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Diverse and proportional size-*l* object summaries using pairwise relevance

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Abstract The abundance and ubiquity of graphs (e.g., online social networks such as Google+ and Facebook; bibliographic graphs such as DBLP) necessitates the effective and efficient search over them. Given a set of keywords that can identify a data subject (DS), a recently proposed keyword search paradigm produces a set of object summaries (OSs) as results. An OS is a tree structure rooted at the DS node (i.e., a node containing the keywords) with surrounding nodes that summarize all data held on the graph about the DS. OS snippets, denoted as size-l OSs, have also been investigated. A size-l OS is a partial OS containing l nodes such that the summation of their importance scores results in the maximum possible total score. However, the set of nodes that maximize the total importance score may result in an uninformative size-l OSs, as very important nodes may be repeated in it, dominating other representative information. In view of this limitation, in this paper, we investigate the effective and efficient generation of two novel types of OS snippets, i.e.,

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diverse and proportional size-l OSs, denoted as DSize-l and PSize-l OSs. Namely, besides the importance of each node, we also consider its pairwise relevance (similarity) to the other nodes in the OS and the snippet. We conduct an extensive evaluation on two real graphs (DBLP and Google+). We verify effectiveness by collecting user feedback, e.g., by asking DBLP authors (i.e., the DSs themselves) to evaluate our results. In addition, we verify the efficiency of our algorithms and evaluate the quality of the snippets that they produce.

Keywords Keyword search · Diversity · Proportionality · Snippets · Summaries

1 Introduction

Keyword search on the Web has dominated our lives, as it facilitates users to find easily and effectively information using only keywords. For instance, the result for query q = "Faloutsos" consists of a set of links to Web pages containing the keyword(s) together with their respective *snippets*. Snippets are short fragments of text extracted from the search results (e.g., Web pages); they significantly enhance the usability of search results as they provide an intuition about which results are worth accessing and which can be ignored. Furthermore, snippets may provide the complete answer to the searcher's actual information needs (if, for example, the user is only interested in whether Michalis Faloutsos is a Professor), thus preventing the need to retrieve the actual result [27].

The keyword search paradigm has also been introduced in relational databases (e.g., [20]), where the objective is to find networks of tuples connected via foreign key links that collectively contain the keywords. For example, the query "Faloutsos" + "Agrawal" over the DBLP database returns



Example 1 The OS for Michalis Faloutsos

Author: Michalis Faloutsos
Paper: On Power-law Relationships of the Internet Topology.
Conference: SIGCOMM. Year: 1999.
Co-Author(s): Christos Faloutsos, Petros Faloutsos.
Cites: Building Shared..., Cited by: The Structure...,
Paper: BLINC: Multilevel Traffic Classification in the Dark.
Conference: ACM SIGCOMM Computer Comm. Review Year:2005.
Co-Author(s): T. Karagiannis, K. Papagiannaki.
Cites: A Parametrizable methodology..., Cited by: P4P: Provider...,
Paper: Transport Layer Identification of P2P Traffic.
Conference: SIGCOMM. Year:2004.
Co-Author(s): T. Karagiannis, A. Broido.
Cites: Their Share: Diversity..., Cited by: Internet Traffic......

tuples Faloutsos and Agrawal from the author table and their associations through co-authored papers. However, relational keyword search may not be very effective when searching information about a particular *data subject* (DS) (e.g., Faloutsos and his papers, co-authors). A DS is an entity (e.g., an individual, paper, product) which has its identity in a tuple which is the result (i.e., subject) of the keyword search. Relational keyword search only returns tuples containing the keywords (in this case, only Faloutsos author tuples) and hence fails to address search for the *context* of most important tuples around a central tuple (i.e., a DS).

In view of this, in [12,13], the concept of object summary (OS) was introduced; an OS summarizes all data held in a database about a particular DS, searched by some keyword(s). More precisely, an OS is a tree with the tuple $n^{\rm DS}$ containing the keywords (e.g., author tuple M. Faloutsos) as the root node and its neighboring tuples, containing additional information (e.g., his papers, co-authors), as child or descendant nodes. The precise definition of an OS is discussed in Sect. 2; in a nutshell, a tuple is included in the OS if it is of high affinity (based on link properties in the tuple network graph of the database) and it is connected to n^{DS} via a short path. For instance, the result for q is a set of OSs: one for each Faloutsos brother. Example 1 illustrates the OS for Michalis Faloutsos. Note that the OS paradigm is in more analogy to Web keyword search, compared to relational keyword search. For instance, Example 1 resembles a Web page (as it includes comprehensive information about the DS). Therefore, for the non-technical users with experience only on Web keyword search, the OS paradigm will be friendlier and also closer to their expectations. In general, an OS is a concise summary of the *context* around any pivot database tuple or graph node, finding application in (interactive) data exploration, schema extraction, etc. Another application of this summarization concept is on semantic knowledge graphs [6, 26].

In [15,16], OS snippets were proposed (denoted as size-l OSs). Size-l OSs are composed of only l important nodes so that (1) the summation of their scores is maximized and (2) all l nodes are connected to the OS root (i.e., n DS). Example 2 illustrates the size-l OS for M. Faloutsos with l = 15 on



Author: Michalis Faloutsos

Paper: On Power-law Relationships of the Internet Topology.

Co-Author: Christos Faloutsos,...

Paper: Power Laws and the AS-Level Internet Topology.

Co-Author: Christos Faloutsos,...

Paper: ACM SIGCOMM' 99. Co-Author: <u>Christos Faloutsos</u>,...
Paper: Information survival thr.... Co-Author: <u>Christos Faloutsos</u>,...
Paper: The Connectivity and Fault... Co-Author: <u>Christos Faloutsos</u>,...
Paper: BGP-lens: Patterns and An... Co-Author: <u>Christos Faloutsos</u>,...
Paper: The eBay Graph: How Do.... Co-Author: <u>Christos Faloutsos</u>,...

Example 3 DSize-15 OS for Michalis Faloutsos

Author: Michalis Faloutsos

Paper: On Power-law Relationships of the Internet Topology.

Conference: SIGCOMM. Year: 1999. Co-Author: Christos Faloutsos,

Paper: Information Survival Threshold in Sensor and P2P Networks.

Co-Author: S. Madden, ..., Conference: INFOCOM.

Paper: Power Laws and the AS-Level Internet Topology.

Conference: IEEE/ACM Tr. Netw. Year: 2003.

Co-Author: Christos Faloutsos,..

Paper: Network Monitoring Using Traffic Dispersion Graphs.
Co-Author: M. Mitzenmacher, ...Conference: SIGCOMM.

Example 4 PSize-15 OS for Michalis Faloutsos

Author: Michalis Faloutsos

Paper: On Power-law Relationships of the Internet Topology.

Conference: SIGCOMM. Year: 1999. Co-Author: Christos Faloutsos,

Paper: Denial of Service Attacks at the MAC Layer...

Co-Author: S. Krishnamurthy, ..., Conference: MILCOM. Paper: Power Laws and the AS-Level Internet Topology.

Conference: IEEE/ACM Tr. Netw.

Co-Author: Christos Faloutsos,...

Paper: Reducing Large Internet Topologies for Faster Simulations Co-Author: S. Krishnamurthy, L. Cui,...Conference: NETWORKING.

the DBLP database. According to [15], a size-l OS should be a standalone subgraph of the complete OS so that the user can comprehend it without any additional information. For this reason, the l nodes should form a connected graph that includes the root of the OS.

However, this selection criterion (i.e., maximizing importance score) can render such snippets ineffective. For instance, in Example 2, the co-authorship of Michalis with Christos Faloutsos, who is a very important author, monopolizes the snippet with papers co-authored only with Christos. Thus, we argue that the diversity of constituent nodes will improve the snippet's effectiveness. In addition, we argue that frequent appearances of nodes in an OS should also be proportionally represented in an effective snippet.

Hence, in this paper, we propose two novel snippets, namely *diverse* and *proportional* size-*l* OSs denoted as *DSize-l* OS and *PSize-l* OS, respectively. More precisely, in a DSize-*l* OS, we favor diversity by penalizing repetitions of relevant nodes. For instance, the DSize-*l* OS of Example 3 includes C. Faloutsos only twice, allowing the appearance of other important co-authors as well. In a PSize-*l* OS, we favor proportionality, i.e., a frequent relevant node should be analogously represented, facilitating diversity at the same time. Similarly, the PSize-*l* OS of Example 4 includes also frequent co-authors S. Krishnamurthy and L. Cui who do not appear



at all in the DSize-l OS. To compute them, we calculate a *combined* score per node, which integrates (1) importance, (2) affinity to the data subject node n^{DS} and (3) diversity or proportionality.

For the diversity and proportionality scores calculation, we employ two types of pairwise relevance: *Similarity* (denoted also as *sim*) and *Equality* (denoted as *equi*). More precisely, *sim* is the textual similarity between nodes (e.g., Jaccard similarity); e.g., two papers with common keywords are similar. Note that textual similarity on author's names makes little sense here; e.g., two authors with a common surname are still two different persons. Thus, we also use *equi* as a *binary* relevance function, i.e., two OS nodes that correspond to the same graph node (e.g., the same author appearing many times in an OS) have *equi*-relevance 1, otherwise their *equi*-relevance is 0. We say that a snippet is an *equi* size-*l* OS if it considers only *equi* relevance (e.g., an *equi* DSize-*l*); we say that a snippet is a *sim* size-*l* OS if it considers both *sim* and *equi* relevance (e.g., *sim* DSize-*l*).

The efficient generation of DSize-*l* or PSize-*l* OSs is a challenging problem since information about the repetitions and frequencies of nodes is required and incremental computation is not possible (as opposed to the original size-*l* OS computation problem [15]). We discuss a brute-force algorithm, BF-*l*, that produces optimal solutions but scales badly. Then, we propose a greedy algorithm (LASP) and its optimization (2-LASP*e*). Both algorithms are general and can address both DSize-*l* and PSize-*l* OS snippet types (with minor modifications), based on either similarity or equivalence relevance. In addition, we propose two preprocessing techniques for the two snippet types (PPrelim-*l* and DPrelim-*l*) that prune the input OSs before processing them.

We conducted an extensive experimental study on the DBLP bibliographic and Google+ social network datasets. We verify effectiveness by collecting user feedback, e.g., by asking DBLP authors (i.e., the DSs themselves) to evaluate our size-l OSs. The users suggested that the results produced by our method are very close to their expectations and that DSize-l and PSize-l are more usable than the respective size-ls OSs, which disregard diversity. In addition, we investigated in detail and verified the efficiency and approximation quality of our algorithms.

The contributions of this paper can be summarized as follows: (1) the introduction of two novel OS snippets, DSize-*l* and PSize-*l* OS, which capture diversity and proportionality, respectively; (2) the introduction of efficient greedy algorithms for their generation; (3) a theoretical analysis for the greedy algorithms including proofs of lower approximations bounds; (4) an extensive experimental evaluation that verifies the proposed concepts and techniques.

A preliminary version of this paper introduces DSize-*l* and PSize-*l* OSs using only equality relevance [17]. Here, we generalize our relevance measure by also considering pairwise

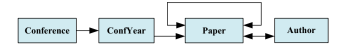


Fig. 1 DBLP database schema

textual similarity (*sim*); this dictates significant amendments of our greedy algorithms. As we demonstrate in Sect. 8, this generalization is significantly better in terms of usability. In addition, in this paper, we enrich the theoretical analysis of our greedy algorithms by including proofs of the lower bounds of their approximations. Finally, we provide a more comprehensive evaluation for the usability, quality and efficiency of size-*l* OSs.

The rest of the paper is structured as follows. Sect. 2 describes background and related work. Section 3 describes the semantics of DSize-*l* and PSize-*l* OSs. Section 4 introduces the optimal solution, whereas Sects. 5 and 6 present the greedy algorithms. Section 7 introduces preprocessing algorithms for DSize-*l* and PSize-*l* OS computation. Section 8 presents experimental results. Finally, Sect. 10 provides concluding remarks.

2 Background work

In this section, we describe background work that we build upon in this paper; namely, we describe the concepts of an *object summary* (OS) and size-*l* OS.

2.1 Object summaries

According to the keyword search paradigm of [13], an object summary (OS) is generated for each node (tuple) n^{DS} found in a graph (database) that contains the query keyword(s) (e.g., "Michalis Faloutsos" node of author relation in the DBLP database). An OS is a tree having n^{DS} as a root, the nodes that link to n^{DS} through foreign keys as its children, and the nodes that link to the children recursively as descendant nodes. To construct an OS, the relation R^{DS} (e.g., the author relation) that holds n^{DS} and those that link to R^{DS} via foreign keys are used. First, a Data Subject Schema Graph G^{DS} is generated. Figure 2 illustrates the G^{DS} for the author relation of the DBLP database, whose schema is shown in Fig. 1. A G^{DS} is a directed labeled tree with a fixed maximum depth that has an R^{DS} as a root node and captures the subset of the schema surrounding R^{DS} ; any surrounding relations participating in loop or many-to-many relationships are replicated accordingly. In other words, a GDS is a "treelization" of the database schema, where R^{DS} becomes the root, R^{DS} 's neighboring relations become child nodes and so on. In order to generate the OS, the relations from G^{DS} which have high affinity with R^{DS} are used. The affinity of a relation R_i to $R^{\rm DS}$ can be calculated by the formula:



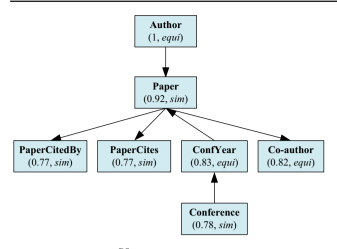


Fig. 2 DBLP author G^{DS} (affinity, relevance type)

$$af(R_i) = \sum_{j} mw_j \cdot m_j \cdot af(R_{Parent}), \tag{1}$$

where j ranges over a set of measures (m_1, m_2, \ldots, m_n) and their corresponding weights $(mw_1, mw_2, \dots, mw_n)$, and $af(R_{Parent})$ is the affinity of R_i 's parent to R^{DS} . The measures' scores range in [0, 1], and the corresponding weights sum to 1; thus, the affinity score of a node is monotonically non-increasing with respect to the node's parent. More precisely, we use four measures: m_1 considers the distance of R_i to $R^{\rm DS}$, i.e., the shorter the distance, the bigger the affinity between the two relations. The remaining measures consider the connectivity of R_i on both the database schema and data graph. m_2 measures the relative cardinality, i.e., the average number of tuples of R_i that are connected with each tuple in R_{Parent} , whereas m_3 measures their reverse relative cardinality, i.e., the average number of tuples of R_{Parent} that are connected with a tuple in R_i . m_4 considers the schema connectivity of R_i (i.e., the number of relations it is connected to in the relation graph). Given a threshold θ , a subset of G^{DS} can be produced that includes only the relations of affinity at least θ to R^{DS} . The OS for a tuple n^{DS} in R^{DS} is generated by traversing the G^{DS} starting from n^{DS} (e.g., by joining n^{DS} with the neighboring relations of R^{DS} ; Algorithm 4). For instance, for q = "Faloutsos" and for $n^{DS} =$ "Michalis Faloutsos" in the author R^{DS} of the DBLP database, the OS presented in Example 1 will be generated.

Every tuple v_i in the database carries a *global importance* weight $gi(v_i)$, calculated using PageRank-inspired measures such as ObjectRank [3] and ValueRank [14]. Due to the "treelization" of the schema graph by G^{DS} , multiple tuples in an OS may correspond to the same tuple in the database. For instance, the same co-author (e.g., Christos Faloutsos) may appear multiple times (e.g., 12) in the OS of Michalis. Formally, for a node n_i of an OS, we use function $g(n_i)$ to denote the corresponding tuple v in the database. Thus, for two OS nodes n_i and n_j , we may have $g(n_i) = g(n_j) = v$.

