

Optimal False-Positive-Free Bloom Filter Design for Scalable Multicast Forwarding

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Abstract—Large-scale information dissemination in multicast communications has been increasingly attracting attention, be it through uptake in new services or through recent research efforts. In these the core issues are supporting increased forwarding speed, avoiding state in the forwarding elements and scaling in terms of the multicast tree size. This paper addresses all these challenges – which are crucial for any scalable multicast scheme to be successful – by revisiting the idea of in-packet Bloom filters and source routing. As opposed to the traditional in-packet Bloom filter concept, we build our Bloom filter by enclosing limited information about the structure of the tree. Analytical investigation is conducted and approximation formulae are provided for optimal length Bloom filters, in which we got rid of typical Bloom filter illnesses such as false-positive forwarding. These filters can be used in several multicast implementations, which is demonstrated through a prototype. Thorough simulations are conducted to demonstrate the scalability of the proposed Bloom filters compared to its counterparts.

Index Terms—Bloom filter, source routing, information centric networking, multicast addressing

I. INTRODUCTION

In the current Internet, IP multicast is hardly deployed beyond single providers due to incentive, security and scalability problems. These issues rendered multicast research a cold topic [1] in the last decades. However, recent trends in networking – such as Information Centric Networking (ICN), which is built on a publish/subscribe service model – urges the need of efficient multicast-based architectures. Common to these trends is that they are leaving the issue of *scalable forwarding* as a major challenge, which makes efficient multicast a core requirement for any successful solution in this space. Therefore, multicast routing research is in its second blossom [2]–[4]. The scalability challenge in these networks is driven by three factors, namely

- Goal 1: the need to scale the speed of the forwarding operation with the growing need for speed in the Internet,
- Goal 2: the scalability in terms of state to be maintained in the routers for the forwarding operation, and
- Goal 3: the scalability in terms of supported sizes of multicast trees.

While solutions such as IP multicast generally support multicast trees that span across the Internet, its realization comes at the cost of lookup-based forwarding operations, hindering Goal 1. These operations require appropriately configured entries in forwarding tables within the individual network elements, which clearly hinders Goal 2. On the other hand, recent trends in networking, such as ICN or Software Defined Networking (SDN), are deeply committed to the clear separation of the data and the control plane in order to achieve this goal [5]–[8].

Tackling Goal 2, an appealing approach to minimize, or even avoid state in the network is that of source routing, which is therefore in its second blossom after decades of ignorance in the IP world. Here, instead of encoding next hop information of the multicast tree at the intermediate nodes, as done in traditional approaches, efficiently encoding link information of the graph in a compact header allows for avoiding any state at these intermediary elements. Such design issues manifest in recent efforts towards using in-packet Bloom filters [2]–[4], [9] for encoding the edges of the multicast tree into the packet header. Bloom filters are originally designed for membership queries i.e., for determining whether an element/edge belongs to a set/tree or not. Placed in packet headers, the *in-packet* Bloom filters can effectively address a set of nodes or links [2]. Conceptually, the LIPSIN method in [2] is closest to source-specific multicast (SSM, RFC3569) where the multicast traffic originates from a dedicated source node. However, this flat type of tree representation in LIPSIN (i.e., the structure of the whole tree is represented in an implicit manner as a set of edges) in a single fixed size (256 bit) filter while keeping the number of *false-positive forwardings* – i.e., sending data on a link which was not involved in the multicast tree – low is very limited (about 20 links). Note, that false-positive forwarding e.g. in IPTV has a severe affect as unnecessarily sending a

This work was supported by the High Speed Networks Laboratory at the Budapest University of Technology and Economics. Research of J. Tapolcai and P. Babarczi were partially supported by the Hungarian Scientific Research Fund (OTKA grant K108947). J. Bíró was supported by the TAMOP-4.2.2.C-11/1/KONV-2012-0001 project. The project has been supported by the European Union, co-financed by the European Social Fund. P. Babarczi was supported by the János Bolyai Research Scholarship of the Hungarian Academy of Sciences (MTA). This document has been produced with the financial assistance of the European Union under the FP7 GÉANT project grant agreement number 605243 as part of the MINERVA Open Call project.

high-quality video stream on a link highly degrades network performance. Thus, a single fixed length header like LIPSIN clearly not suitable to reach Goal 3.

