBitSliceSig(); it uses page signatures

Psig ADT

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ADT to represent page signatures

Bsig ADT

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ADT to represent bit-sliced signatures

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What to do now?

... Bsig ADT

- review the notes on superimposed codewords
- read the spec carefully
- read the code for the commands (to see how they use the ADTs)
- read the ADT *.h files; read the ADT *.c files
- write and test your code (suggest: tsig, then psig, then bsig)

Testing script available next week.

Don't wait. It's easy to devise your own tests.

N-d Tree Indexes

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Over the last 20 years, from a range of problem areas

Multi-dimensional Tree Indexes

- different multi-d tree index schemes have been proposed
- varying primarily in how they partition tuple-space

Consider three popular schemes: kd-trees, Quad-trees, R-trees.

Example data for multi-d trees is based on the following relation:

```
create table Rel (
    X char(1) check (X between 'a' and 'z'),
    Y integer check (Y between 0 and 9)
);
```

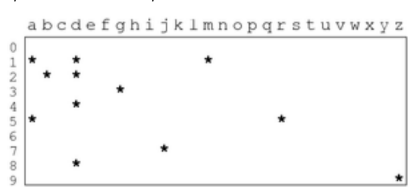
... Multi-dimensional Tree Indexes

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Example tuples:

```
Rel('a',1) Rel('a',5) Rel('b',2) Rel('d',1) Rel('d',2) Rel('d',4) Rel('d',8) Rel('g',3) Rel('j',7) Rel('m',1) Rel('r',5) Rel('z',9)
```

The tuple-space for the above tuples:

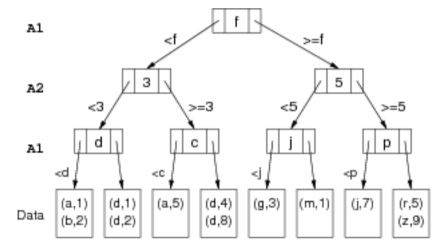


```
Q1 ... Q2 ... Q3 ... Q4
```

kd-Trees 43/73

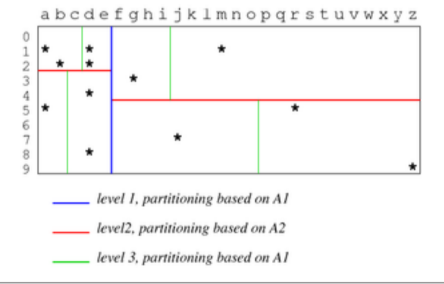
kd-trees are multi-way search trees where

- each level of the tree partitions on a different attribute
- each node contains n-1 key values, pointers to n subtrees



... kd-Trees

How this tree partitions the tuple space:



Searching in kd-Trees

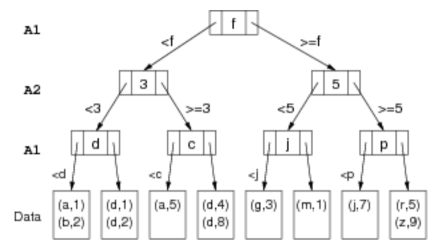
```
// Started by Search(Q, R, 0, kdTreeRoot)
Search(Query Q, Relation R, Level L, Node N)
{
   if (isDataPage(N)) {
     Buf = getPage(fileOf(R),idOf(N))
     check Buf for matching tuples
} else {
   a = attrLev[L]
```

```
if (!hasValue(Q,a))
    nextNodes = all children of N
else {
    val = getAttr(Q,a)
    nextNodes = find(N,Q,a,val)
}
for each C in nextNodes
    Search(Q, R, L+1, C)
}
}
```

Exercise 5: Searching in kd-Trees

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Using the following kd-tree index



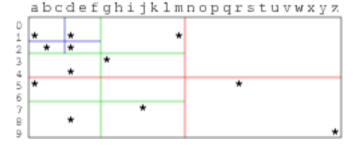
Answer the queries (m,1), (a,?), (?,1), (?,?)