We also denote as fr(v) (i.e., $fr(g(n_i))$, or simply $fr(n_i)$) the frequency of tuple v in the given OS.

2.2 Size-l OSs

According to [15], given an OS and an integer l, a candidate size-l OS is any subset of the OS composed of l nodes, such that the l nodes form a tree rooted at n^{DS} . In [15], we argue that a good size-l OS should be a standalone and meaningful synopsis of the most important and representative information about the particular DS (so that users can understand it as is, without any additional nodes). In particular, any intermediate nodes that connect n^{DS} (e.g., M. Faloutsos) with other important nodes (e.g., C. Faloutsos) in the size-l OS guarantee that the size-l remains standalone, since these connecting nodes (e.g., co-authored papers) include the semantics of the associations. For instance, in Example 2, if we exclude the paper "On Power-law ..." but only include the co-authors, we exclude the semantic association between n^{DS} and co-author(s), which in this case is their common paper.

The holistic importance Im(Sl) of any candidate size-l OS Sl is defined as the sum of the local importance scores of its nodes, i.e., $Im(Sl) = \sum_{n_i \in Sl} li(n_i)$. The local importance of a node n_i is the affinity-weighted global importance of n_i , i.e., $li(n_i) = af(n_i) \cdot gi(n_i)$. The affinity $af(n_i)$ of a node n_i equals the affinity of the relation where n_i belongs (Equation 1); global importance was defined in Sect. 2.1. A candidate size-l OS is an *optimal* size-l OS, if it has the maximum Im(Sl) over all candidate size-l OSs.

The generation of a size-l OS is a challenging task because we need to select l nodes that are connected to $n^{\rm DS}$ and at the same time result in the maximum score. An optimal dynamic programming algorithm (requiring $O(nl^2)$ time where n is the amount of nodes in the OS) and greedy algorithms were proposed in [15].

3 DSize-l and PSize-l snippets

We propose two types of size-l OSs, namely diverse DSize-l OSs and proportional PSize-l OSs, which extend the size-l OS definition [15] to capture diversity and proportionality, respectively. Similarly to size-l OSs, both DSize-l OSs and PSize-l OSs should be standalone sub-trees of the OS, composed of l important and representative nodes only, so that the user can understand them without any additional information. Thus, the l nodes should form a connected graph that includes the root of the OS (i.e., n^{DS}). We argue that an effective DSize-l (PSize-l) OS should gracefully combine diversity (proportionality) and the local importance scores of constituent nodes. Hence, we propose that for each OS node n_i , we estimate a respective score for diversity, proportionality and local importance, denoted by $dv(n_i)$, $pq(n_i)$ and



Table 1 Notations

Notation	Definition
$gi(v_i)$	The global importance of a graph node v_i
$li(n_i)$	The local importance of an OS node n_i
$fr(v_i)$	The <u>fr</u> equency a graph node v_i appears in an OS
$z(v_i)$	The amount of times that a graph node v_i has been added on the snippet
$dv(n_i)$	The <u>diversity</u> score of an OS node n_i
$dw(n_i)$	The diversity weight score: $li(n_i) * dv(n_i)$
$pq(n_i)$	The proportionality quotient of an OS node n_i
$pw(n_i)$	The proportional weight score: $li(n_i) * pq(n_i)$
$w(n_i)$	A weight score that may represent either $dw(n_i)$ or $pw(n_i)$ (when the context is clear)
$ap(n_i)$	The average $w(.)$ score of nodes of path n_i to root
Im(Sl)	The <u>Importance</u> score of a size- <i>l Sl</i>

 $li(n_i)$, respectively. $dv(n_i)(pq(n_i))$ and $li(n_i)$ are combined to a single score $dw(n_i)(pw(n_i))$ for a DSize-l (PSize-l) OS, simply denoted by $w(n_i)$ when the context (i.e., diversity or proportionality) is clear. The notations frequently used throughout this section and in the rest of the paper are summarized in Table 1.

Then, the objective is to select the l nodes that (1) include n^{DS} and form a connected subtree of the OS and (2) the sum of their w(.) scores is maximized. Local importance (i.e., li(.)) can be calculated as in the original size-l OS problem (i.e., by multiplying affinity with global importance [15]), thus hereby, we discuss only diversity and proportionality.

We investigate two relevance types among OS nodes, namely similarity (denoted as sim) and equality (denoted as equi). More precisely, we consider relevance among nodes belonging to the same relation; thus, we classify each $G^{\rm DS}$ relation either as sim or equi. We can use a domain expert to classify each relation in these types. When two nodes belong to different relations, then they have relevance 0 (e.g., the similarity between a conference and a paper is always 0). For nodes belonging to the same sim relation, we use Jaccard similarity (i.e., $sim(n_i, n_j) = \frac{|n_i \cap n_j|}{|n_i \cup n_j|}$; we treat n_i and n_j as sets of words). We can use an expert to define which attributes to compare per relation. For instance, paper is a sim relation (author G^{DS} , Fig. 2), and we define similarity between papers using their titles (e.g., "On Power-law relationships of the Internet Topology" vs. "Power laws and the AS-Level Internet topology"). (Note that Jaccard similarity is symmetric (i.e., $sim(n_i, n_j) = sim(n_j, n_i)$), and the respective distance (i.e., $1 - sim(n_i, n_j)$) is a metric.) However, we observe that textual similarity cannot be applied on all relations meaningfully. For instance, consider Authors; it is not meaningful to define textual similarity between author names "Christos Faloutsos" vs. "Michalis Faloutsos." Thus, for such cases (e.g., DBLP relations Author and ConfYear), we consider equality relevance, where two different OS nodes may either correspond to the same tuple (e.g., $g(n_i) = g(n_j) = v_p$) or to two different tuples (e.g., $g(n_i) \neq g(n_j)$). In the former case, we have $sim(n_i, n_j) = 1$, whereas in the latter case, we have $sim(n_i, n_j) = 0$. Note that in both relevance types, we measure relevance between two nodes using the same notation, i.e., $sim(n_i, n_j)$ (even for equi relations). Also note that $sim(n_i, n_j) = 1$ indicates that the two nodes are equal even in the case of sim relations; e.g., although conference is a sim relation, an author may have papers appearing in the same conference more than once.

3.1 Diversity (DSize-l OSs)

We suggest that the l nodes should be diversified by preventing the domination of very important nodes. For example, in the Michalis Faloutsos OS, the co-authorship with the very important author Christos Faloutsos dominates the snippet, and this renders the snippet not representative. A natural criterion objective toward measuring diversity is to maximize the sum of dissimilarities between nodes. Thus, for a given graph node n_i in a DSize-l DSl, we suggest to estimate diversity as follows:

$$dv(n_i) = \begin{cases} 1 - \frac{\sum_{n_j \in DSl, n_i \neq n_j} sim(n_i, n_j)}{l - 1} & R(n_i) \text{ is sim relation} \\ 1 - \frac{z(g(n_i)) - 1}{l - 1} & R(n_i) \text{ is equi relation} \end{cases},$$
(2)

where $R(n_i)$ is the relation n_i belongs to, n_j is any other node in DSI, and $sim(n_i, n_j)$ is the similarity between n_i and n_j . When $g(n_i) = g(n_j)$ (i.e., n_i and n_j correspond to the same tuple), then $sim(n_i, n_j) = 1$ for both sim and equi relations. For an equi relation, if $g(n_i) \neq g(n_j)$, then $sim(n_i, n_j) = 0$. Recall also that if n_i and n_j do not belong to the same relations (i.e., $R(n_i) \neq R(n_j)$), then $sim(n_i, n_j) = 0$. The summation of similarities of n_i to the rest of the nodes in the snippet will give us the respective $dv(n_i)$ score. For an equi relation, that will be $z(g(n_i)) - 1$, where $z(g(n_i))$ is the amount of times $g(n_i)$ appears in the snippet (since for all nodes n_j such that $g(n_i) = g(n_j)$, $sim(n_i, n_j) = 1$). Dividing by l - 1, we normalize $dv(n_i)$ in the range [0,1].

Given a set of nodes, $n_{j1} ldots n_{jx}$, that have been added to the DSize-l OS, we denote as $dv(n_i|n_{j1}, \ldots n_{jx})$ the diversity score of an unselected node n_i considering these added nodes. For instance, the score $dv(P_1|P_5)$ in Table 2 denotes the score of P_1 after the addition of P_5 in the snippet. For short, when the context is clear, we also denote as $dv(n_i|n_{jx})$ the score of n_i given that n_{jx} has been appended as the last (i.e., xth) node on the snippet. We also denote as $dv(n_i|\varnothing)$ the maximum diversity score a node n_i can get, i.e.,



 Table 2
 Sim
 Relevance of Selected Papers of Michalis Faloutsos

Name	li(.) dv(.	$dv(. \varnothing)$	$dw(. \varnothing)$	$dv(. P_5)$	$dw(. \varnothing) dv(. P_5) dw(. P_5) dv(. P_8) dw(. P_8)$	$dv(. P_8)$	$dw(. P_8)$	$pq(. \varnothing)$	$pw(. \varnothing)$	$pq(. P_5)$	$pq(. P_5)$ $pw(. P_5)$	$pq(. P_8)$	$pw(. P_8)$
<i>P</i> ₁ : Aggregated Multicast for Scalable QoS Mu 0.17 1.00	0.17	1.00	0.17	1.00	0.17	66.0	0.16	69.0	0.11	69.0	0.11	0.59	0.10
P ₂ : Aggregated Multicast with Inter-Group Tr	0.18 1.00	1.00	0.18	1.00	0.18	66.0	0.18	0.56	0.10	0.56	0.10	0.48	0.09
P_3 : BGP-lens: Patterns and Anomalies in Inte	0.16	0.16 1.00	0.16	0.99	0.16	86.0	0.16	0.58	0.09	0.49	0.08	0.43	0.07
P_4 : Bounds for the On-line Multicast Problem	0.19 1.00	1.00	0.19	1.00	0.19	66.0	0.19	89.0	0.13	89.0	0.13	0.59	0.11
P_5 : On Power-law Relationships of the Intern	0.61 1.00	1.00	0.61	I	I	ı	ı	1.15	0.71	I	ı	I	ı
P_6 : Power Laws and the AS-Level Internet Top 0.19 1.00	0.19	1.00	0.19	0.93	0.17	0.92	0.17	1.15	0.21	0.49	0.09	0.46	80.0
P_7 : QoS-aware Multicast Routing for the Inte	0.18 1.00	1.00	0.18	0.99	0.18	96.0	0.18	1.15	0.21	0.99	0.18	69.0	0.13
<i>P</i> ₈ : QoSMIC: Quality of Service Sensitive Mu	0.31 1.00	1.00	0.31	0.99	0.31	ı	ı	0.93	0.29	0.79	0.24	ı	ı
P_9 : Reducing Large Internet Topologies for F.	0.27 1.00	1.00	0.27	0.98	0.26	0.97	0.26	69.0	0.19	0.48	0.13	0.43	0.12
P_{10} : The Effect of Asymmetry on the On-Line 0.17 1.00	0.17	1.00	0.17	1.00	0.17	66.0	0.16	98.0	0.14	98.0	0.14	0.74	0.12

 $dv(n_i|\varnothing) = 1$, e.g., when n_i is the first to be added. This notation will be useful when describing our greedy algorithms, where after each node addition, the score of unselected nodes is affected accordingly.

For equi nodes for short, we also denote as $dv[z](g(n_i))$ the diversity score of a graph node n_i considering it appears for the zth time in the snippet. For instance, in Table 3, dv[1] indicates the score of a node assuming it appears for the first time, where $dv[1](n_i) = 1$ is the maximum diversity score (which corresponds to $dv(n_i|\varnothing) = 1$). As another example, consider l = 10 and that C. Faloutsos appears two times (i.e., z = 2); $dv[2] = 1 - \frac{2-1}{10-1} = \frac{8}{9} = 0.89$ (Table 3). Note that this score corresponds to the graph node $g(n_i)$; thus, both nodes will have the same dv(.), i.e 0.89 (an alterative way would be to score the first occurrence as 1 and the second as 0.78, since 1 + 0.78 = 0.89 + 0.89).

Our equation is inspired by (1) max-sum diversification that maximizes the sum of the relevance and dissimilarity of a set and by (2) the use of a mono-objective formulation, which, similarly to our equation, combines relevance and dissimilarity to a single value for each document [18]. Note that, in general, diversification approaches trade-off (1) the similarity of results with the given query and (2) the dissimilarity among such results using a similarity measure (e.g., IR techniques). For instance, given a query "Internet Topology," papers "On Power-law relationships of the Internet Topology" and "Power laws and the AS-Level Internet topology" have some similarity to this query, but they also have some similarity among them, both types can be estimated using a common IR measures such as Jaccard similarity. This is not the case here, since we do not consider the similarity of nodes to the query, but a local importance score in relation to n^{DS} . Thus, local importance and similarity are not meaningfully comparable. Note also that their respective values may not be in the same range (e.g., local importance may range in [0,10], whereas dv(.) always ranges in [0,1]). Hence, unlike most diversification approaches, in the combining function dw(.) (to be defined in Sect. 3.3), we do not sum up local importance and dissimilarity, but multiply them.

3.2 Proportionality (PSize-l OSs)

We observe that in an OS, we often find equi graph nodes (i.e., database tuples) multiple times. For instance, in the Michalis Faloutsos OS (see Table 3), we have 37 instances of S. Krishnamurthy, 12 instances of C. Faloutsos, 18 papers in INFOCOM. We denote the frequency of a graph node v_i in an OS as $fr(v_i)$ (or simply by fr when the context is clear). Graph nodes appear in an OS multiple times could sometimes be comparatively weak in terms of importance, but still given their frequency in the OS, they should be represented analogously in an effective snippet. Thus, we suggest that in a PSize-l snippet, disregarding local importance (i.e., assum-



Name	li(.)	<i>fr</i> (.)	dv[1](.)	dw[1](.)	dv[2](.)	dw[2](.)	pq[1](.)	pw[1](.)	pq[2](.)	pw[2](.)
Srikanth V. Krishnamurthy	0.60	37	1.00	0.60	0.89	0.53	12.33	7.40	7.40	4.44
Christos Faloutsos	1.80	12	1.00	1.80	0.89	1.60	4.00	7.20	2.40	4.32
Jun-Hong Cui	0.81	11	1.00	0.81	0.89	0.72	3.67	2.97	2.20	1.78
Thomas Karagiannis	0.70	10	1.00	0.70	0.89	0.62	3.33	2.33	2.00	1.40
Michael Mitzenmacher	1.40	3	1.00	1.40	0.89	1.24	1.00	1.40	0.60	0.84
George Varghese	1.38	2	1.00	1.38	0.89	1.23	0.67	0.92	0.40	0.55
Konstantina Papagiannaki	0.61	4	1.00	0.61	0.89	0.54	1.33	0.81	0.80	0.49
Samuel Madden	1.61	1	1.00	1.61	0.89	1.43	0.33	0.54	0.20	0.32
Marek Chrobak	0.33	4	1.00	0.33	0.89	0.29	1.33	0.44	0.80	0.27
Jakob Eriksson	0.15	7	1.00	0.15	0.89	0.13	2.33	0.35	1.40	0.21

Table 3 Equi relevance of selected co-authors of Michalis (Ranked descending their pw[1])

ing that all nodes have the same li(.), we should include nodes in proportion of their frequency. For example, if a graph node v_i appears 37 times in the total of 1,259 OS nodes, then v_i should ideally appear $\lfloor l \cdot 37/1, 259 \rfloor$ times in the respective PSize-l OS (note that this may not practically possible as in-between nodes may also be required, i.e., the co-authored papers in our example).