In this paper we show that Goal 3, i.e., the scalability problem in terms of multicast tree size can be tackled by using some topology-related information when composing the in-packet Bloom filter, which was not present in the previous approaches. Although the excellent scalability of our solution would make it suitable for Internet-wide adoption, some technological concerns (e.g. simultaneous deployment of the required functionality at networking hardware) render an incremental domain-by-domain deployment scenario much more realistic. This promotes the thorough intra-domain investigation of the problem, which will be made in this paper. First, we propose the so called *multi-stage Bloom filter (MSBF)* which consists of consecutive Bloom filters encoding only the membership of the edges residing at a given hop-distance (*stages*) from the source. Second, optimal (varying) length *false-positive-free* Bloom filters are introduced in order to eliminate false-positive forwardings, which is of utmost importance in any practical realization of in-packet Bloom filter based multicast forwarding. Our analytical investigations and simulations prove that our MSBF solution not only provides excellent space efficiency but it also remedies besides false-positive forwarding several other anomalies arising when Bloom filters are at use, like forwarding loops and packet storms [4].

The rest of the paper is organized as follows. Section II expands on the related work in this area. In Section III, we define the architectural background in which we embed our solution and introduce the concept of our new multicast architecture. As the main contribution of this paper, the analytical results on creating minimal length false-positive-free Bloom filters are presented in Section IV. In Section V we discuss the implementation details of multi-stage Bloom filters in SDNs, IP routers and in an ICN prototype followed by Section VI, where we present our simulation and measurement results on the proposed false-positive-free scalable multicast architecture. Finally, Section VII concludes the paper.

II. RELATED WORK

A. Multicast Routing

Standard IP multicasting can provide a way of transmitting packets from a source to many recipients in a bandwidth economical way. Multicast routing protocols like DVMRP [10], [11] and PIM-SM [12] for performing the transmission maintain Multicast Forwarding Tables (MFT) in the routers along the multicast trees. This stateful approach is not only in full contradiction of the stateless design of unicast IP, but is also the Achilles' heel of IP multicast. Due to the potentially high number of multicast groups in the network according to massive multi-party applications like video conferencing or networked games, the states to be maintained in a router could be untenable due to the unaggregability of the MFT's.

For better scalability in terms of Goal 1 explicit multicast solutions have been suggested in the literature, in which

forwarding information (e.g. list of destination IP addresses) on the targeted group is encoded in the packet headers. This flat type of stateless multicast routing protocols (like Xcast, Xcast+ [13]) undoubtedly has the drawback that every intermediate node along the multicast tree should process the header, even in the case the node is not a branching point of the tree. To avoid mandatory packet header processing, tree encoding schemes have been proposed to integrate into multicast protocols, like in ERM and Linkcast [14], [15]. In these methods, the entire tree structure is encoded into the headers, moreover, in the latter case (and its successors [13]) forwarding at intermediate nodes is based on interpreting the tree code, unicast lookups are completely eliminated. Nevertheless, the tree encoding scheme in Linkcast uses fixed length link codes by the assumption on the maximum size of multicast groups and imposes large computational overhead when interpreting the encoded tree.

B. In-packet Bloom Filters

The *Bloom filter* [16] is a simple yet efficient data structure to answer membership queries. Let \mathbb{S} be a set of elements, assigned with b bit long binary codes, in which a maximum of k bits are set in positions indicated by k different *hash functions*. Each of the k hash functions maps the given element onto one of the b bit positions. The hash functions are assumed to be independent and each position is selected with equal probability. The Bloom filter representing $\mathbb{T} \subseteq \mathbb{S}$ is a b bit long binary array consisting of the bitwise OR of the codes of the elements in \mathbb{T} . As the main feature a membership test can simply be performed by checking if all bit positions that are set in the code of the underlying element s are also set in the filter. The performance of the filter is measured by the false-positive rate that is the probability that an element s appears to be included (by test) but actually it is never added ($s \in \mathbb{S} \setminus \mathbb{T}$).

