







Overview: The pq-Gram Index

• pq-Gram profile:



- Algorithm
- Fanout Weighting and Lower Bound
- Experiments

The pq-Gram Distance Definition

The pq-Gram Distance

Definition (pq-Gram Distance)

The pq-gram distance between two trees, T and T', is defined as

$$\delta_{g}(\mathsf{T},\mathsf{T}') = |\mathcal{I}(\mathsf{T}) \uplus \mathcal{I}(\mathsf{T}')| - 2|\mathcal{I}(\mathsf{T}) \cap \mathcal{I}(\mathsf{T}')|$$

• Metric normalization to [0..1]: $\delta'_g(\mathsf{T},\mathsf{T}') = \frac{\delta_g(\mathsf{T},\mathsf{T}')}{|\mathcal{I}(\mathsf{T}) \uplus \mathcal{I}(\mathsf{T}')| - |\mathcal{I}(\mathsf{T}) \cap \mathcal{I}(\mathsf{T}')|}$ • Pseudo-metric properties hold for normalization [ABG10]: Self-identity: $x = y \not\models \Rightarrow \delta_{g}(x, y) = 0$ Symmetry: $\delta_{\sigma}(x, y) = \delta_{\sigma}(y, x)$ • triangle inequality: $\delta_{\mathfrak{g}}(x,z) \leq \delta_{\mathfrak{g}}(x,y) + \delta_{\mathfrak{g}}(y,z)$ а а / \ / \ b b b • Different trees may have identical indexes: С С Augsten (Univ. Salzburg) Similarity Search WS 2019/20 14 / 32

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The pq-Gram Distance Algorithm
Main Memory Algorithm (I)
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CREATEINDEX($\mathsf{T}, \mathsf{r}, \mathcal{I}, stem, p, q$)

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stem := shift(stem, \lambda(\mathbf{r}))
base: shift register of size q (filled with *)
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if r is a leaf then
   \mathcal{I} := \mathcal{I} \cup \{ stem \circ base \}
else
   for each child c (from left to right) of r do
         base := shift(base, \lambda(c))
         \mathcal{I} := \mathcal{I} \cup \{ stem \circ base \}
         \mathcal{I} := CREATEINDEX(T, c, \mathcal{I}, stem, p, q)
   for k := 1 to q - 1
         base := shift(base, *)
         \mathcal{I} := \mathcal{I} \cup \{ stem \circ base \}
```

return \mathcal{I}

The pq-Gram Distance Algorithm The pq-Gram Distance Algorithm Main Memory Algorithm (II) Main Memory Algorithm (III) • Input of CREATEINDEX(T, r, *I*, *stem*, *p*, *q*): • pq-GRAM-INDEX(T, p, q) computes the pq-gram index for a • a subtree of T rooted in r complete tree T: • the *pg*-gram index \mathcal{I} computed so far • the stem *stem* of r's parent pq-GRAM-INDEX (T, p, q) • the parameters p and q stem: shift register of size p (filled with *) • Output of CREATEINDEX(T, r, *I*, *stem*, *p*, *q*): \mathcal{I} : empty index *pq*-gram index including $\mathcal{I} = \text{CREATEINDEX}(\mathsf{T}, \text{root}(\mathsf{T}), \mathcal{I}, stem, p, q)$ • the input index \mathcal{I} return \mathcal{I} • the pg-gram index of r and all its descendants i.e., the pq-grams (label tuples) with anchor node r or a descendant of r Augsten (Univ. Salzburg WS 2019/20 17/32 Augsten (Univ. Salzburg) Similarity Search WS 2019/20 Similarity Search 18 / 32 The pq-Gram Distance Algorithm The pq-Gram Distance Algorithm Complexity of the pq-Gram Index Algorithm Size of the *pq*-Gram Index Theorem (Size of the pq-Gram Index) Let T be a tree with I leaves and i non-leaves. The size of the pg-gram index of T is Theorem (pq-Gram Index Complexity) $|\mathcal{I}^{pq}(\mathsf{T})| = 2l + qi - 1.$ The pg-gram index of a tree T with size |T| can be computed in O(|T|)Proof. time. • We count all pq-grams whose leftmost leaf is a dummy node: Each

Proof.

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Each recursive call of *createIndex*() processes one node in constant time, and each node is processed exactly once. $\hfill \square$

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19/32

Overall number of pq-grams: l + i(q - 1) + (l + i - 1) = 2l + qi - 1.

leaf is the anchor node of exactly one pq-gram whose leftmost leaf is a

dummy node, giving *l* pq-grams. Each non-leaf is the anchor of q-1

pq-grams whose leftmost leaf is a dummy, giving i(q-1) pq-grams.

We count all *pq*-grams whose leftmost leaf is not a dummy node: Each node of the tree except the root is the leftmost leaf of exactly

one pq-gram, giving l + i - 1 pq-grams.



The pq-Gram Distance Fanout Weighting and Lower Bound pq-Gram Distance Lower Bound Outline

Theorem

Proof.

Let p = 1 and $c \ge \max(2q - 1, 2)$ be the cost of changing a leaf node. The pq-gram distance provides a lower bound for the fanout weighted tree edit distance, i.e., for any two trees, T and T',

 $\frac{\delta_g(\mathsf{T},\mathsf{T}')}{2} \leq \delta_f(\mathsf{T},\mathsf{T}').$

See [ABG10] (ACM Transactions on Database Systems).

• pq-Gram index size: linear in the tree size

• Experiment:

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• compute *pq*-gram index for trees with different number of nodes

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The pq-Gram Distance Experiments

• compare tree and index size

Size of the *pq*-Gram Index



Why is the *pq*-gram index smaller than the tree?

- hash values are smaller than labels
- duplicate *pq*-grams of a tree are stored only once



(Artificial tree with 144 nodes, 102 leaves, fanout 2-6 and depth 6. Average over 100 runs.)

WS 2019/20

25 / 32

The *pq*-Gram Distance Experiments

Sensitivity to Structure Change — Non-Leaf

- Cost for non-leaf change \rightarrow controlled by p
- Experiment:
 - delete non-leaf nodes
 - measure normalized *pq*-gram distance



(Artificial tree with 144 nodes, 102 leaves, fanout 2-6 and depth 6. Average over 100 runs.)

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Scalability to Large Trees

- pq-gram distance \rightarrow scalable to large trees
- compare with edit distance
- Experiment: For pair of trees
 - compute tree edit distance and *pq*-gram distance
 - vary tree size: up 5×10^5 nodes
 - measure wall clock time



The pq-Gram Distance Experiments

Influence of *p* and *q* on Scalability

- Scalability (almost) independent of p and q.
- Experiment: For pair of trees
 - compute pq-gram distance for varying p and q
 - vary tree size: up 10⁶ nodes
 - measure wall clock time



pg-Grams vs. other Edit Distance Approximations

The pq-Gram Distance Experiments

Effectiveness: *pq*-grams outperform all other approximations

Experiment: two sets of address trees (299 and 302 trees)

- compute distances between all tree pairs
- find matches (symmetric nearest neighbor)

Distance	Correct	Recall	Precision	f-Measure	Runtime
fanout edit dist	259	86.6%	98.5%	0.922	19 min
unit edit dist	247	82.6%	96.5%	0.890	14 min
node intersection	197	65.9%	93.8%	0.774	4.3s
p,q-grams	236	78.9%	98.7%	0.877	8.1s
tree-embedding	206	68.9%	96.3%	0.803	7.1s
binary branch	193	64.5%	93.2%	0.763	7.4s
bottom-up	148	49.6%	92.5%	0.645	67.0s

WS 2019/20 29/32