Analogously, we observe that the topic of sim nodes may appear multiple times; a node may be very similar to many other nodes in the OS. For instance, in Table 2, we find six out of ten papers including the word "Multicast" (e.g., P_1 , P_2 , P_4) and two papers including a pair of words "Aggregated Multicast." Thus, papers have some similarity due to the frequent common topics (e.g., "Aggregated Multicast"), and hence, they should also be analogously represented.

For this purpose, for a given graph node n_i in a PSizel PSI, we propose the use of the *proportional quotient* as follows:

$$pq(n_i) = \begin{cases} \frac{\sum_{n_j \in OS} sim(n_i, n_j)}{\alpha \cdot \sum_{n_j \in PSl} sim(n_i, n_j) + 1} & R(n_i) \text{ is sim relation} \\ \frac{fr(g(n_i))}{\alpha \cdot z(g(n_i)) + 1} & R(n_i) \text{ is equi relation} \end{cases}$$

(3)where $R(n_i)$ is the relation where n_i belongs, $sim(n_i, n_i)$

where $R(n_i)$ is the relation where n_i belongs, $sim(n_i, n_j)$ is the similarity between the two nodes (as defined in Equation 2), and α is a constant that can tune proportionality. For equi relations, $z(g(n_i))$ is the amount of times that node n_i appears in the snippet and $fr(g(n_i))$ is the frequency that the node appears in the OS. We use analogous notations as in $dv(n_i)$. We denote by $pq(n_i|n_{j1}, \dots n_{jx})$ the proportional quotient of n_i when nodes $n_{j1}, \dots n_{jx}$ have been appended to the snippet. For equi nodes, we also denote as $pq[z](n_i)$ the proportional score considering n_i appears z times (Table 3).

This equation is inspired by the Sainte-Laguë Algorithm [8] (with $\alpha=2$), and empirically we found that it is very effective for our problem (other equations can also be con-

sidered, e.g., [7,29]). The rationale of this quotient is to favor a frequent node (or nodes including frequent topical words), and each time a node is added to the snippet, its proportional score is significantly decayed so that other frequent nodes will be selected, in turn. This way, diversification is also facilitated. For instance, considering fr = 12 and $\alpha = 2$ for C. Faloutsos, by adding this node once we get $pq[1](n_i) = 12/3 = 4$ and twice we get $pq[2](n_i) = 12/5 = 2.4$.

3.3 DSize-l and PSize-l definitions

Based on the above discussion, for DSize-*l* OSs, we propose the following combining score per node:

$$dw(n_i) = li(n_i) \cdot dv(n_i), \tag{4}$$

where $li(n_i) = af(n_i) \cdot gi(n_i)$ is the local importance of n_i and $dv(n_i)$ is the diversity factor (Equation 2). Tables 2 and 3 depict examples of how these scores can be obtained by constituent scores for l=10. For instance, consider the simplified example where we need to select five authors (and thus an intermediary paper), then we will select twice C. Faloutsos (i.e., $0.89 \cdot 1.8 + 0.89 \cdot 1.8 = 1 \cdot 1.8 + 0.78 \cdot 1.8$) and once S. Madden (1 · 1.6), M.Mitzenmacher (1 · 1.4) and G.Varghese (1 · 1.4). Note that a third addition of C. Faloutsos cannot compete the total 1.4 score, as the additional score is only 1.12 (i.e., $(3 \cdot 0.78 - 1 - 0.78) \cdot 1.8 = 0.56 \cdot 1.8 = 1.08$).

Definition 1 (DSize-l **OS**) Given an OS and l, a DSize-l OS is a subset of OS that satisfies the following:

- (1) The size in nodes of DSize-l OS is l (where $l \leq |OS|$)
- (2) The l nodes form a connected tree rooted at n^{DS}
- (3) Each node n_i carries a weight $dw(n_i)$ (obtained by Equation 4)
- (4) The aggregated score of a DSize-*l* OS *DSl* can be calculated by:

$$Im(DSl) = \sum_{n_i \in DSl} dw(n_i).$$
 (5)



Let a candidate DSize-l OS be any OS subset satisfying conditions 1-3; then, the optimal DSize-l OS is the candidate snippet that has the maximum Im(DSl) among all such candidates.

Problem 1 (*Find an optimal DSize-l* **OS**) Given an OS and *l*, find a candidate DSize-*l* OS of maximum score (according to Definition 1).

Analogously, we define the proportionality score per node (i.e., $pw(n_i) = li(n_i) \cdot pq(n_i)$, instead of Equation 4), PSize-l OS and the optimal PSize-l OS problem (a formal definition is omitted due to the interest of space). For instance, we observe that our selection policy will favor first the addition of S. Krishnamurthyan author and the respective co-authored paper; then, the addition of author C. Faloutsos with a co-authored paper; then, another round with these two authors, etc.

3.3.1 Problem definitions and algorithms

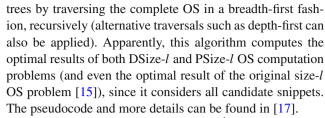
We have two types of problems, namely PSize-*l* and DSize-*l* generation. In addition, an OS may have only *equi* relevance or both *equi* and *sim* relevance. Thus, we have four combinations of problems, and thus, we propose algorithms that are general to address all these combinations. Firstly, we propose a brute-force algorithm which is prohibitively slow. Then, we propose two greedy algorithms LASP and 2-LASP*e* (which is LASP's optimization). Finally, we propose pruning algorithms that can produce pruned preliminary results. We can apply all aforementioned algorithms on these preliminary OSs.

3.3.2 Notation

For simplicity, we unify dw(.) and pw(.) into a single notation w(.) and use $w(n_i)$ to refer to the corresponding diversity or proportionality score of a node n_i in a DSize-l OS or PSize-l OS, respectively. Analogously to diversification and proportionality scores notations, we denote $w(n_i|n_{j1},\ldots n_{jx})$ as the score given $n_{j1},\ldots n_{jx}$ have been added and $w[z](n_i)$ as the score when n_i is added for the zth time. In the rest of the paper, whenever the context is clear, we drop n_i or $(n_i|n_{j1},\ldots n_{jx})$ from the notation and denote the diversity/proportionality score of a node simply by w(.).

4 Brute-force (BF-*l*) algorithm

A brute-force (BF-*l*) algorithm for computing the optimal DSize-*l* (or PSize-*l*) OS would consider all candidate size-*l* trees, compute the respective scores, and eventually find the optimal solution. BF-*l* generates all possible candidate



The time complexity of BF-*l* is $O(\frac{n!}{(n-l)!l!})$, where *n* is the number of nodes in OS and *l* is the required size. Note that, an application of a dynamic programming algorithm [15] that could reduce this cost is not possible here, because the score of a diversified size-l OS (either DSize-l or PSize-l) is not distributive w.r.t. the scores of the subtrees it is composed of; the reason is that the two subtrees that contain one or more common tuples are not independent (i.e., they affect each others' score due to the diversity components of the scoring formulae). In other words, given an OS tree T that can be decomposed to subtrees T_1 , T_2 , etc., the optimal diversified size-l OS is not necessarily composed by some optimal diversified size-l' OSs (l' < l) of the subtrees T_1, T_2 , etc. (which is the case for size-l OSs [15]). This is because the computation of an optimal diversified size-l' OS in a subtree T_i disregards the diversity of its constituent nodes w.r.t. the snippets chosen from the other subtrees $T_i \neq T_i$. If we were to consider all possibilities of graph node frequencies in the snippets of other subtrees, this would require exponential space.

5 Largest averaged score path (LASP)

The BF-l algorithm can be very expensive even for moderate values of l or |OS|. Thus, we propose LASP, a greedy algorithm, that can produce a size-l OS of high quality at a much lower cost. In a nutshell, LASP firstly generates the OS. It also calculates for each node n_i an initial $w(n_i)$ score (i.e., $w(n_i|\varnothing)$), using Eq. 4) and its corresponding average $w(n_i)$ score per node (denoted as $ap(n_i)$) of the path from n_i to the root. Then, the algorithm iteratively selects and adds to the size-l OS the path p_i of nodes with the largest ap(.). The rationale behind selecting paths instead of single nodes with the largest score is that we can include nodes of very large importance while their ancestors have less importance as their score is averaged. Algorithm 1 is a pseudocode of the heuristic, and Fig. 3 illustrates an example.

LASP is a general algorithm that (1) can compute both types of size-*l* OSs (i.e., DSize-*l* and PSize-*l* OSs) and (2) can process both relevance types (i.e., equality and similarity). The difference between the two size-*l* types is that the proportionality equation also considers the similarity/equality (frequency) of each node against all other nodes, which is calculated during the OS generation process (to be described in more detail shortly). LASP can process both relevance types by using the pre-calculated *sim matrix* (a matrix stor-



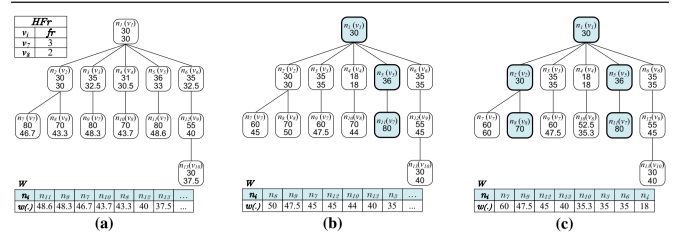


Fig. 3 LASP algorithm: the size-5 OS (annotated with OS and (graph) node ID, w(.) and ap(.); selected nodes are shaded). a The initial OS. b First update. c The final update