The application of Bloom filters is becoming increasingly popular in future Internet architectures [2], [3] for tackling Goals 1 and 2. Placed in packet headers, the *in-packet Bloom filters* can effectively address a set of nodes or links, hereby qualifying themselves as a strong candidate solution for efficient stateless multicast addressing. When a packet with in-packet Bloom filter arrives to a router, membership testing is performed on the outgoing link identifiers. The bitwise AND and compare (CMP) operation on the Bloom filter placed in the header and on the address of the outgoing link is extremely fast when implemented on a digital signal processor, which leads to a simpler router architecture compared to current IP routers. Realizations of this solution, e.g. in [7], demonstrate that 1 GB/s throughput can be easily achieved in prototypes. Moreover, *Bloom filters are favored for their space efficiency* since the filter requires much less space than listing the identifiers for each links/hosts in the multicast tree. Furthermore, this allows routers to stay quasi-stateless, because the routers only need to know the address of the neighboring nodes and links.

A clear weakness of such an in-packet Bloom filter concept is the total ignorance of

- (i) the topology information and
- (ii) the tree structure in addressing.

In [4] the first issue (i) was touched by generating the Bloom filter addresses of each link according to the number of adjacent links, which is intuitively beneficial because the links with fewer adjacent links has a smaller chance to give false-positive. However, it still not able to fully support Goal 3, which is addressed in this paper.

C. Our Contribution

In this paper, we propose an addressing scheme for arbitrary size trees and multicast groups which lies somewhat between the two extreme approaches of explicit routing, i.e., the completely flat solution and the complete tree encoding based ones, and suits well for the requirements of ICNs by tackling Goal 3. In our solution link identifiers at equal hop-distances from the source of the multicast tree (referred to as *stages* hereafter) are to be confined into Bloom filters, but without any exact encoding of the tree topology hereby providing an appealing mixture of space and forwarding efficiency.

Furthermore, we address both weaknesses (i) and (ii) of in-packet Bloom filters by introducing (i) optimal length varying size (ii) multi-stage Bloom filters, as opposed to the traditional (single stage) fixed size in-packet Bloom filter concept [2] (which represent the trees flatly as sets in a fixed length header). The idea of multi-stage Bloom filters was first introduced in [17], followed by an ICN prototype implementation in [18]. However, these works contained nor clear guidelines for the proper dimensioning of the filters from the theoretical side, neither any tool for processing the required parameters (e.g. the number of hash functions) in an implementation. Thus, in this paper we also introduce the concept of false-positive-free Bloom filters and provide approximation and analytical results for filters with minimal length which can be directly used in practice. The effectiveness and applicability of the derived approximation formulae are demonstrated through simulations and in our ICN prototype implementation as well.

III. ARCHITECTURAL BACKGROUND

For the reasons discussed earlier assuming a domain-by-domain introduction of our MSBF approach, in the rest of the paper we assume that information is disseminated across a single domain. As *link identifiers are used instead of end-point addresses*, from an overall network design perspective, an important consideration is the scheme being used for encoding the link information. Usually the usage of fixed size identifiers is favored, enabling line-speed execution of the forwarding operation (see the 256 bit long identifier in [2] for an example). Such fixed size, however, limits the ability to appropriately encode any given size tree within the limited size of the identifier. While varying size header approaches have been clearly at a disadvantage compared to fixed size approaches, it is our contribution in this paper to provide a solution that closes this gap in possible forwarding speed while providing the previously outlined design advantages. For this, we contrast our work against existing source routing

approaches and provide insight into the performance and implementation issues of our scheme.

Crucial to our approach is the knowledge of the overall topology over which the multicast tree is formed. Such topology knowledge is not unreasonable to assume, as can be seen in technologies like Multi-Protocol Label Switching (MPLS) or software defined networks as well as in proposals envisioning future information-centric solutions. Common here is the existence of a topology manager (or network controller) with knowledge of link information between forwarding elements under the management of this entity. It is the role of this topology manager to compute an appropriate forwarding header (MSBF in our multicast architecture) that can be used within its topology. We argue that the existence of such topology management in today's networks points towards the possibility to improve information delivery in existing IP networks as well.