Quad Trees

Quad trees use regular, disjoint partitioning of tuple space.

- for 2d, partition space into quadrants (NW, NE, SW, SE)
- each quadrant can be further subdivided into four, etc.

Example:



... Quad Trees

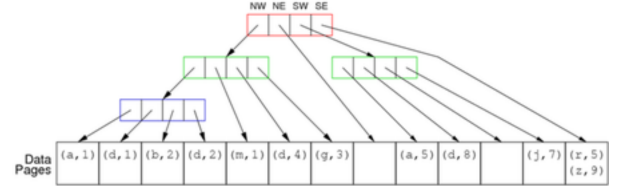
Basis for the partitioning:

- a quadrant that has no sub-partitions is a leaf quadrant
- each leaf quadrant maps to a single data page
- subdivide until points in each quadrant fit into one data page
- ideal: same number of points in each leaf quadrant (balanced)
- point density varies over space
 - ⇒ different regions require different levels of partitioning
- this means that the tree is not necessarily balanced

Note: effective for d≤5, ok for 6≤d≤10, ineffective for d>10

... Quad Trees

The previous partitioning gives this tree structure, e.g.

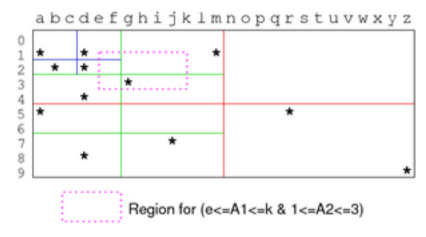


In this and following examples, we give coords of top-left,bottom-right of a region

Searching in Quad-tree

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Space query example:

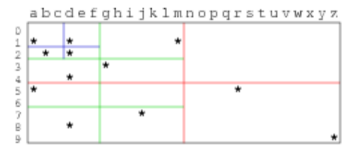


Need to traverse: red(NW), green(NW,NE,SW,SE), blue(NE,SE).

Exercise 6: Searching in Quad-trees

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Using the following quad-tree index



Answer the queries (m,1), (a,?), (?,1), (?,?)

R-Trees 52/73

R-trees use a flexible, overlapping partitioning of tuple space.

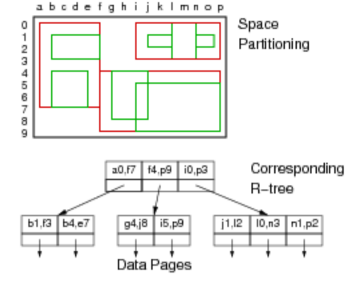
- each node in the tree represents a kd hypercube
- its children represent (possibly overlapping) subregions
- the child regions do not need to cover the entire parent region

Overlap and partial cover means:

- can optimize space partitioning wrt data distribution
- so that there are similar numbers of points in each region

Aim: height-balanced, partly-full index pages (cf. B-tree)

... R-Trees 53/73



Insertion into R-tree 54/73

Insertion of an object R occurs as follows:

- start at root, look for children that completely contain R
- if no child completely contains R, choose one of the children and expand its boundaries so that it does contain R
- if several children contain R, choose one and proceed to child
- · repeat above containment search in children of current node
- once we reach data page, insert R if there is room
- if no room in data page, replace by two data pages
- partition existing objects between two data pages
- update node pointing to data pages
 (may cause B-tree-like propagation of node changes up into tree)

Note that R may be a point or a polygon.

Query with R-trees

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Designed to handle space queries and "where-am-I" queries.