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Algorithm 1 Largest Averaged Path Algorithm
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LASP(l, n^{DS})
1: OS Generation (n^{DS}) \triangleright generates OS, w(.), ap(.), HFr and W
2: while (|\text{size-}l| < l) do
     p_i=path from maximum W node
4:
     add first (l - |\text{size-}l|) nodes of p_i to size-l
5:
     if (|\text{size-}l| < l) then
        remove selected path p_i from the OS tree and from W
6:
7.
        UpdateRemPaths (p_i)
        UpdateRelScores (p_i)
9: return size-l
UpdateRemPaths (p_i)
1: for each child v of nodes in p_i do
2:
     for each node n_i in the subtree rooted at v do
3:
        update ap(n_i) on the OS tree
4:
        update ap(n_i) on W
UpdateRelScores (p_i)
1: for each node n_i in p_i do
     for each unselected OS node n_i do
2:
3:
        if (sim(n_i, n_i) > 0) then
4:
          update w(n_i) considering sim(n_i, n_i)
5:
          for each node n_k in the subtree rooted at n_i do
6:
             update ap(n_k) on OS tree using w(n_i)
```

ing the similarity among all nodes), which facilitates the initial calculation of w(.) and ap(.), and the consequent updates (to be described in more detail shortly). Thus, given an OS annotated with w(.) and ap(.) scores, the problem of determining either DSize-l or PSize-l using either equality or similarity relevance type remains the same for LASP.

update $ap(n_k)$ on W

7:

More specifically, the LASP algorithm firstly generates the OS (line 1). During **OS Generation**(), LASP also calculates the w(.) score and the respective ap(.) score per node. For the DSize-l, the calculation of dv(.) (and thus w(.)) is straightforward, whereas for the PSize-l, the calculation of

pq(.) is more demanding as it requires the comparison of each node against all other nodes. Thus, for equi relations, in order to facilitate faster calculation of pq(.) scores, we also maintain a hash table of graph nodes (denoted as HFr) containing the frequency of a graph node in the OS tree (denoted as fr). HFr can easily be compressed by excluding nodes appearing only once in the OS; thus, if a node does not exist in HFr, we can infer that it only appears once. For sim relations, we compare each node against all other nodes as to obtain their pq(.). Note that sim comparisons are more expensive (i.e., $\frac{(n+1)\cdot n}{2}$ time) in contrast to equi HFr-based comparisons that require only n time. During the OS generation, we also generate a priority queue W of the initial ap(.), in order to better manage nodes.

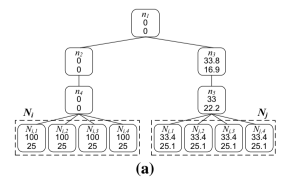
We then select the node with the largest ap(.) and add the corresponding path to the size-l OS. We remove this p_i from the OS and from W (lines 6). By removing the nodes of p_i from the OS, the tree now becomes a forest; each child of a node in p_i is the root of a tree. Accordingly, the ap(.) of affected nodes is updated (1) to disregard the removed nodes in the path (UpdateRemPaths()) (2) and also to consider the revised w(.)s due to relevance to newly added nodes (**UpdateRelScores**()). Note that the ap(.) of an unselected node corresponds to the w(.) score the node will have if included in the snippet (considering also the similarity loss of already added nodes for the diversity case, i.e., dv). Thus, w(.) and ap(.) scores of all unselected nodes should be updated each time a new node is added (by function **UpdateRelScore**()). Also note that this general LASP can be significantly more expensive than the version of LASP presented in [17], since for the latter, it suffices to simply count fr and z of added nodes for the estimation of ap(.) and w(.)scores.

This process (i.e., the selection of the path with the largest ap(.), the addition to the size-l OS, and the required updates)



continues iteratively as long as the selected nodes are less than l. If less than $|p_i|$ nodes are needed to complete the size-l OS, only the top nodes of the path are added to the size-l OS (because only these nodes are connected to the current size-l OS). Note that each time a path is selected, it is guaranteed to be connected with the previously selected paths (as the root of the selected path should be a child of a previously selected path); therefore, the selected paths will form a valid size-l OS.

Take for instance the example of Fig. 3, where nodes at level one have similarity relevance and nodes at level two have equality relevance. More precisely, consider sim $(n_3, n_4) = sim(n_4, n_5) = 0.6$, whereas equality relevance is annotated on the example of figure (e.g., $g(n_7) = g(n_9) =$ $g(n_{11}) = v_7$). Node n_{11} (i.e., graph node v_7) has $ap(n_{11}) =$ 48.6, because its path includes nodes n_1 , n_5 and n_{11} with average w(.) being (30+36+80)/3 = 48.6. Assuming that l=5, at the first iteration, the algorithm selects and appends to size-l OS the path comprising nodes n_1 , n_5 and n_{11} with the largest ap(.), i.e., 48.6. For the remaining nodes, w(.) and ap(.) are updated to disregard the removed nodes and also to consider the inclusion of newly added nodes (Fig. 3b). For instance, the revised $ap(n_{12})$ is (35 + 55)/2 = 45, because its path now includes only n_6 and n_{12} . Also, nodes n_7 and n_9 which correspond to the same graph node as n_{11} and node n_4 which has similarity with node n_5 which have just been added to the size-l also need to be updated with new w(.) and ap(.) scores. In general, if such nodes have descendants, then their descendants should also be updated because both their ap(.)s and w(.) are affected. The next path to be selected is that ending at n_8 , which adds two more nodes to the size-lOS (Fig. 3c). Note that $ap(n_i)$ for each node n_i corresponds to the path starting from n_i to the root of the corresponding unselected tree (from the unselected forest). For instance, during the second update, p_8 comprises n_2 and n_8 . Note also that the path's root (e.g., n_2) is always the child of a node (in the OS) which already exists in the current size-l OS, e.g., n_1 in this case. Thus, each time we select a path to append to the size-l OS, we always get a valid OS.



5.1 Analysis

The time complexity of the algorithm is $O(nl \log(n))$, where n is the size of the complete OS, as at each step the algorithm may choose only one node which causes the update of O(n) paths twice (firstly for the path size update and secondly for the z updates). Each update costs log(n) time using the priority queue W. In terms of approximation quality, this algorithm empirically produces very good results. Hereby, we prove the approximation lower bound and cases where this algorithm will return the optimal DSize-l and PSize-l OSs.

5.1.1 Lower bound of LASP approximation

Theorem 1 LASP is a d-approximation algorithm, where d is the maximum depth of the G^{DS} tree. Namely, the ratio of the optimal Im(.) (denoted as OPT) over the Im(.) of the solution by LASP (denoted also as LASP) is at most d.

Proof We analyze the worst case for LASP, by defining two rival sets of nodes N_i and N_j . As we will show in the sequel, our definitions of N_i and N_j sets correspond to the worst case for LASP, which maximizes the Im(.) ratio between the optimal result (by including nodes from N_i) and the approximate solution by LASP (by including nodes from N_j).

Let N_i be a set of sibling nodes in the OS tree, which are at depth d_i . Let $N_{i,p}$ be the pth node in N_i . Moreover, (i) $|N_i| \ge l - d_i$, i.e., the amount of nodes in N_i is at least $l - d_i$ and (ii) $sim(N_{i,p}, N_{i,q}) = 0$ for all p, q, i.e., none of N_i nodes have any similarity. Let us also assume that all nodes in N_i have the same $w(N_{i,p})$ score (denoted as $w(N_i)$ for simplicity). Let ap(.)[z] be the ap(.) of a node after z paths additions (iterations).

In addition, let N_j be also a set of nodes, where $|N_j| \ge l$, with a common $w(N_j)$ for all nodes and a common $ap(N_j)[z]$ for all nodes such that $ap(N_j)[z] = w(n_j)$ for a $z \ge 1$. Again, we assume that none of these nodes are similar. An example of N_j set will be the analogy of the N_i set, i.e., a set of siblings at depth d_j , where $|N_j| \ge l - d_j$, with a common $w(N_j)$ for all nodes (Fig. 4). Another example of a

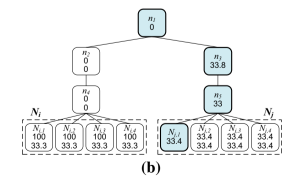


Fig. 4 LASP: example of rival N_i and N_j sets (annotated with node ID, w(.) and ap(.)[z]; selected nodes are *shaded*). **a** The initial OS. **b** First update



valid N_j would be as follows: The $N_{j,1}$ is as defined in Fig. 4 (i.e., at path n_1 , n_3 , n_5 , $N_{j,1}$), whereas $N_{j,2}$, ..., $N_{j,l}$ are children nodes of the root n_1 instead (since after the first addition of $N_{j,1}$, the $ap(N_j)[z]$ of $N_{j,2}$, ..., $N_{j,l}$ becomes $w(n_j)$).

When, (1) $ap(N_i)[z] > ap(N_i)[z]$ for any z, (2) $w(N_i) >$ $ap(N_i)[z]$ for $z \ge 1$ and (3) $w(N_i) > w(N_i)$, the above assumptions dictate that LASP would wrongly prefer to iteratively select nodes from N_i over nodes from N_i . As illustrated by Fig. 4, there are two states of the first condition: the initial state before any updates, i.e., z = 0 and the state after the first update, i.e., $z \ge 1$. The latter is after the first selection of an N_i node where its ancestors are removed from the OS, leading to new aps, i.e., $ap(N_i)[z] = w(N_i)/d_i$ (to be explained shortly) and $ap(N_i)[z] = w(N_i)$, which leads to the second condition: $w(N_i) > ap(N_i)[z]$. The bigger the difference between $w(N_i)$ and $w(N_i)$, the bigger the approximation loss of LASP over the optimal solution, because N_i nodes will repeatedly be selected (due to $w(N_i) > ap(N_i)[z]$) over N_i nodes. To see this in an example, consider the OS tree of Fig. 4, where each node includes its w(.) score and the corresponding ap(.)[z] score for z = 0 and 1 (where $d_i = d_i = 3$). All nodes in group N_i (four leftmost leaf nodes) have w(.) scores 100, all nodes in the path from the OS root to the parent of N_i have 0 w(.) scores, and all nodes in group N_i (four rightmost leaf nodes) have w(.) scores 33.4. $ap(N_i)[z]$ are always marginally larger than the respective $ap(N_i)[z]$ for all z. E.g., initially, we have $ap(N_i)[0] = 25.1$ and $ap(N_i)[0] = 25$, and after the first addition of a path to $N_{i,1}$, we have $ap(N_i)[z] = 33.4$ and $ap(N_i)[z] = 33.3$ for $z \ge 1$. In this example, LASP will iteratively select all N_i nodes before it can pick any N_i node. On the other hand, the optimal solution would include $l - d_i N_i$ nodes. Thus, if $|N_i| \ge l - d_i$, the approximation loss of LASP will be $(PPar_i + (l - d_i) \cdot w(N_i)) - (PPar_j + (l - d_j) \cdot w(N_i)),$ where $PPar_i$ and $PPar_i$ are the sum of w(.) scores at the nodes along the path from the OS root to the parent of N_i and N_i , respectively.

Sets N_i and N_j are rivals, since they compete in the selection and LASP wrongly chooses nodes from N_j over nodes from N_i , resulting to an approximation loss. We say that two such sets are not valid rivals if they do not meet the conditions: (1) $ap(N_j)[z] > ap(N_i)[z]$, (2) $w(N_j) > ap(N_i)[z]$ and (3) $w(N_i) > w(N_j)$ for any z.

Thus, considering an OS contains two rival sets N_i and N_j , we can estimate the approximation ratio of LASP compared to OPT as follows:

$$\frac{OPT}{LASP} = \frac{PPar_i + (l - d_i) \cdot w(N_i)}{PPar_j + (l - d_j) \cdot w(N_j)}$$
 (6)

Lemma 1 For a fixed $ap(N_i)[0]$ and $l > d_i$, the summation of the scores of the nodes of a snippet that includes a subset

of N_i (i.e., $Im(DSl(N_i)) = PPar_i + (l - d_i) \cdot w(N_i)$) is maximized when $PPar_i = 0$.

Proof For a fixed $ap(N_i)[0]$, obviously $w(N_i)$ is maximized when $PPar_i = 0$, as $ap(N_i)[0] = (PPar_i + w(N_i))/d_i$ and $ap(N_i)[1] = w(N_i)$. Hence, $Im(DSl(N_i)) = PPar_i + (l - d_i) \cdot w(N_i)$ is also maximized when $PPar_i = 0$ and $l > d_i$, for a fixed $ap(N_i)[0]$.

For instance, for $ap(N_i)[0] = 25$, $d_i = 3$ and l = 6, consider the two N_{i1} and N_{i2} cases with $PPar_{i1}$: $ap(N_{i1}) = (25 + 25 + 25 + 25)/4 = 25$ and $PPar_{i2}$: $ap(N_{i2}) = (0 + 0 + 0 + 100)/4 = 25$. Thus, we have $w(N_i)$ scores 25 and 100 and holistic scores $Im(DSl(N_{i1})) = 6 \cdot 25 = 150$ and $Im(DSl(N_{i2})) = 0 + 0 + 0 + 100 + 100 + 100 = 300$, respectively (i.e., $Im(DSl(N_{i1})) < Im(DSl(N_{i2}))$).

According to Lemma 1, for a fixed $ap(N_i)[0]$, the total score $Im(DSl(N_i))$ due to nodes from N_i is maximized when $PPar_i = 0$. In this case, $ap(N_i)[z] = w(N_i)/(d_i + 1)$ for z = 0 and $ap(N_i)[z] = w(N_i)/(d_i)$ for $z \ge 1$. As mentioned above, for LASP to repeatedly select nodes from N_i , we should have $ap(N_i)[z] > ap(N_i)[z]$, (i.e., $ap(N_i)[z] >$ $w(N_i)/(d_i+1)$ for z=0 and $ap(N_i)[z]>w(N_i)/(d_i)$ for $z \ge 1$) and $w(N_i) > ap(N_i)[z]$ (i.e., $w(N_i) > w(N_i)/(d_i + d_i)$ 1) for z = 0 and $w(N_i) > w(N_i)/d_i$ for $z \ge 1$). Following the same reasoning as Lemma 1, the contribution of the N_i nodes in the score of the snippet computed by LASP is minimized when $w(N_i)$ is minimized (since $PPar_i$ is considered only once for all chosen N_i nodes). Thus, as to minimize the holistic score of the LASP computed snippet over the optimal score, we should minimize $w(N_i)$ subject to $w(N_i) > w(N_i)/d_i$ (i.e., $w(N_i) > ap(N_i)[z]$ for any z); thus, set $w(N_i) \approx w(N_i) \cdot d_i$. This corresponds to the worst case for LASP. Now, in order to compute an upper bound for the optimality ratio OPT/LASP shown in the fraction above, we (i) set $PPar_i = 0$, as discussed in Lemma 1, (ii) set $PPar_i = 0$, as this can only increase the ratio, (ii) set $d_i = d_i$, as $l - d_i \ge l - d_i$, i.e., $d_i \le d_i$ (in order to select at least the same number of nodes from N_j as from N_i) and setting $d_i = d_i$ minimizes the denominator. Thus, we get:

$$\frac{OPT}{LASP} \le \frac{0 + (l - d_i) \cdot w(N_j) \cdot (d_i)}{0 + (l - d_i) \cdot w(N_j)} \approx d_i \tag{7}$$

Obviously, the above ratio is maximized for $d_i = d$, i.e., N_i is at the maximum depth of the OS tree; thus, d is the worst-case ratio between the qualities of the snippet computed by LASP compared with the optimal snippet. Note that in all our case studies, $d \le 3$.

5.1.1.1 Discussion on Rival N_i and N_j Sets We now show that the case of rival sets N_i and N_j is the worst case for LASP and that the relaxation of the assumptions about these rival sets can only be in favor of LASP, thereby establishing the d optimality ratio.



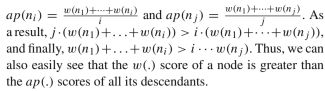
When the rival sets have at least $l-d_i$ (resp. $l-d_i$) siblings with common w(.) scores, this will result in the worst case. In our proof, we assume that the two rival sets consist of at least $l - d_i$ (resp. $l - d_i$) siblings carrying the same score. This is because only such sets can maximize the maximal difference between two sets. Recall that N_i and N_j represent the optimal and worst case, respectively. For instance, let us assume that we have two N_i sets N_{i1} and N_{i2} consisting of l/2nodes carrying equal score (i.e., not all N_i nodes are siblings) corresponding to two respective $PPar_{i1}$ and $PPar_{i2}$. Then, the optimal solution will include the two respective sub-trees where the additional $PPar_{i2}$ path will be an overhead we did not have in the original case. Similarly, if we assume we have only one set N_i where its nodes do not have the same score, then this may dictate the selection of nodes from other sub-trees (similarly to the case with two N_i sets), and we will end up again with an additional $PPar_{i2}$ overhead. Analogously, we can show that if the N_i nodes have different parents belonging to different paths, such that $ap(N_i)[z] <$ $w(n_i)$ for a $z \ge 1$, this can only improve the score of LASP's solution and thus reduce its approximation loss.

Rival sets should consist of diverse graph nodes (i.e., impact of Diversity and Proportionality). Let us assume the case where some nodes in the rival N_i or N_j sets correspond to the same graph node, e.g., $g(N_{j,p}) = g(N_{j,q})$. The inclusion of a node (from either set N_i or N_j) will still have no impact on the proved lower bound. The reason is that the addition of a node in the result can only result in the possible reduction of the w(.) scores of the unselected nodes and the respective ap(.) scores. This reduction will disqualify them against unselected unique nodes which will retain the same $ap(N_i)$ and $w(N_i)$ scores (resp. $ap(N_j)$ and $w(N_j)$ scores). Since, we can always construct a worst case with unique $(l-d_i)$ -sized subsets of N_i and N_j , the upper bound of the approximation loss by LASP remains unchanged.

5.1.2 Optimality of LASP

Theorem 2 For equi relevance, if the nodes of an OS have monotonically decreasing initial w(.) ($w(n_i|\varnothing)$) scores with respect to their distance from the root (i.e., when the score of each parent is not smaller than that of its children), then LASP returns the optimal equi PSize-l OS or equi DSize-l OS. We denote such an OS as monotonic OS(w).

Proof The optimal PSize-l or DSize-l should include the l nodes with the largest possible w(.) score. Thus, we need to prove that our algorithm can achieve this goal when monotonicity on the w(.) scores hold. Firstly, we show that since monotonicity holds on w(.), the respective ap(.) scores should also be monotonic. For instance, consider the path n_1, n_2, \ldots, n_k with $w(n_1) > w(n_2) > \cdots > w(n_k)$. Then, $ap(n_1) > ap(n_2) > \cdots > ap(n_k)$ since for i < j we have



Note that Equation 4 is monotonic for both size-*l* types to the repetitions of the same graph nodes; i.e., $w(n_i)[1] >$ $w(n_i)$ [2]. Also for equi relevance, we maintain final monotonicity of descendants after repetitions. E.g., consider sibling nodes, n_1 and n_2 , which correspond to the same graph node (i.e., $sim(n_1, n_2) = 1$). Note that, since these two nodes are equal, they have common descendants (sub-trees). E.g., descendants $dn_{1,1}, \ldots, dn_{1,l}$ and $dn_{2,1}, \ldots, dn_{2,l}$, respectively, where $sim(dn_{1,i}, dn_{2,i}) = 1$ for all i. Thus, if we add n_1 first, this will result in the reduction of $w(n_2)$ score which can lead to two cases: In the first case, $w(n_2)$ score is still larger than all its descendants, and thus, we still have monotonicity on w(.) of the tree. In the second case, the new score $w(n_2)$ is smaller than its descendant's score, e.g., $w(n_2) < w(dn_{2,i})$ which also implies that $w(n_2) <$ $w(dn_{1i})$. In that case, a child node of n_1 will be selected since it has larger w(.) than n_2 ; e.g., $w(dn_{1,i})$. Then, the respective descendant of n_2 will be reduced $(w(dn_{2,i}))$ which should give $w(n_2) > w(dn_{2,i})$. This way, LASP will still include nodes with monotonicity on w(.). (Note that for sim relevance, the property of descendant's equality may not hold, and thus, we no longer have monotonicity on w(.), and thus, LASP will not provide the optimal solution.)