A. Architectural Concept of Information Centric Networks

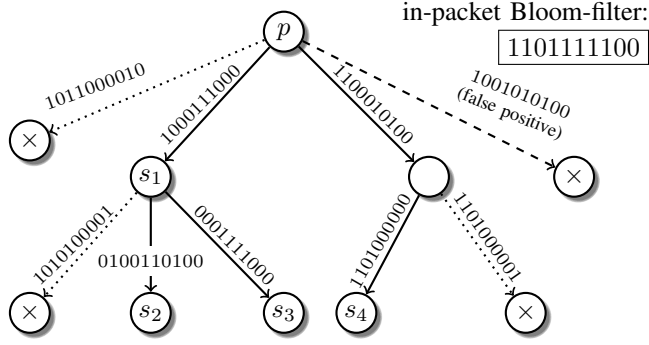
The architectural context of information-centric networks is based on the work in [7], [19]. Here we summarize the main design principles onto which we set the foundations for our proposed addressing scheme:

Spatially and temporally decouple communicating parties. A publish/subscribe service model is exported to all applications and ancillary network components. Hence, the producer of information (publisher) does not need to coexist in time with the consumers (subscribers), i.e., a subscriber can receive information even if the initial information producer is not online, since the information can be replicated or cached throughout the network. Moreover, communicating entities do not know the location of each other, spatially decoupling them.

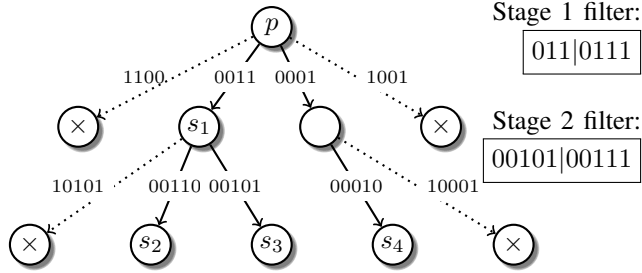
Clearly separate network functions. As discussed in [19], the core network functions are cleanly separated with each node supporting three network functions. The first one, *rendezvous*, matches demand for and supply of information. This results in some form of (location) information that is used for binding the provisioning of information to a network location. This information is used by the second function, *topology management and formation*, to determine a suitable multicast tree for the transfer of the information, this transfer being executed by the third function, *forwarding*.

B. Traditional Bloom Filter Based Forwarding (ISBF)

Based on the previous discussion on the ICN architecture, similarly to previous works on Bloom filters [2]–[4], [9], the existence of the *rendezvous* point is assumed in the network. However, it is important to note that this function can be easily implemented in a distributed fashion, e.g., target nodes subscribe to source nodes directly. Similarly to other source routing approaches we assume that the up-to-date topology of the network is available at the *topology manager*. It is also the task of this topology manager to compute the multicast trees (or DAGs) for each multicast group after consulting with the rendezvous point. Following the architecture of [2] the links are assigned by binary addresses and the topology manager



(a) Traditional (fixed length) in-packet Bloom filter with false-positive forwarding, owing to the large number of $\alpha = 5$ in-tree and $\beta = 4$ out tree links.



(b) Multi-stage (varying length) Bloom filter on the same topology without false-forwarding. At Stage 1 $\alpha_1 = 2$ and $\beta_1 = 2$, while at Stage 2 $\alpha_2 = 3$ and $\beta_2 = 2$.

Fig. 1. Traditional and multi-stage in-packet Bloom filters at the forwarding phase (AND & CMP at each interface) on a multicast tree with four subscribers s_1, s_2, s_3 and s_4 .

computes the multicast packet headers by combining the addresses of the links residing in the corresponding multicast tree. In order to better understand the details of the proposed multi-stage filter let us first recall how this computation is done using the traditional in-packet Bloom filters.

- Each link is assigned by a binary link address (consisting of b bits of which at most k are set to 1).
- The topology manager computes the multicast header by bitwise OR-ing the addresses of the links in the corresponding multicast tree.

At each router outgoing link addresses are tested against the Bloom filter (i.e., testing at each bit-position where the link address is set if the filter is also set with a binary AND and CMP function) and forwarded if positive. Note that this approach is totally topology unaware since it simply traces back the forwarding decision to a classical membership testing problem. As an example in the multicast tree of Fig. 1a the in-packet Bloom filter contains the bitwise OR of 5 link addresses (with fixed length $b = 10$ and $k = 4$ hash functions). The filter is tested 9 times (against every outgoing interface) during the forwarding process, and producing false positive forwarding even with such low number of links.