"Where-am-I" query: find all regions containing a given point P:

- start at root, select all children whose subregions contain P
- if there are zero such regions, search finishes with P not found
- otherwise, recursively search within node for each subregion
- once we reach a leaf, we know that region contains P

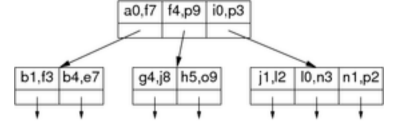
Space (region) queries are handled in a similar way

we traverse down any path that intersects the query region

Exercise 7: Query with R-trees

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Using the following R-tree:



Show how the following queries would be answered:

```
Q1: select * from Rel where X='a' and Y=4 Q2: select * from Rel where X='i' and Y=6 Q3: select * from Rel where 'c' \le X \le 'j' and Y=5 Q4: select * from Rel where X='c'
```

Note: can view unknown value X=? as range $min(X) \le X \le max(X)$

Up to version 8.2, PostgreSQL had an R-tree implementation Superseded by GiST = Generalized Search Trees GiST indexes parameterise: data type, searching, splitting • via seven user-defined functions (e.g. picksplit()) GIST trees have the following structural constraints: every node is at least fraction f full (e.g. 0.5) the root node has at least two children (unless also a leaf) all leaves appear at the same level Details: src/backend/access/gist 58/73 **Costs of Search in Multi-d Trees** Difficult to determine cost precisely. Best case: pmr query where all attributes have known values in kd-trees and quad-trees, follow single tree path cost is equal to depth D of tree in R-trees, may follow several paths (overlapping partitions) Typical case: some attributes are unknown or defined by range need to visit multiple sub-trees how many depends on: range, choice-points in tree nodes **Similarity-based Selection** 60/73 **Relational vs Similarity Selection** Relational selection is based on a boolean condition C evaluate C for each tuple t (or a likely subset of tuples) if C(t) is true, add t to result set if C(t) is false, t is not part of solution result is a set of tuples $\{t_1, t_2, ..., t_n\}$ all of which satisfy C Uses for relational selection: precise matching on structured data using individual attributes with known, exact values 61/73 ... Relational vs Similarity Selection Similarity selection is used in contexts where cannot define a precise matching condition but can identify a notion of (S similar to T) requires a measure d of "distance" between tuples d=0 is an exact match, d>0 is less accurate match result is a list of pairs [(t_1,d_1) , (t_2,d_2) , ..., (t_n,d_n)] (ordered by d_i)

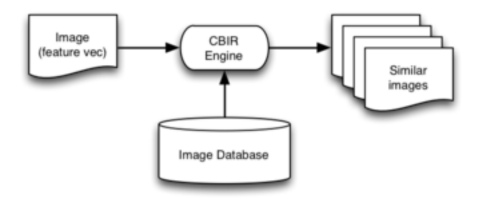
Uses for similarity matching:

- text or multimedia (image/music) retrieval
- ranked queries in conventional databases

Example: Content-based Image Retrieval

User supplies a description or sample of desired image.

System returns a ranked list of similar images from database.



... Example: Content-based Image Retrieval

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At the SQL level, this might appear as ...

Similarity-based Retrieval

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Database contains media objects, but also tuples, e.g.

- id to uniquely identify object (e.g. PostgreSQL oid)
- metadata (e.g. artist, title, genre, date taken, ...)
- value of object itself (e.g. PostgreSQL BLOB or bytea)

BLOB = Binary Large OBject

- BLOB stored in separate file; tuple contains reference (cf. TOAST)
- BLOBs are typically MB in size (1MB..2GB)

... Similarity-based Retrieval

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Similarity-based retrieval requires a distance measure

• $dist(x,y) \in 0..1$, dist(x,x) = 0, dist(x,y) = dist(y,x)

where x and y are two objects (in the database)

Note: distance calculation often requires substantial computational effort

How to restrict solution set to only the "most similar" objects:

- threshold d_{max} (only objects t such that $dist(t,q) \le d_{max}$)
- count k (k closest objects (k nearest neighbours))

BUT both above methods require knowing distance between query object and all objects in DB