Thus, this algorithm will always select the unselected node with the maximum w(.) score which is always a child of an already selected node (i.e., a root of a tree of the unselected forest). Note that because of the monotonicity properties mentioned above, only child nodes of already selected nodes can have the largest score. Thus, progressing iteratively, the l nodes will comprise the optimal PSize-l or DSize-l OSs since this set of l nodes will give the maximum score.

Theorem 3 For equi relevance, if the nodes of an OS have monotonically decreasing local importance (li(.)) to their distance from the root (we denote such an OS as monotonic OS(li)), then LASP returns the optimal equi DSize-l OS.

Proof The dv(.) equation (in contrast to pq(.)) is monotonic to li(.). That is $dv(n_i)[z] \leq li(n_i)$ for any z. Thus, if an OS is monotonic w.r.t. li(.), it will also be monotonic w.r.t. w(.). Thus, based on Theorem 2, the algorithm will give the optimal result.

Theorem 4 For equi relevance, and when we have a monotonic OS(li) and all nodes have $fr \leq \alpha + 1$, then LASP returns the optimal equi PSize-l OS.

Proof The pq(.) equation (in contrast to dv(.)) is not always monotonic to li(.). It is only monotonic when $fr \le \alpha + 1$



that gives a w(.) in [0, 1], and thus, for these frequencies, $pq[z] \le li$. Thus, since the OS is monotonic w.r.t. li(.), it is also monotonic w.r.t. pq(.). Thus, based on Theorem 2, the algorithm will compute the optimal result also in this case.

6 2-LASPe

The runtime cost of LASP is dominated by the numerous updates it applies; each time we add a node (or path) to the snippet, we have to update up to twice each of the remaining nodes. Thus, we introduce the 2-LASPe algorithm, an enhancement of LASP, that aims to reduce where possible such updates. In a nutshell, 2-LASPe facilitates update reductions at both **UpdateRemPaths()** and **UpdateRelScores()** phases. The algorithm remains, like LASP, general and can address both types of size-l and relevance. Algorithm 2 illustrates the differences of the two functions from the respective original functions of the LASP algorithm, whereas Fig. 5 illustrates an example.

We propose to relax LASP by averaging w(.) pairs of nodes (hence the prefix 2 to the name of the algorithm). Namely, we take the average between the current node and the parent instead of the whole path from the current node to the root. As a consequence of this relaxation, updates will be required only on the affected pairs rather on the whole path to the root. Recall that the rationale of considering the average from each node to the root in LASP was to exploit nodes lower on the tree with larger scores than their ancestors. We observe that because of the proposed equations, we expect recurrent monotonicity in the OSs; i.e., recurrent cases where the w(.) of the parent is bigger than that of its child. Recall that $li(.) = af(.) \cdot gi(.)$ where af(.) is monotonic by definition, and additionally, both dv(.) and pq(.) equations are monotonic to z (and additions of similar nodes), where z is expected to be larger at the bottom levels of the OS tree.

Algorithm 2 2-LASPe Algorithm

 $2-LASPe(l, n^{DS})$

UpdateRemPaths (p_i)

- 1: for each child n_i of nodes in p_i do
- 2: update $ap(n_j)$ on the OS tree
- 3: update $ap(n_i)$ on W

UpdateRelScores (p_i)

5:

- 1: for each node n_i in p_i do
- 2: **for each** similarity edge $se(n_i, n_j)$ (of n_i with an unselected n_j node) **do**
- 3: update $w(n_i)$ considering $sim(n_i, n_i)$
- 4: deleteEdge($se(n_i, n_i)$)
 - **for each** child n_k of n_j **do**
- 6: update $ap(n_k)$ on OS tree using $w(n_i)$
 - update $ap(n_k)$ on W

Secondly, we introduce a similarity index on the OS tree, denoted as sim OS tree (see Fig. 5) (instead of using the HFr of LASP and 2-LASP [17]). This sim OS tree is generated during the **OS Generation**() function (line 1). For each pair of nodes that there exists a similarity, we add a similarity edge carrying the similarity value, denoted as $se(n_i, n_j)$. Using the similarity edges, we can limit checks of newly added nodes against only unselected nodes that we know they have a similarity (thus the suffix e for edge to the name of the algorithm). This is in contrast to LASP which checks against all unselected nodes.

Let us first demonstrate the updates due to removals of paths. At each addition, we update only pairs of scores, i.e., ap(.) between the affected node and its parent (instead of all remaining paths toward the root). For example, after adding path p_{11} , in 2-LASPe, we only need to update nodes at level 1, except the included n_5 node (since node n_1 is removed). Let us now demonstrate the updates due to relevance between the selected path and unselected nodes. We can easily determine from the sim OS tree that p_{11} path's nodes have similarity

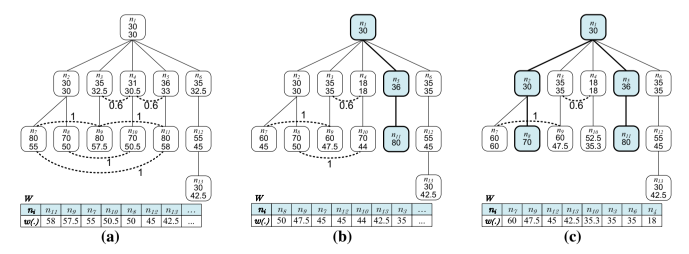


Fig. 5 2-LASPe algorithm (annotated with similarity edges of unselected nodes). a The initial OS (and simOS). b First update. c Final update



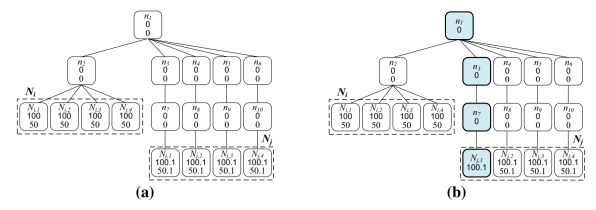


Fig. 6 2-LASPe: example of rival N_i and N_i sets (with $d_i = 2$). a The initial OS. b First update

with n_4 , n_7 and n_9 nodes and thus update them according to their similarities. Since at each iteration, we only need to check the similarity between a newly selected node against all unselected nodes, we delete all similarity edges of a newly selected node (line 4). Thus, after the first update and removal of p_{11} path, similarity edges of n_5 and n_{11} nodes are deleted from the sim OS tree.

6.1 Analysis

The worst-case complexity of this algorithm remains the same as that of LASP. However, in practice 2-LASPe is much faster; for instance, for an exemplary OS of size 735 and l=50, LASP conducts 24,267 updates (in 145 ms), whereas 2-LASPe performs only 214 updates (in 14 ms), resulting to snippets of similar quality. We can easily show that all optimality theorems that hold for LASP also hold for 2-LASPe. Empirically, 2-LASPe provides snippets of almost the same quality as LASP. Below, we prove the approximation lower bound of this algorithm.

We also investigated the 3-LASPe algorithm (i.e., averaging the score of a node with that of its parent and grandparent). We found that 3-LASPe is slower than 2-LASPe as it requires more operations, while again giving results of similar effectiveness and quality as 2-LASPe and LASP. Note that LASP is in fact a 4-LASPe algorithm for the DBLP and Google+ G^{DS} s that we use in our experiments, as both graphs have a maximum path length 4.

6.1.1 Lower bound of 2-LASPe approximation

Theorem 5 The 2-LASPe is a $d \cdot \frac{l-2}{l-1-d}$ -approximation algorithm for l > d+1, where l and d are the required size of snippets and the maximum depth of G^{DS} , respectively. Namely, the ratio of the optimal Im(.) (denoted as OPT) over the lower bound of an Im(.) generated by the 2-LASPe (denoted also as 2-LASPe) is only $d \cdot \frac{l-2}{l-1-d}$ times larger.

Proof We follow the same strategy as in LASP lower bound calculation; namely using rival sets N_i (for the optimal result) and N_i (for the approximate result) (Fig. 6).

Let N_i have the following common properties as in the LASP proof, i.e., it consists of more than l sibling and diverse nodes, and all nodes have a common score, $w(N_i)$. In contrast to the LASP algorithm, where the approximation error is maximized when N_i is at the lowest depth d of the OS tree, for 2-LASPe, the error is maximized when N_i is at depth 2. Since, when d_i is minimized, the number of N_i nodes to be included is maximized, and thus, the optimal snippet score is also maximized. $d_i = 2$ is the minimum depth that facilitates $ap(N_i)[z] = w(N_i)$ for all z (even after the addition of the first path to an N_i or N_j node). Note that for a smaller d_i , e.g., for $d_i = 1$, we have $ap(N_i)[z] = ap(N_i)[0]/2 = w(N_i)/2$ for $z \ge 1$ which reduces the holistic score. Thus, we assume that N_i is at depth 2. As before, we use $PPar_i$ (resp., $PPar_i$) to refer to the sums of w(.) scores of all nodes from the OS root until the parents of N_i (resp. N_i) set. The OPT score is maximized when $PPar_i = 0$, since as discussed in LASP proof, for a fixed $ap(N_i)[0]$, the $PPar_i = 0$ maximizes the score of the optimal snippet (Lemma 1). Thus, we get the maximum score for the optimal snippet when $PPar_i =$ $0, d_i = 2$ and $l \ge 2$. For instance, for l = 5 and $d_i = 2$, we have $Im(DSl(N_i)) = 0 + 0 + 100 + 100 + 100 = 300$.

Unlike in LASP algorithm, N_j will result in the worst case when the N_j nodes are leaves at maximum depth d and belong to separate paths with $PPar_j = 0$, instead of being siblings (see the example of Fig. 6). We assume that the w(.) scores of N_j nodes are all equal and marginally larger than the w(.) scores of the N_i nodes (so that they are selected over N_i nodes), i.e., we assume $w(N_i) \approx w(N_j)$. Analogously, when d_j is maximized, the number of N_j nodes to be included is minimized, and thus, the approximate score is also minimized. Similarly, we have $ap(N_j)[z] > ap(N_i)[z]$ for any z (even after the addition of the first path to an N_j node since it does not now affect ap(.) scores). And also,



2-LASPe score is minimized when $PPar_j = 0$, since apart from N_j nodes no other node contributes to the score.

We can easily see that the two sets are rivals and 2-LASPe will wrongly select N_j nodes. As we can include only $\lfloor \frac{l-1}{d} \rfloor$ nodes from the N_j set, we have:

$$\frac{OPT}{2 - LASPe} = \frac{0 + (l - 2) \cdot w(N_i)}{0 + \lfloor \frac{l - 1}{d} \rfloor \cdot w(N_j)}$$

$$\approx \frac{l - 2}{\lfloor \frac{l - 1}{d} \rfloor} \le \frac{l - 2}{\frac{l - 1}{d} - 1} \approx d \cdot \frac{l - 2}{l - 1 - d}.$$
(8)

Assuming that l > d+1, it makes the denominator in the above formula positive. Recall that in most cases $d \le 3$ and l > 10; therefore, the assumption is valid in typical settings.

We can see that the discussion in LASP about the effect of alternative rival sets and non-diverse graph nodes hold here: Rival sets N_i and N_j must consist of (1) at least l-2 and l-d nodes, respectively, with common score w(.) and (2) diverse nodes in order to maximize the approximation loss of 2-LASPe. Also, the N_i set should consist of sibling nodes.

7 Prelim-l algorithms

The aforementioned algorithms operate on the complete OS. Inspired by the prelim-l approach [15], we propose to produce a subset of the OS, denoted as DPrelim-*l* (or PPrelim-*l*) OS, that prunes nodes from the OS which have low probability to be considered for the size-l OS; this saves a lot from the OS generation time and the consequent size-l OS computation time. Note that the prelim-l generation approach of [15] considers the inclusion of the top-l nodes, i.e., the lnodes with the highest li(.) in the OS (allowing their repetitions); for clarity, we refer to this algorithm as VPrelim-l. The direct application of VPrelim-l is inappropriate here, especially for the PPrelim-l OS, for the following three reasons: (1) it allows the consideration in the top-*l* set of nodes repeatedly which is against the diversification requirements of PSize and DSize; (2) it fails to manage the non-monotonic relationship between $w(n_i|.)$ and $li(n_i)$ of proportionality (e.g., $w(n_i|\varnothing) > li(n_i)$), which requires the challenging estimation of similarity among nodes (and frequency per node) in the OS; and (3) it does not facilitate further pruning of nodes that have similarity with already added nodes (or are included multiple times).

Recall, however, that the two properties, proportionality and diversity, are based on different equations and thus have different properties in measuring the w(.) score. More precisely, the proportionality equation is more challenging as it requires the apriori knowledge of similarity between nodes (and frequency of nodes) in an OS in order to produce w(.)

scores. Thus, we first present more comprehensively a generalized version of the VPrelim-*l* approach, which addresses proportionality. This algorithm, denoted as PPrelim-*l*, considers the similarity/frequency and respective upper bounds of nodes in an OS, in order to produce the so called PPrelim-*l* OS. Then, we present more synoptically the DPrelim-*l* algorithm with the required specializations and simplifications in order to produce DPrelim-*l* OS.

7.1 PPrelim-l

The computation of the optimal PSize-l OS is very expensive and, as a consequence, so is the computation of any PPrelim-l OS that is guaranteed to include the optimal PSizel OS. Thus, we resort to a heuristic that aims to generate a PPrelim-l OS that includes at least the l diverse graph nodes (i.e., nodes that their similarity is less than a threshold $d(\theta)$) with the largest $w(n_i|\varnothing)$ scores (denoted as the $top_{w}l$ set). The rationale is that while searching for $top_{w}l$ and by appending the retrieved nodes to the PPrelim-l OS, we will generate a good superset of the PSize-l OS. The constraint of including only diverse nodes in top_wl is necessary in order to facilitate eventual diversity. We generate the PPrelim-l OS by extending the complete OS generation algorithm (Algorithm 4, described in Sect. 2.1 and in [13]) to include three pruning conditions. We traverse the G^{DS} graph in a breadth-first order, according to Algorithm 3. For this purpose, cheap pre-computed indexes, variables and data structures are employed. Hereby, we describe the algorithm by introducing the (1) pre-computed indexes per relation and $n^{\rm DS}$, (2) variables and data structures, and (3) the pruning conditions that are used as to construct the algorithm. We try to describe these terms, where possible, in the order they appear in the algorithm. Figures 7 and 8 illustrate an example and Table 4 summarizes the notation we are using.

The calculation of sim relevance requires more time (i.e., quadratic; as we have to compare each node against all remaining nodes); in comparison with equi relevance which simply requires counting the frequency of graph nodes. Thus, we address the two relevance types separately. For instance, for sim relations, we rely mainly our pruning on pre-computed indexes (e.g., $mw(n^{DS}, R_i)$ to be described shortly), whereas for equi relations, we achieve further pruning by using online retrieved information (e.g., $cmFr(R_i)$ to be described shortly).

7.1.1 Index per GDS relation

Our PPrelim-l OS generation technique uses three indexes (i.e., bounds) for each G^{DS} relation R_i , which are precomputed. $max(R_i)$ is the maximum value of li(.) in R_i . $mmax(R_i)$ is the maximum value of $max(R_i)$ of all R_i 's descendant nodes in G^{DS} or 0 if R_i has no descendants



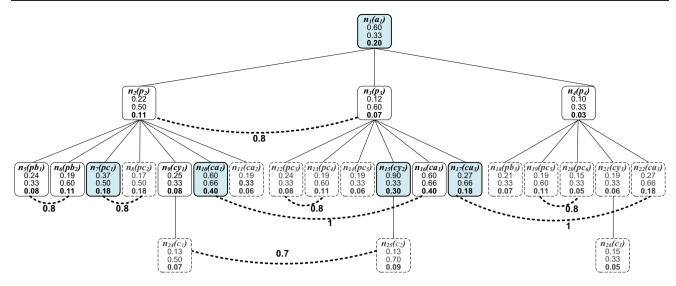


Fig. 7 PPrelim-l example for l = 5 (the complete OS, the PPrelim-l OS and the $top_w l$ set). Edges between nodes indicate their similarity. Nodes are annotated with their li(.), pq(.) and w(.). Nodes with

low transparency are pruned nodes (e.g., n_8 , n_{11}), shaded nodes are the $top_w l$ set (e.g., n_1 , n_7) and the rest are the remaining tuples of the PPrelim-l OS (e.g., n_2 , n_3)

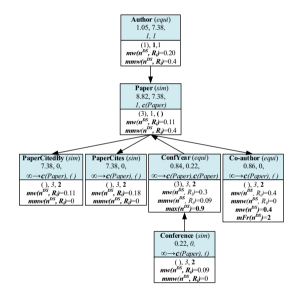


Fig. 8 DBLP author G^{DS} , annotated with relation indexes: (relevance type), $max(R_i)$, $mmax(R_i)$, UBFr (R_i) , $c(R_i)$, and n^{DS} indexes for the example of Fig. 7: $(c(R_i))$, UBFr (R_i) , $mFr(n^{DS})$, $mw(n^{DS}, R_i)$, etc.)

(i.e., R_i is a G^{DS} leaf node). Finally, UBFr(R_i) is the upper bound of joins a node in R_i can have with any n^{DS} . During preprocessing, we can determine only for some cases these bounds, e.g., when up to 1 node from a relation (e.g., R_{Paper}) can only join with n^{DS} . Otherwise, we assume infinite joins (e.g., the same co-author may appear in an unbounded number of papers) and set UBFr(co-author)= ∞ . In order to facilitate the calculation of UBFr(R_i), we also introduce the $c(R_i)$ variable which is the summation of tuples from R_i that can join with n^{DS} . Thus, for N:1 relationships, $c(R_i) = c(RPar_i)$, where $c(R^{DS}) = 1$ and $RPar_i$ is the parent relation of R_i . Thus, given $c(RPar_i)$ for cases where

Table 4 Notations of the PPrelim-l Algorithm

Notation	Definition				
$top_w l$ set	The <i>l</i> diverse graph nodes with the largest <i>w</i> scores				
$max(R_i)$	The $\underline{\text{max}}$ imum value of $li(.)$ in R_i				
$mmax(R_i)$	The <u>maximum $max(R_i)$ in all R_i's descendant nodes or 0 if R_i has no descendants</u>				
$\mathrm{UBFr}(R_i)$	The upper bound of joins a node in R_i can have with any n^{DS}				
$mw(n^{DS}, R_i)$	The <u>maximum</u> w score of R_i nodes in the n^{DS} OS				
$mmw(n^{DS},R_i)$	The <u>maximum</u> $\underline{mw}(n^{DS}, R_i)$ of all R_i 's descendents or 0 if R_i has no descendants				
$mw(n^{DS})$	The <u>maximum</u> \underline{w} score of nodes in the n^{DS} OS				
$mFr(n^{DS})$	The maximum frequency of any node in the OS				
$R(n_i)$	The relation n_i belongs to				
$R_i(n_j)$	The subset of R_i that joins with n_j				
$c(R_i)$	The summation of nodes in R_i that can join n^{DS}				
$\mathrm{UBw}(n_j,R_i)$	The <u>upper bound of \underline{w} in $R_i(n_j)$</u>				
$dUBw(n_j, R_i)$	The <u>upper bound</u> of \underline{w} from all R_i 's <u>descendants</u> that join with n_j				

UBFr(co-author)= ∞ , we estimate UBFr(R_i) as a function of $c(RPar_i)$, i.e., UBFr(R_i) = $c(RPar_i)$ (denoted as $\infty \rightarrow c(R_i)$ in Fig. 8); this association will be useful later, during the online calculation of tighter frequency bounds. Also note that since we only need $c(RPar_i)$, we do not need to calculate $c(R_i)$ for leaf nodes (thus, we denote their c(.) as () in Fig. 8). Finally, we define and use index dUBFr(R_i), which is the upper bound of joins of a node belonging to any descendant relation of R_i that can have with any n^{DS} .



Algorithm 3 The PPrelim-*l* OS Generation Algorithm