C. Varying Length False-Positive-Free Multi-Stage Bloom Filters (MSBF)

In-packet Bloom-filters are an efficient representation of the links of a multicast tree in an ICN context. However,

it was shown in [2] that the multicast tree size which can be efficiently represented in a single fixed size filter while keeping the false-positive rate low is very limited (about 20 links), contradicting with Goal 3.

Our contribution in this paper is to use a sequence of shorter, so called *stage filters* instead of a single large one. Each stage filter contains only the forwarding information for the edges with the same hop-distance from the source node in the multicast tree. Thus, we improve the completely flat addressing structure of the original in-packet Bloom filter concept, and some topology-related information is added to the packet header through the stage filters for tackling Goal 3. The efficiency of this approach will be demonstrated analytically in Section IV, while empirically in Section VI.

Another desired property of the introduced *Multi-Stage Bloom Filters (MSBF)* approach is that when forwarding a packet the *already used stage filters can be truncated* at each hop one-by-one, as it contains only information for the already traversed links. That is, a tree of h hops is represented by h stage Bloom filters, where the i^{th} one contains only the links residing at i hop-distance from the source. When leaving the source the multicast in-packet Bloom filter header consists of h stage filters, which then shrinks as the packet travels along the tree, highly reducing the overhead introduced by the in-packet Bloom filter.

In ICNs and also in traditional multicast applications, unnecessarily forwarding a high-bandwidth video stream on a link could result in degraded network performance owing to congestion. In terms of security and business considerations, it is also not desirable if a node receives a stream unintendedly. Thus, in order to keep the number of false-positive forwardings on a tolerable level in arbitrary size multicast trees, applying *false-positive-free (FPF) varying length stage filters* could be a solution, discussed in Section IV. For ensuring space efficiency, the length of the FPF filters are optimized at each stage as the function of the number of elements (i.e., links) it contains, which clearly results in varying size stage filters. Thus, for identifying filter boundaries we propose to store the length of each filter in the header e.g. by applying the commonly used Elias gamma universal code [20], where the length γ of a b bit long filter is coded as follows (notations can be found in Table I):

- first $\lceil \log_2(b) \rceil - 1$ zero bits are written,
- followed by the binary representation of b on $\lceil \log_2(b) \rceil$ bits.

As it is shown in the example of Fig. 1b, the packet header at Stage 1 consists of two parts, a $\gamma_1 = 3$ bit long Elias gamma code, and a $b_1 = 4$ bit long Bloom filter. This filter at Stage 1 is tested against four links, out of which two are expected to be chosen. Similarly, at Stage 2 the second filter ($\gamma_2 = 5, b_2 = 5$) is tested five times with three expected positive outcomes.

The benefit of our MSBF approach with respect to the introduced overhead is clear, for example, if we investigate Fig. 1b. Note that, the average Bloom filter length is $b_{avg} = 10$ for 1SBF, while $b_{avg} = 6.6$ for MSBF. The average overhead bits for our MSBF approach is maximally $m_{avg} = 17$, while

applying the header truncation mechanism it could be reduced to $m_{avg} = 12.8$ (as the number of overhead bits is 17 on the two links of Stage 1, while 10 for the three links of Stage 2). One can observe that even on this small multicast tree (where ISBF still performs well), the number of extra overhead bits paid for varying length multi-stage filters is negligible (12.8 in contrast with 10 of the ISBF approach), while with the application of the MSBF filter false-positive forwarding was completely eliminated.

Note that the main idea behind eliminating false positive forwarding is to move the uncertainty from the forwarding phase to the Bloom filter generation phase, i.e., generating an FPF filter can be considered as a randomized algorithm (with controllable uncertainties). However, once an appropriate filter is found, the forwarding will be surely FPF, i.e., the forwarding phase is deterministic. Thus, in order to characterize the properties of the FPF filter generation process, in Section IV we will derive and analyze the distribution and the expected value of FPF filter length in several settings, and provide some practical guidelines which can be used in an implementation.

IV. ANALYSIS AND DESIGN OF FPF BLOOM FILTERS

In case of the original Bloom filter, elements of a certain subset \mathbb{T} of a much larger (often unknown) set \mathbb{S} can be collected (encoded) into the filter in such a way that membership queries can justify the inclusion of the elements of \mathbb{T} . Unfortunately, other elements from $\mathbb{T} \setminus \mathbb{S}$ can also happen to be tested positively, although they were never added. The frequency of the events of these false reportings of the membership test is called *false-positive probability*. The design goal is to generate a Bloom filter for \mathbb{T} which can provide the false-positive probability under a prescribed value. After generation, the operation of such a filter can be erroneous; however, the false-positive rate can be kept under a small value.