... Similarity-based Retrieval

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Naive approach to similarity-based retrieval

```
// query object
dmax = ... // dmax > 0 =>
                            using threshold
knn = \dots // knn > 0
                            using nearest-neighbours
                       =>
Dists = [] // empty list
foreach tuple t in R {
    d = dist(t.val, q)
    insert (t.oid,d) into Dists // sorted on d
n = 0: Results = []
foreach (i,d) in Dists {
    if (dmax > 0 && d > dmax) break;
    if (knn > 0 && ++n > knn) break;
    insert (i,d) into Results // sorted on d
return Results:
```

Cost = fetch all r objects + compute distance() for each

For some applications, Cost(dist(x,y)) is comparable to T_t ⇒ computing dist(t.val,q) for every tuple t is infeasible. compute feature vector to capture "critical" object properties store feature vectors "in parallel" with objects (cf. signatures) compute distance using feature vectors (not objects) i.e. replace dist(t,q) by dist'(vec(t),vec(q)) in previous algorithm. Further optimisation: dimension-reduction to make vectors smaller 68/73 ... Similarity-based Retrieval Feature vectors ... often use multiple features, concatenated into single vector represent points in a very high-dimensional (vh-dim) space Content of feature vectors depends on application ... • image ... colour histogram (e.g. 100's of values/dimensions) music ... loudness/pitch/tone (e.g. 100's of values/dimensions) text ... term frequencies (e.g. 1000's of values/dimensions) Query: feature vector representing one point in vh-dim space Answer: list of objects "near to" query object in this space 69/73 ... Similarity-based Retrieval Inputs to content-based similarity-retrieval: • a database of r objects $(obj_1, obj_2, ..., obj_r)$ plus associated ... • r × n-dimensional feature vectors (v_{obj1}, v_{obj2}, ..., v_{obj}) a query image q with associated n-dimensional vector (v_q) a distance measure $D(v_i,v_j):[0..1)$ $(D=0 \rightarrow v_i=v_j)$ Outputs from content-based similarity-retrieval: a list of the k nearest objects in the database [a₁, a₂, ... a_k] • ordered by distance $D(v_{a_1}, v_q) \le D(v_{a_2}, v_q) \le \dots \le D(v_{a_{k'}}, v_q)$ 70/73 Approaches to kNN Retrieval Partition-based use auxiliary data structure to identify candidates space/data-partitioning methods: e.g. k-d-B-tree, R-tree, unfortunately, such methods "fail" when #dims > 10..20 absolute upper bound on d before linear scan is best d = 610Approximation-based use approximating data structure to identify candidates signatures: VA-files projections: iDistance, LSH, MedRank, CurveIX, Pyramid 71/73 ... Approaches to kNN Retrieval Above approaches try to reduce number of objects considered. cf. indexes in relational databases Other optimisations to make kNN retrieval faster reduce I/O by reducing size of vectors (compression, d-reduction)

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- reduce I/O by placing "similar" records together (clustering)
- reduce I/O by remembering previous pages (caching)
- reduce cpu by making distance computation faster

Similarity Retrieval in PostgreSQL

PostgreSQL has always supported simple "similarity" on strings

```
- for most SQL implementations
select * from Students where name like '%oo%';
 - and PostgreSQL-specific
select * from Students where name ~ '[Ss]mit';
```

Also provides support for ranked similarity on text values

- using tsvector data type (stemmed, stopped feature vector for text)
- using tsquery data type (stemmed, stopped feature vector for strings)

• using @@ similarity operator

... Similarity Retrieval in PostgreSQL

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Example of PostgreSQL text retrieval:

```
create table Docs
    ( id integer, title text, body text );
// add column to hold document feature vectors
alter table Docs add column features tsvector;
update Docs set features =
    to_tsvector('english', title||''||body);
// ask query and get results in ranked order
select title, ts_rank(d.features, query) as rank
from Docs d,
    to_tsquery('potter|(roger&rabbit)') as query
where query @@ d.features
order by rank desc
limit 10;
```

For more details, see PostgreSQL documentation, Chapter 12.

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