```
PPrelim-l (l, G^{DS})
 1: t = 0; n_i = n^{DS}; W_l = \{\}; Q = \{\}
 2: addNode(n<sub>i</sub>)
 3: while !IsEmptyQueue(Q) do
              Xn_i = n_i; n_i = deQueue(Q)
              for each child relation R_i of R(n_i) in G^{DS} do
 5:
                    if (UBFr(R_i) > 1) then
 6:
 7:
                           if (R(n_i) \neq R(Xn_i)) then
                                                                                                                               \triangleright n_i \in \text{new relation}
                                cmFr(R_i) = 0; i = c(RPar_i)
 8:
                           UBFr(n_j, R_i) = min\{-i + cmFr(R_i), mFR(n^{DS})\}\
 9.
 10:
                      else
                            UBFr(n_j, R_i) = 1
 11:
                    UBw(n_j, R_i) = \min\{mw(n^{DS}, R_i), f_1(\min\{max(R_i), max(n^{DS})\}, \min\{(mFr(n^{DS}), UBFr(R_i), UBFr(n_j, R_i)\})\}
 12:
                      dUBw(n_j, R_i) = \min\{mmw(n^{DS}, R_1), f_2(\min\{mmax(R_i), f
 13:
                    max(n^{DS})}, min{(mFr(n^{DS}), dUBFr(R_i)})}
 14:
                      if !(t \ge UBw(n_i, R_i) \&\& (t \ge dUBw(n_i, R_i))) then \triangleright Prun.
 15:
                            if (t \ge dUBw(n_i, R_i)) then
                                                                                                                                          ⊳ Prun. Cond.2
                                   R_i(n_i): get up to diverse l nodes with UBw(.) > t
 16:
                                where UBw(.) = f_3(.)
 17:
 18:
                                  get R_i(n_i)
 19:
                            for each tuple n_i of R_i(n_i) do
20:
                                  if R_i is equi relation then
21:
                                        if (UBFr(R_i) > 1) then
 22:
                                              UpdateHFr(n_i)
23:
                                              UpdatecmFr(n_i)
24:
                                              w(n_i)[1] = f(li(n_i), HFr[g(n_i)].fr, 1)
25:
                                        if !((HFr[g(n_i)].fr > 1) && (w(n_i)[2] < t) && (t \ge 1)
                                      dUBw(n_i, R_i)) then
                                                                                                                                          ⊳ Prun. Cond.3
26:
                                              addNode(n_i)
                                  else if R_i is sim relation then
27:
28:
                                        addOnHSi(n_i)
 29:
                                        for each n_k in HSi do
 30:
                                              if sim(n_i, n_k) then
 31:
                                                    Update (w(n_i|\varnothing), sim(n_i, n_k))
32:
                                                    Update (w(n_k|\varnothing), sim(n_k, n_i))
33:
                                                   cSim(n_i) = true
 34:
                                        w(n_i)[2] = f(w(n_i|\varnothing), 2)
                                                                                                                                                                 ⊳ Eq.4
35:
                                        if !(cSim(n_i) \&\& (w(n_i)[2] < t) \&\& (t \ge 1)
                                                                                                                                          ⊳ Prun. Cond.3
                                      dUBw(n_i, R_i)) then
36:
                                              addNode(n_i)
37: return PPrelim-l
addNode(n_i)
 1: EnQueue (O, n_i)
 2: add n_i on PPrelim-l as child of n_i or as root if n_i is n^{DS}
 3: if (w(n_i)[1] > t) then
              if n_i is not similar with any nodes in W_l then
                    EnQueue (W_l, n_i)
 6:
                    if (|W_l| > l) then
 7:
                           DeQueue(W_l)
                           if (|W_l| < l) then
 8:
 9:
                               t = 0
 10:
                            else
 11:
                                  t = \min(W_l)
```

7.1.2 Index per n^{DS} node

During preprocessing, we also maintain a number of indexes per n^{DS} . $mw(n^{DS}, R_i)$ is the maximum $w(n_i|\varnothing)$ of any

node in R_i (note that this can be prohibitively expensive to calculate online and can render the pruning ineffective). For instance, in our running example for paper, we have $mw(n^{DS}, R_i) = 0.11. mmw(n^{DS}, R_i)$ is the maximum value of $mw(n^{DS}, R_i)$ of all R_i 's descendants or 0 if R_i has no descendants (leaf node) (it can cheaply be obtained from descendants' $mw(n^{DS}, R_i)$. For example, for the paper relation, $mmw(n^{DS}, R_i) = 0.4$ due to $mw(n^{DS}, R_i) = 0.4$ of the co-author relation. $max(n^{DS})$ is the maximum li(.)for all nodes in an OS (excluding n^{DS}); e.g., in Fig. 8, $max(n^{DS}) = 0.9$ is found in the ConfYear relation. Note that this score overrides the maximum li(.) score of all G^{DS} relations (i.e., $max(R_i)$ and $mmax(R_i)$). $mw(n^{DS})$ is the maximum $w(n_i|\varnothing)$ of any node in the OS (excluding $n^{\rm DS}$). Similarly to $max(n^{\rm DS})$, this score is considered as the upper bound of w(.) of all nodes of all relations (e.g., in our running example, $mw(n^{DS}) = 0.4$ is found in relation coauthor). $mFr(n^{DS})$ is the maximum frequency of any node in the OS that belongs to a relation with $UBFr(R_i) > 1$, where $mFR(n^{DS}) < UBFr(R_i)$. For instance, in our example, $mFr(n^{DS}) = 2$ (since ca_1 and ca_3 appear twice); which is less than $UBFr(R_i) = 3$; thus, we can use this as a tighter bound and thus override the $UBFr(R_i)$ bound.

7.1.3 Variables and data structures

Let $R_i(n_i)$ be the subset of R_i that joins with n_i , and $R(n_i)$ be the relation where to n_i belongs. While processing n_i (in $R(n_i)$) against a relation R_i with UBFr(R_i) > 1, we try to get a tighter bound than $UBFr(R_i)$ and $mFr(n^{DS})$, denoted as $UBFr(n_i, R_i)$. For this purpose, we maintain the current maximum frequency, denoted as $cmFr(R_i)$; a node was found so far from R_i (i.e., from processing predecessor nodes, $n_1, ..., n_{j-1}$, of n_j against R_i , i.e., from their respective $R_i(n_1), ..., R_i(n_{i-1})$ sets). For instance, consider we are processing node n_4 (p_4) against the coauthor relation, node ca_1 is the most frequent among all $R_i(n_1), ..., R_i(n_{i-1})$ sets that were found so far since it was found twice; thus, $cmFr(R_i) = 2$. Given $cmFr(R_i)$, UBFr(n_i , R_i) assumes that ca_1 will appear in all the remaining sets $R_i(n_i), ..., R_i(n_{|R_i|})$ after processing n_i . At the beginning, $UBFr(n_i, R_i)$ can be very loose, so we compare it with $mFr(n^{DS})$, to keep the minimum of the two (lines 6-11).

Another bound we use is $UBw(n_j, R_i)$, which is the upper bound of the $w(n_i|\varnothing)$ score that can be obtained from $R_i(n_j)$ (line 12) (this value will be useful as to facilitate Pruning Condition 1). It is defined as the minimum of $f_1(.)$ and $mw(n^{DS}, R_i)$. The $mw(n^{DS}, R_i)$ index can be a very effective pruning tool for both relevance types. $f_1(.)$ aims to facilitate further pruning for equi relations thus is defined as follows: for sim relations are set to ∞ (as we do not expect to achieve a better bound than $mw(n^{DS}, R_i)$) that can be practi-



cally useful), whereas for equi relations are calculated using Equation 4 for z = 1.

We denote as $dUB(n_i, R_i)$ the upper bound of $w(n_i|\varnothing)$ of all nodes from R_i 's descendant relations that can join with n_i or 0 if R_i has no descendants (and it will be useful in facilitating Pruning Condition 2). Similarly to $UBw(n_i, R_i)$ calculation, $dUBw(n_i, R_i)$ can be defined as the minimum of $mmw(n^{DS}, R_i)$ and $f_2(.)$. $f_2(.)$ is also defined by Equation 4 for z = 1 (line 13). The $mmw(n^{DS}, R_i)$ index can be a very effective pruning tool for both relevance types. Analogously, $f_2(.)$ aims to facilitate further pruning for equi relations thus is defined as follows: for sim relations are set to ∞ , whereas for *equi* relations are calculated using the given parameters. Also note that, if R_i is a leaf node on G^{DS} then $mmax(R_i) = 0$, and thus, $dUBw(n_i, R_i) = 0$. $UBw(n_i, R_i)$ and $dUBw(n_i, R_i)$ bounds are specializations of $max(R_i)$ and $mmax(R_i)$ that have been used in prelim-l [15] in pruning conditions 1 and 2, respectively; however, they are tighter bounds as they are specific for the given n^{DS} .

Finally, we define the upper bound of $w(n_i|\varnothing)$ score of a node as $UBw(n_i)$ (which will be useful during Pruning Condition 2; line 16). We calculate $UBw(n_i)$ using the respective $pq(n_i|\varnothing)$ produced by function f_3 which is defined as follows. For an R_i being an equi relation, $pq(n_i)[1] = f_3(li(n_i), UBFr(n_j, R_i))$. Whereas, for an R_i being a sim relation, $pq(n_i|\varnothing) = f_3(((|OS(R_i)| - UBFr(n_j, R_i)) \cdot msim(n_i) + UBFr(n_j, R_i) \cdot 1) \cdot li(n_i))$, where $|OS(R_i)|$ is for $UBFr(R_i) = 1$ the amount of nodes of R_i in the OS and for $UBFr(R_i) > 1$ $c(RPar_i)$. $msim(n_i)$ is the maximum similarity of an n_i node with any other node in the OS. Namely, we assume that n_i appears $UBFr(n_j, R_i)$ times (thus $UBFr(n_j, R_i) \cdot 1$ similarity) and has the maximum similarity with all $OS(R_i)$ nodes.

As we have already explained, we process sim and equi relations separately. Thus, while processing equi relations, we maintain HFr, a hash table, which indexes for each graph node the computed frequency in the OS so far (lines 22 and 23). And while processing sim relations, we maintain HSi hash table which indexes the similarity of each OS node against all other OS nodes (i.e., $\sum_{n_i \in OS} sim(n_i, n_j)$). The update of HSi requires quadratic time as we need to compare the similarity of each node against all previous nodes already on HSi (lines 29-32). We also use the $cSim(n_i)$ flag variable to indicate whether n_i is similar to other nodes (line 33). We also denote as $w(n_i)[2]$, the $w(n_i)$ score given another node with maximum similarity with n_i has previously been added (lines 16 and 34); maximum similarity is the equality similarity (thus the common use of the equi notation).

We manage the retrieval of $top_w l$ set as follows. Let t be the current smallest value of the $top_w l$ set (or 0 if $top_w l$ does not contain l values yet). If the current tuple n_i is greater than t (line 3, function AddNode) and if n_i is diverse to all

 $top_w l$ nodes, then is added to the PPrelim-l and the l-sized priority queue W_l which manages the $top_w l$ set (AddNode(), lines 4–11). For instance, in Fig. 7, the shaded nodes comprise the final $top_w l$ set for the given OS. Note also that, although node n_{16} has score larger than t=0.18, it is excluded as it is similar (equal) to node n_{10} . Note that by considering the computed similarity/frequency of a node n_i so far, which is less than or equal to actual similarity/frequency of n_i , in fact, we consider the lower bound of w(.).

7.1.4 Pruning conditions

Each time we further process a node n_j , we employ three pruning conditions:

- Pruning Condition 1. If t is greater than or equal to the $w(n_i|\varnothing)$ of all tuples of the current relation R_i and all its descendants (i.e., $t > \text{UBw}(n_j, R_i)$) and $t > \text{dUBw}(n_j, R_i)$), then there is no need to traverse the sub-tree starting at R_i (line 14).
- Pruning Condition 2. We can limit the amount of tuples returned by an $R_i(n_j)$ join (i.e., by avoiding computing the entire join of n_j with R_i), if we can infer that none of R_i 's descendants (if any) can be fruitful for the $top_w l$ (i.e., when $t > dUB(n_j, R_i)$; line 15). Then, we can extract only nodes that their upper bound $w(n_i|\varnothing)$ score, $UBw(n_i)$, is greater than t (line 16).
- Pruning Condition 3. When Pruning Condition 2 holds, we can safely extract only part of the join. However, it is still possible that we extracted nodes which are similar to already added on the PPrelim-l nodes, and thus, their $w(n_i|n_{i1}, \dots n_{ix})$ will be actually used. Thus, we introduce a new pruning condition that checks first if a node n_i is similar to an added node and then considers adding it. For equi relations, we can easily detect equality by accessing HFr. Whereas for sim relations, this is more demanding (requiring quadratic time; lines 29-33), thus we use the $cSim(n_i)$ flag variable to detect similarity. For both cases, we use a safe bound i.e., $w(n_i)[2]$, and we do not add the node unless it is greater than t (lines 25 and 35). Recall that these scores are actually lower bounds as they are produced by comparison against only already retrieved nodes.

7.2 DPrelim-l OS

We simplify the previous algorithm by excluding all work concerning calculating or upper bounding the frequencies of nodes. For instance, indexes such as $UBFr(R_i)$, $mFr(n^{DS})$ and calculations of $UBFr(n_j, R_i)$ are not required. We adjust accordingly our algorithm to include these alterations (e.g., exclude calculations of $UBFr(n_j, R_i)$; functions (lines 12 and 13) use Equations 2 and 4, etc.).



7.3 Analysis

In terms of cost, in the worst case, we need up to *n* extractions of nodes, where n is the amount of nodes in the complete OS. In practice, however, there can be significant savings if the top_wl tuples are found early and large sub-trees of the complete OS are pruned. The PPrelim-l (resp. DPrelim-l) OS does not essentially contain the optimal PSize-l (resp. DSize-l) OS; in practice, however, we found that this is the case in most problem instances. This means that the PSize-l (resp. DSize-1) OS computation algorithms most likely give the same results when applied either on the PPrelim-l (resp. DPrelim-l) OS or the complete OS. Similarly to LASP and 2-LASPe algorithms, DPrelim-l and PPrelim-l algorithms will produce optimal results (i.e., supersets of the respective optimal size-l OSs) when we have a monotonic OS(li) and monotonic OS(w), respectively. Note that PPrelim-l cannot return an optimal solution when we have simply a monotonic OS(li(.)) because the proportionality equation (in contrast to the diversification equation) is not monotonic to li. The following theorem proves the lower approximation bound and that if the li(.) scores of nodes are monotonic then the computed DPrelim-l OS will be optimal.