As opposed to this, in multicast communication both $\mathbb{T} = \mathbb{L}$ and $\mathbb{S} = E$ are known and their cardinality are quite comparable. This fact may be utilized in the filter generation process in such a way that after generating a filter for \mathbb{L} , all the elements of $E \setminus \mathbb{L}$ can be tested against the false inclusion. If the inclusion of one or more of the elements of $E \setminus \mathbb{L}$ are reported, then another filter can be generated for \mathbb{L} and tested against for false inclusion again. In principle, it is possible to find a filter with appropriate length which does not contain any of the elements of $E \setminus \mathbb{L}$. As opposed to the original Bloom filter, such a filter will be false-positive-free in operation and thus inherently suitable for error-free multicast forwarding.

The generation of such filters at the topology manager for any given multicast tree \mathbb{L} is not a trivial issue and will be discussed in this section. Shortly foreshadow this, the main tasks in an FPF filter generation can be summarized as follows:

- computing expected filter lengths based on the number of elements α to be included in and the number of elements β to be excluded from the filter (called **Scalability**, discussed in Section IV-A and Section IV-B),

TABLE I
NOTATION LIST

Notations	Description
$G = (V, E)$	the directed graph representation of the topology with nodes V and edge set E
\mathbb{L}	multicast tree
α	number of in-tree links ($= \mathbb{L} $)
β	number of out-tree links ($\leq E \setminus \mathbb{L} $)
b	Bloom filter length
γ	total length of the Elias-gamma function in the varying length in-packet Bloom filters
$m_l = \gamma_l + b_l$	total overhead bits at link (or stage) l
m_{avg}	average number of overhead bits per link in multicast tree \mathbb{L}
$\eta(\mathbb{L})$	filter compactness, m_{avg} divided by the number of links in the multicast tree
b_{avg}	average number of in-packet Bloom filter bits per link in multicast tree \mathbb{L}
$\lambda(\mathbb{L})$	Bloom filter compactness, b_{avg} divided by the number of links in the multicast tree
δ	test range where FPF Bloom filters are most probable in a practical implementation ($[E[L] - \delta, E[L] + \delta]$)
h	number of stages (maximal hop distance from source)
L	random variable representing the Bloom filter length
$\bar{L}(\alpha, \beta)$	approximation of the expected FPF Bloom filter length
k^{opt}	optimal (real) number of hash functions
$k^*(\alpha, \beta)$	approximation (integer) number of hash functions
\mathcal{D}	set of multicast trees given in the network
η_{avg}	average filter compactness for all multicast trees \mathcal{D} (with header abbreviation, if MSBFs are considered)
μ_{avg}	average filter compactness for all multicast trees \mathcal{D} (without header abbreviation, if MSBFs are considered)
λ_{avg}	average Bloom filter compactness for all multicast trees \mathcal{D}

- computing/estimating the required test range δ based on the expected filter length (called **Insensitivity**, discussed in Section IV-C),
- testing the filter lengths in increasing order in the test range against the FPF property, stop when the filter tested is FPF using the appropriate number of hash functions (called **Implementability**, discussed in Section IV-D).

We will also demonstrate that breaking down the Bloom filter into stages with optimal length can significantly reduce the overall filter length b compared to

- the traditional (single stage) Bloom filter with optimal length, and
- to the LIPSIN approach using fixed length in-packet Bloom filters.

The notations are summarized in Table I.

A. The Expected Length of the False-Positive-Free Bloom Filter

Let α and β denote the number of in-tree (which are contained in the multicast tree \mathbb{L} , denoted by bold lines in Fig. 1) and out-tree (which are not contained in the multicast tree $E \setminus \mathbb{L}$, dotted lines in Fig. 1) links in a given stage, respectively, and we assume that there are h stages. For the sake of simplicity, we conduct our analysis for identical α and β values in each stage. However, in Section VI general settings are used.