7.3.1 Lower bound of the PPrelim-l (resp. DPrelim-l) algorithms

Theorem 6 The PPrelim-l (resp. DPrelim-l) algorithm is a $d \cdot \frac{l-1}{l-1-d}$ -approximation algorithm for l > d+1, where l and d are the required size of snippets and the maximum depth of G^{DS} , respectively. Namely, the largest possible score ratio of the optimal size-l OS (denoted as OPT) that is computed from the complete OS over the optimal size-l OS that is computed from the PPrelim-l (or DPrelim-l) OS (denoted as prelim-l) is $d \cdot \frac{l-1}{l-1-d}$.

Proof We follow an analogous strategy as in LASP and 2-LASPe lower bound calculation, namely, using rival sets N_i (included in the optimal solution, but not in the PPrelim-l OS) and N_j (affecting the approximate solution). Recall that according to the PPrelim-l computation algorithms, the PPrelim-l OS contains the l diverse nodes with the maximum w(.), denoted as $top_w l$ and all nodes along the paths from the OS root to these nodes. The worst case happens when none of the nodes of the $top_w l$ set is part of the optimal size-l OS at and at the same time this results to the maximum loss. Thus, we define N_i and N_j as follows.

 N_i is defined as a set of l sibling nodes with common score at depth 1. This minimum depth $(d_i = 1)$ allows the inclusion of the maximum amount of N_i nodes. Based on our analysis for 2-LASPe, this results in the maximum total score for the snippet. N_j consists of l nodes with common scores, and these are the $top_w l$ nodes. Similarly to 2-LASPe, N_j will

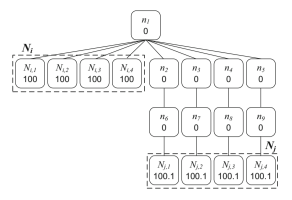


Fig. 9 PPrelim-l: example of rival N_i and N_j sets (with $d_i = 1$, $d_j = d$ and N_j ($top_w l$))

result to the worst case when the N_j nodes are leaves at maximum depth d and belong to separate paths with $PPar_j = 0$, instead of being siblings (see the example of Fig. 9). We assume that the w(.) scores of N_j nodes are all equal and marginally larger than the w(.) scores of the N_i nodes (so that they are selected over N_i nodes in the Prelim-l OS), i.e., we assume that $w(N_i) \approx w(N_j)$. As we can include only $\lfloor \frac{l-1}{d} \rfloor$ nodes from the $top_w l$ set in the Prelim-l OS, we can easily see that the score difference between OPT and prelim-l is maximized when (i) $PPar_i = 0$, such that the root does not count both in the optimal snippet and in the snippet computed from the Prelim-l OS, (ii) $PPar_j = 0$, such that besides N_j nodes, no other node contributes to the score of the snippet from the Prelim-l OS, and thus, prelim-l is minimized. Thus, we have:

$$\frac{OPT}{prelim-l} = \frac{0 + (l-1) \cdot w(N_i)}{0 + \lfloor \frac{l-1}{d} \rfloor \cdot w(N_j)}$$

$$\approx \frac{l-1}{\lfloor \frac{l-1}{d} \rfloor} \le \frac{l-1}{\frac{l-1}{d}-1} = d \cdot \frac{l-1}{l-1-d} \tag{9}$$

For instance, consider the example of Fig. 9, $w(N_i) = 100$, d = 3, and l = 5; we have OPT: 0 + 100 + 100 + 100 + 100 = 400 and prelim-l: two paths to N_{j1} and to n_3 , respectively: 0 + 0 + 0 + 100.1 + 0 = 100.1.

We assume that l>d+1, which makes the denominator in the above formula positive. If $l\leq d+1$, then the N_j nodes should be chosen at depth l in order to include a single node from N_j , as opposed to multiple N_i nodes. In this case, the ratio will be $\frac{l-1}{1}$, i.e., l-1. We can easily see that as discussed in LASP, rival sets must consist of (1) diverse nodes (thus, PPrelim-l approximation also holds for DPrelim-l) and (2) have a common score w(.).

Finally, we can bound the approximation ratios of LASP and 2-LASPe which apply on a PPrelim-l OS (or DPrelim-l) OS, over the optimal solution in the complete OS, by multiplying the approximation ratios of LASP and 2-



LASP, respectively, stated in theorems 1 and 5 by the ratio OPT/prelim-l stated in Theorem 6. For example, a lower upper bound for the approximation loss of LASP which applies on a PPrelim-l OS compared to the optimal solution on the complete OS is $d \cdot (d \cdot \frac{(l-1)}{l-1-d}) = d^2 \cdot \frac{(l-1)}{l-1-d}$.

7.3.2 Optimality of PPrelim-l (resp. DPrelim-l) algorithms

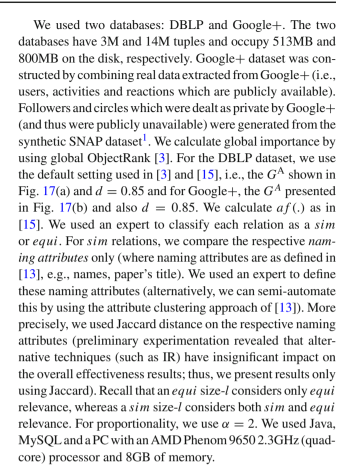
Theorem 7 For equi relevance, if the local importance scores of nodes (li(.)) are monotonically non-increasing with respect to the distance of the nodes to n^{DS}, then DPrelim will produce the optimal DPrelim-l OS (i.e., a superset of the optimal equi DSize-l OS).

Proof The DPrelim OS will include the (1) $top_w l$ set, i.e., ldistinct nodes with the largest w(.)[1] scores where t is the minimum $(top_w l)$ and (2) also all repetitions of $top_w l$ nodes with w(.)[2] > t, denoted as $rep_w l$ (since we only prune nodes with w(.)[1] < t and w(.)[2] < t in pruning conditions 2 and 3, respectively). When we have a monotonic OS, we can produce the optimal DSize-l OS by using the LASP algorithm (Theorem 3) as follows. Initially, we will include jdistinct nodes with the largest li(.) scores (as they also correspond to the largest ap(.) since w(.)[1] = li(.), where j < l), where a subset of these j nodes may have fr > 1. All these nodes are members of the $top_w l$ by definition. Then, consider that if the next node to be added (according to LASP) is a repetition of a node (i.e., we include it considering its w(.)[2]); then, this node is member of $rep_w l$, as by contradiction it should have w(.)[2] > t (as otherwise another distinct node would have been selected). Thus, we conclude that the optimal DSize-l will comprise nodes from either $top_w l$ or $rep_w l$ nodes which are included in DPrelim-l by definition. (Note that for the same reason as in Theorem 2, this algorithm cannot provide an optimal solution for *sim* relevance.)

Theorem 8 *Similarly, based on theorems 4 and 7 we can easily see that the PPrelim can produce an optimal PPrelim-* l *OS for equi relevance if we have a monotonic OS(li(.)) and all nodes have* $fr(n_i) < \alpha + 1$.

8 Experimental evaluation

We experimentally evaluate the proposed snippets and algorithms. We emphasize on effectiveness comparisons between the two types of diversified snippets, the two types of relevance and also against the non-diversified size-l snippets [15]. Firstly, we thoroughly investigate the effectiveness and usability of the produced snippets with the help of human evaluators. Then, we evaluate the quality of the size-l OSs produced by the greedy heuristics. Finally, we comparatively investigate the efficiency of the proposed algorithms.



8.1 Effectiveness

We conducted an effectiveness evaluation with the help of human evaluators. The evaluators were professors and researchers from our universities. None of our evaluators were involved in this paper. Because of the complexity of the evaluation (we have to compare five different types of snippets), we used evaluators with expertise in the topics we investigate. In particular, since the DBLP database includes data about real people, we asked the DSs themselves where possible (i.e., eleven authors or students of authors listed in DBLP) to participate in this evaluation. The rationale of this evaluation is that the DSs themselves (even their students) have the best knowledge of their work and can therefore provide accurate summaries. For Google+, we presented ten random OSs to nine evaluators. First, we familiarized them with the concepts of OSs in general and the five types of size-l OSs. Specifically, we explained that a good size-l OS should be a standalone and meaningful synopsis of the most important information about the particular DS. In addition, we explained that DSize-l OSs and PSize-l OSs consider diversity and proportionality, respectively, and the difference between the two relevance types. However, we avoided to dis-



¹ http://snap.stanford.edu/data/egonets-Gplus.html.

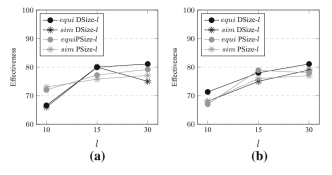


Fig. 10 Effectiveness (i.e., recall = precision). **a** DBLP author. **b** Google+ user

cuss the advantages or disadvantages of these combinations of types as to avoid any bias. In order to assist them with their tasks, we provided them useful information per node, such as $fr(.), li(.), dv(n_i|\varnothing), pq(n_i|\varnothing)$ and $w(n_i|\varnothing)$. For instance, we provided them, with the amount of times the co-author C. Faloutsos appears in the M. Faloutsos OS, his li(.), etc. We also provided summarized ranked tables (similar to Tables 2 and 3 at the end of each OS) with the top-10 most frequent and top-10 most important nodes and their respective $w(n_i|.)$ scores.

8.1.1 Precision and recall

We provided evaluators with OSs and asked them to DSize-l and PSize-l them using both types of relevance (i.e., equality and similarity) for l = 10, 15, 30. Figure 10 measures the effectiveness of our approach as the average percentage of the nodes that exist in both the evaluators' size-l OS and the computed size-l OS by our methods. This measure corresponds to recall and precision at the same time, as both the OSs compared have a common size. Figure 10a, b plot the recall of the DSize-l and PSize-l for DBLP author and Google+ user G^{DS} 's Fig. 18. On average, the effectiveness of DSize-l and PSize-l OSs ranges from 67 to 82% for all cases, which is very encouraging. The results of Fig. 10 are obtained using the LASP algorithm (as the BF-l algorithm was prohibitively expensive). We omit results obtained by our other approximate algorithms as they do not vary from these results. For instance, the 2-LASPe algorithm gave almost identical results as LASP and the use of DPrelim-l OSs or PPrelim-l OSs had no impact on effectiveness. As we show later, they have very minor impact on the quality of the computed snippets.

8.1.2 Usability test

We conducted a comparative study of the usability of the five types that verifies users' preference for *sim* over *equi* relevance and DSize-*l* and PSize-*l* OSs over size-*l* OSs. In

summary, the evaluation reveals the usability superiority of *sim* PSize-*l* OSs over all other types. Usability is the ease of use and learnability of a human-made object; namely, how efficient it is to use (for instance, whether it takes less time to accomplish a particular task), how easy it is to learn and whether it is more satisfying to use.² More precisely, for a given OS, we measured the ease of use of all types through a usability test. We presented to users the various versions of size-*l* OSs in a random order to avoid any bias, and we also gave them six tasks to complete for each OS. Then, we asked them to give a score in a scale of one to ten and also to justify in their answers, where possible, the usability of the five approaches when completing these tasks. Namely, to score them considering (1) the ease of accomplishing each task, (2) how easy and (3) satisfying are to learn and use.

More precisely, the first task (T1) was to score the general use of all types; namely, which one they prefer as a representative and informative snippet. For this purpose, we emphasized again that a snippet should be short, stand-alone and a meaningful synopsis of the most important and representative information about the particular DS; we avoided to discuss any advantages/disadvantages. The rest of the tasks were to extract information about the DSs. For the DBLP author, Task 2 (T2) was to determine the most frequent coauthors of a given author (e.g., whether C. Faloutsos and S. Krishnamurthy are among the most frequent collaborators of M. Faloutsos). Task 3 (T3) was to determine the most important co-authors (e.g., whether C. Faloutsos and S. Madden are among the most important co-authors of M. Faloutsos). Task 4 (T4) was to determine the most frequent journal/conference the DS has published. Task 5 (T5) was to determine the most frequent topic of an author's papers (i.e., repeated set of keywords appearing in author's papers). Finally, Task 6 (T6) was to determine the most frequent topic appearing in papers citing an author's papers. Analogous tasks were used for the Google+ user. Namely, T2 was to determine a couple of the most frequent users in the DS's circles; T3 was to determine a couple of the most important users in DS's circles; T4 was to determine the most frequent user making comments on DS's activities; T5 was to determine the most frequent topic of the DS's comments and T6 was to determine the most frequent topics of DS's activities. Note that for comparison purposes, we maintain an analogy between the respective tasks of the two databases, e.g., T2 of both databases aim to determine the most frequent co-authors/users associated with the DS, whereas T3 to determine the most important co-authors/users, etc.

Figures 11 and 12 average the evaluators' usability scores of all methods per $G^{\rm DS}$, per task and per l. More precisely, respective subfigures (a) represent scores for l=15, (b) for l=30, and (c) the average of all tasks per l. The results show



² www.wikipedia.org/wiki/Usability.

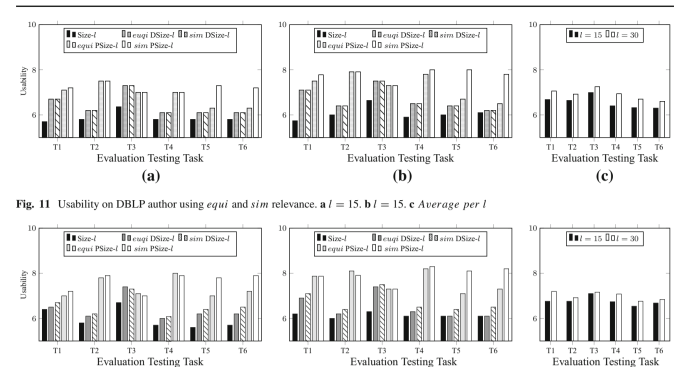


Fig. 12 Usability on Google+ user using equi and sim relevance. a l = 15. b l = 15. c Average per l

that evaluators preferred firstly sim PSize-l OSs, secondly equi PSize-l OSs, then sim and equi DSize-l OSs and lastly size-l OSs for both datasets. They also preferred size l = 30over l = 15. For instance for the author G^{DS} , the average scores of all tasks and both values of l, for sim PSize-l OSs is 7.5, for equi PSize-l OSs is 7.1, for sim DSize-l OSs is 6.7, for equi DSize-l OSs is 6.6 and finally for size-l OSs is 6.0. Evaluators expressed very similar preference for equi and sim DSize-l OSs because the snippets of these two types are almost identical (i.e., their constituent nodes are almost the same). The reason is that for the specific DBLP author G^{DS} , both relevance types equi and sim result in the same DSize-l OSs, as the sim relevance will only impact toward the avoidance of including papers or conferences with frequent textual similarity to already added nodes (which was not very often in these cases).

(a)

The evaluators also provided justifications for their scores. We summarize them for each type and l, and we also analyze their reflection on the given tasks. The evaluators explained that in general, they prefer the concept of PSize-l OS as it also considers frequent nodes and topics; this is a property other types do not consider. This is evidenced by the superiority of the usability of PSize-l for tasks T2, T4, T5 and T6, since these tasks consider the frequency of nodes and topics. In addition, the evaluators explained that they found useful the results considering the frequency of keywords (i.e., frequent topics); this is evidenced by high scores of sim PSize-l for tasks T5 and T6 which address the frequency of topics in the

results. However, as they pointed out, although the inclusion of repeated frequent items or topics is informative, it comes at the cost of excluding other important nodes. They found that a DSize-l OS is very useful in covering the most important elements of an OS (i.e., evidenced by high scores of DSize-l for Task 3); however, they pointed out that rare but important elements may appear which again can be misleading to some extent. They found the non-diversified size-l summaries [15] more misleading as very important nodes are too dominant in them. The evaluators stated that l values of around 30 are the most appropriate, since the corresponding snippets include sufficient descriptive information about the corresponding OSs, giving a better representation of frequent and important information, and without being overwhelmingly large. This is also evidenced by Figs. 11c and 12c.

(c)

8.2 Quality of snippets

(b)

We now compare the holistic importance Im(.) scores of DSize-l and PSize-l OSs produced by the greedy methods. More precisely, the results of Fig. 13 represent the average holistic scores for ten random OSs per G^{DS} . The average size (i.e., the amount of nodes) of OSs is also indicated (denoted as (|OS|)). The results show that in most cases, the results of LASP and 2-LASPe are of very similar (or even identical) quality, i.e., they have similar (or equal) holistic Im(.) scores. The evaluation also reveals that using the DPrelim-l



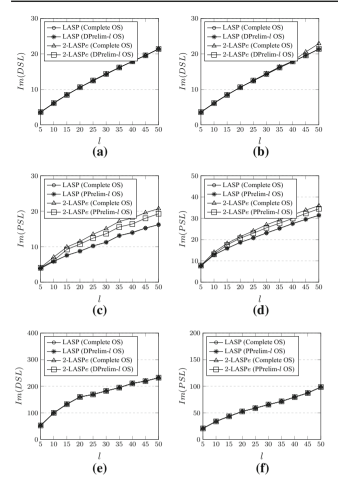


Fig. 13 Quality on DBLP and Google+. **a** equi DSize-l author $(|\overline{OS}| = 707)$. **b** sim DSize-l author $(|\overline{OS}| = 707)$. **c** equi PSize-l author $(|\overline{OS}| = 707)$. **d** sim PSize-l author $(|\overline{OS}| = 707)$. **f** equi DSize-l user $(|\overline{OS}| = 132K)$. **g** equi PSize-l user $(|\overline{OS}| = 132K)$

and PPrelim-l OSs results to very minor (even to zero) quality loss compared to using the complete respective OSs; e.g., by using LASP on either the complete OS or on the corresponding DPrelim-l OS, we obtain a DSize-l OS of the same Im(DSl). More precisely, for the case of the DBLP author equi PSize-l OSs, we get the maximum score loss by our algorithms; i.e., 2-LASPe complete and LASP PPrelim-l algorithms return scores 20 and 15.5, respectively, for l=50. In the Google+ user case, respective quality remains the same for all (combinations of) algorithms (thus, we omit sim DSize-l and sim PSize-l results). We did not compare with the optimal results, as the BF-l algorithm is too expensive.

8.3 Efficiency

We compare the run-time performance of our greedy algorithms in Figs. 14, 15 and 16. We used the same OSs as in Sect. 8.2 (i.e., the same ten OSs per G^{DS}). Figures 14 and 15 show the costs of our algorithms for computing DSize-l (resp.

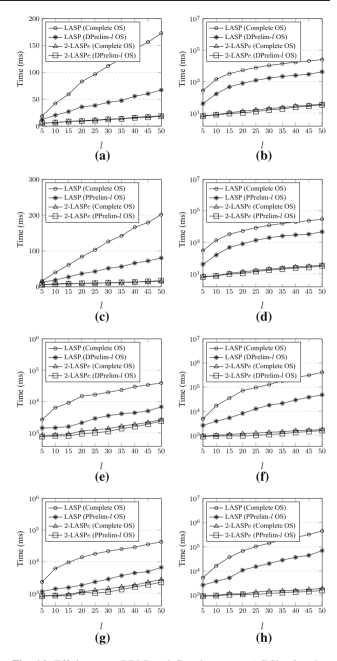


Fig. 14 Efficiency on DBLP and Google+. **a** equi DSize-l author (|OS| = 707). **b** sim DSize-l author (|OS| = 707). **c** equi PSize-l author (|OS| = 707). **d** sim PSize-l author (|OS| = 707). **e** equi DSize-l user (|OS| = 132K). **f** sim DSize-l user (|OS| = 132K). **g** equi PSize-l user (|OS| = 132K). **h** sim PSize-l user (|OS| = 132K).

PSize-l) for both types of relevance (equi and sim), excluding the time required to generate and preprocess OSs (i.e., the generation of w(.), ap(.) scores), where each algorithm operates on. More precisely, Fig. 14 show the costs of our algorithms for computing size-ls from OSs of the two $G^{\rm DS}$ s with various sizes and using a range of l values. The average sizes of the OSs on which the algorithms operate are indicated in brackets for each $G^{\rm DS}$. Figure 15a, b shows the scalability for author PSize-l of different sizes, after fixing l=10 (anal-



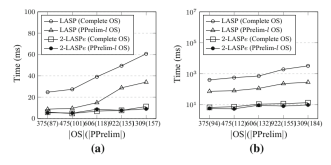


Fig. 15 Efficiency (varying OS size). a equi PSize-l author (size-l = 10). b sim PSize-l author (size-l = 10)

ogous results were obtained from User G^{DS} and DSize-l and thus we omit them). Each value on the x-axis represents an OS size (and the corresponding PPrelim-10 size). Comparing these numbers, we can get an indication of preliminary OSs savings; e.g., the OS with size 1,309 has a PPrelim-10 size 157 (i.e., 11% of the size of the complete OS). From Figs. 14 and 15, we can see that the use of 2-LASPe on a preliminary OS is the fastest approach. For instance, equi DPrelim-l 2-LASPe for l = 50 requires only 18.3ms. The results also verify that the use of sim relevance is more expensive than the use of equi relevance as it dictates the comparison of each node against all other OS nodes. For instance, sim DPrelim-l 2-LASPe for l = 50 requires up to 33.9ms (which remains a practical time).

Finally, Fig. 16a and b break down the cost to OS generation and preprocessing time (bottom of the bar) and size-l computation (top of the bar) for each method for PSize-1. The figures also show (on the x-axis) the average sizes of the complete OSs and the PPrelim-l OSs for l = 10 and l = 50, respectively. For instance, the average size of the complete OS is 707, whereas the average sizes of the corresponding equi PPrelim-10 and PPrelim-50 OSs are 119 and 272. Evidently, the preliminary OS generation is always faster than that of the complete OS; for instance, the PPrelim-5 OS's size is approximately 10% of the size of the complete OS and its generation can be done up to 2.5 times faster. Also, 2-LASPe is always faster at both phases (i.e., during OS generation and preprocessing and during size-l calculation) as at both phases, more operations are required by LASP (recall that during preprocessing, the ap(.) of a node in LASP corresponds is the path to the root, whereas in 2-LASPe is the node with its parent only). The comparison of Fig. 16a, b verifies again that sim relevance is more demanding than equi relevance. In general as expected, the OS size, l and sim relevance negatively affect the cost.

The cost of the BF-l algorithm becomes unbearable for moderate OSs sizes and values of l. For instance, although using BF-l we could get results for l=5 (e.g., 16 ms for the author $R^{\rm DS}$), we had to terminate the algorithm for $l\geq 10$ as it exceeded 30min of running. In summary, the BF-l algo-

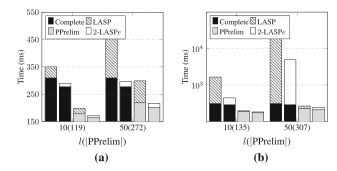


Fig. 16 Efficiency (cost breakdown). a equi PSize-l author ($\overline{|OS|}$ = 707). b sim PSize-l author ($\overline{|OS|}$ = 707)

rithm is not practical at all, whereas our greedy algorithms are very fast, and as we showed in Sect. 8.2, their results are snippets of high quality. In addition, the use of preliminary OSs and 2-LASPe is constantly a better choice over the complete OSs and LASP, respectively, since they are always faster with a negligible quality loss.

9 Related work

We present and compare related work in relational keyword search, ranking and summarization. To the best of our knowledge, no previous work has focused on the computation of diverse and proportional size-*l* OSs.

9.1 Keyword search and ranking

Relational keyword search facilitates the discovery of joining tuples (i.e., Minimal Total Join Networks of Tuples (MT-JNTs) [20]) that collectively contain all query keywords and are associated through their keys; hence, the concept of candidate networks is introduced (e.g., DISCOVER [19,20]). Relational keyword search paradigms differ from OSs semantically, since they search for connections of keywords. Précis Queries [23,25] resemble size-l OSs as they append additional information to the nodes containing the keywords, by considering neighboring relations. However, a précis query result is a logical subset of the original database (see [13] for a detailed comparison to size-l OSs). Other works in this context also investigate indexing and ranking techniques to facilitate efficiency [24]. Recent related work investigated keyword search on tree structured data, e.g., [9].

Related ranking paradigms consider *Importance*, which weights the authority flow through relationships (e.g., ObjectRank [3], ValueRank [14], PageRank [4]). In this work, we use nodes' importance to model $gi(n_i)$ and more precisely global ObjectRank. Our algorithms are orthogonal to how importance of nodes is defined (alternative methods could also be investigated).



9.1.1 Document summarization

Web snippets [21,27] are examples of document summaries that accompany search results of Web keyword search in order to facilitate their quick preview. They can be either static (e.g., the first words of the document or metadata) or query-biased (e.g., containing the query keywords). However, the direct application of such techniques on OSs and databases in general is ineffective as they disregard the relational associations (e.g., for q = "Faloutsos," papers authored by Faloutsos will be disregarded as they do not include the "Faloutsos" keyword).

9.2 Diversity

Diversification of query results has attracted a lot of attention recently as a method for improving the quality of results by balancing similarity (relevance) to a query q and dissimilarity among results. Typically, given a query q and a desired number of results k, firstly, we get a ranked list of results S in descending order of their similarity to q, denoted as $sim(s_i, q)$; namely, $S = \langle s_1, ..., s_n \rangle$ where $n \geq k$. Then, the objective of diversification is to find a subset of S, $R \subseteq S$ of size k, such that the elements in R are similar to q (w.r.t $sim(q, s_i)$) and at the same time dissimilar to each other (w.r.t. $dis(s_i, s_i)$). The definition of sim and dis scores is orthogonal to the diversification problem per se, and a variety of IR-based or probabilistic approaches have been used to define such functions (e.g., PageRank-based similarity). In most cases, the same function is used to estimate both scores (i.e., $dis(s_i, s_i) = 1 - sim(s_i, s_i)$). One of the earliest and most influential diversification functions is maximal marginal relevance (MMR) [5], which trades off between the novelty (a measure of diversity) and relevance of search results; a parameter is used to control this trade-off. A general framework for result diversification appears in [18] with eight axioms. In [18,28], max-sum, max-min and monoobjective objective functions and algorithms are proposed. Our proposed diversity definition is inspired by this monoobjective approach, where for each document, a single score trades off the relevance to the query and the dissimilarity from other documents. In [1,2], probabilistic interpretations of sim and dis functions and objective functions are proposed. In [10,11], an intuitive definition of diversity, called DisC diversity, is proposed where the computed diverse subset R covers all elements of S in the sense that for each element in S there should be a similar element in R, and at the same time, the elements in R should be dissimilar to each other (i.e., diverse). In [22], LogRank is proposed, a principled authority-flow-based algorithm that computes a representative summary of the user's activities by selecting activities that are simultaneously important, diverse and time-dispersed.

9.2.1 Proportionality

[7,8,29] investigate proportional diversification. More precisely, in [8], an election-based method is proposed to address the problem of diversifying searched results proportionally (our work is inspired by this approach). However, this method disregards the similarity (or importance) of the computed set R to the query q and thus may result in including irrelevant objects into R. In [29], this limitation is addressed by considering relevance in the objective function.

9.2.2 Differences

Our problem has a significant difference from the existing related works that renders their straightforward application inappropriate. Related work considers diversity and proportionality of a set of mutually independent results (i.e., S and R sets). Whereas, we aim at finding a diverse/proportional l-sized connected subtree of the OS, which is required to include the root n^{DS} .

10 Conclusion and future work

In this paper, we introduced and investigated the effectiveness and efficiency of two novel types of size-l OSs, namely DSize-l OSs and PSize-l OSs. For this purpose, we employed two types of nodes pairwise relevance, i.e., similarity and equality. We proposed a brute-force algorithm, two efficient greedy heuristics and a preprocessing strategy that restricts processing on only a subset of the OS. We also provide extensive theoretical analysis of these greedy algorithms. Finally, we conducted a systematic experimental evaluation on the DBLP and Google+ datasets that verifies the effectiveness, approximation quality and efficiency of our techniques. The evaluation verified that the two novel snippets are preferred by human evaluators over non-diversified size-l OSs [15]. The evaluation also verified preference for results produced using sim relevance over results produced by using equi relevance that was proposed in [17].

A direction of future work concerns the investigation of inter-diversity and inter-proportionality among a set of query results. For instance, for q, we get three OSs, one per Faloutsos brother; we can diversify M. Faloutsos DSize-l by avoiding information included in the C. Faloutsos DSize-l. Another challenging problem is the combined size-l and top-k ranking of OSs.

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Appendix

Algorithm 4 OS Generation Algorithm

OS Generation (n^{DS}, G^{DS}) 1: $enQueue(Q, n^{DS})$ ightharpoonup Queue Q facilitates breadth-first traversal 2: add n^{DS} as the root of the OS 3: while !(isEmptyQueue(Q)) do $n_i = deQueue(Q)$ **for each** child relation R_i of $R(n_i)$ in G^{DS} do 5: 6: get $R_i(n_i)$ 7: **for each** tuple n_i of $R_i(n_i)$ **do** $enQueue(Q, n_i)$ g. add n_i on OS as child of n_i 10: return OS

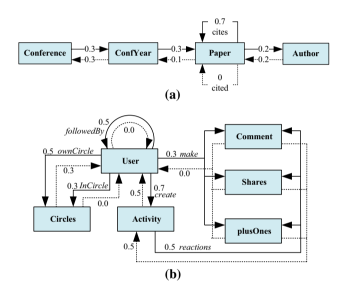


Fig. 17 The G^A s for the DBLP and Google+ datasets. a The DBLP G^A . b The Google+ G^A

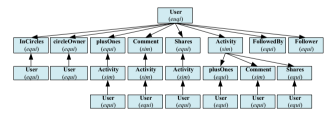


Fig. 18 The Google+ User G^{DS} (relevance type)

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