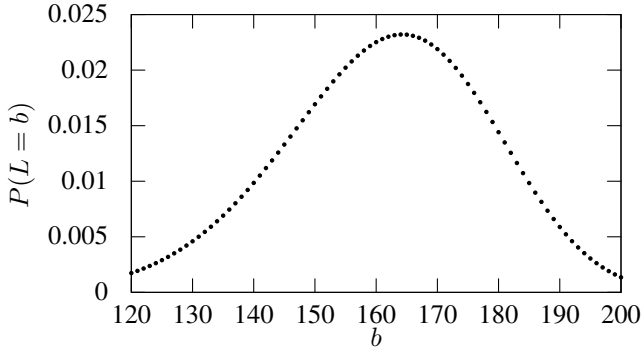


Fig. 2. The probability distribution of FPF filter lengths for $\alpha = 30$ and $\beta = 40$ ($E[L]^{MS} = 161.2$).

TABLE II
COMPARISON OF TRADITIONAL (1S) AND MULTI-STAGE (MS) BLOOM FILTERS IN EXPECTED FILTER LENGTH WHEN $\alpha_i = 10$, $\beta_i = 30$, $i = 1, \dots, 5$.

No. of hops	$E[L]^{1S}$ from Eq. (8)	$E[L]^{MS}$ from Eq. (9)
1	54.31	54.31
2	127.94	108.62
3	210.01	162.93
4	297.69	217.24
5	389.61	271.55

First we recall how to design an optimally sized Bloom filter to meet a prescribed false-positive requirement [21]. Without loss of generality, the false-positive probability for a Bloom filter with length b can be calculated as:

$$P_f = \left(1 - \left(1 - \frac{1}{b}\right)^{\alpha k}\right)^k. \quad (1)$$

Introduce the notation $p = \left(1 - \frac{1}{b}\right)^{k\alpha}$ and minimizing P_f using the first derivative transforms to

$$\frac{\partial P_f}{\partial k} = (1-p)^{k-1} \cdot [(1-p) \cdot \ln(1-p) - p \ln p] = 0. \quad (2)$$

It is easy to see that the optimal k , that is the optimal number of hash functions is attained when $p = 0.5$:

$$k^{opt} = \frac{1}{\alpha} \frac{\ln(0.5)}{\ln(1 - \frac{1}{b})}, \text{ and } P_f^{opt} = 0.5^{\frac{\ln(0.5)}{\alpha \ln(1 - \frac{1}{b})}}, \quad (3)$$

or using

$$\left(1 - \frac{k}{b}\right)^{\alpha} \approx e^{-\frac{k\alpha}{b}} \quad (4)$$

we get the approximation

$$P_f^{opt} \approx 0.5^{\frac{-b \ln 0.5}{\alpha}} = c^{\frac{b}{\alpha}}, \text{ where } c = \frac{1}{2^{\ln 2}}. \quad (5)$$

In the following the length of false-positive-free Bloom filters is discussed, i.e., a large enough filter is created for which the expected number of false-positives will be zero. For this let us define the following probability:

$$P(b, \alpha, \beta) = \left(1 - c^{\frac{b}{\alpha}}\right)^{\beta}. \quad (6)$$

This expresses the probability, that in a b length filter containing α elements none of the β out-tree links are included in the filter. The probability distribution of the shortest FPF filter length L is shown in Figure 2, which can be calculated as follows:

$$P(L=b) = P(b, \alpha, \beta) \prod_{i=1}^{b-1} (1 - P(i, \alpha, \beta)). \quad (7)$$

Now the length L of the false-positive-free Bloom filter comes from the random process of trying with increasing size filters and stop when false-positive cannot be found. That is this expected length in case of the traditional (single stage) filter is

$$E[L]^{1S} = \sum_{b=1}^{\infty} b \cdot P(b, h\alpha, h\beta) \prod_{i=1}^{b-1} (1 - P(i, h\alpha, h\beta)), \quad (8)$$

because there are $|\mathbb{L}| = h\alpha$ links to be included and $E \setminus \mathbb{L} = h\beta$ links are to be excluded in the filter. The expected length of the false-positive-free multi-stage filter is

$$E[L]^{MS} = h \sum_{b=1}^{\infty} b \cdot P(b, \alpha, \beta) \prod_{i=1}^{b-1} (1 - P(i, \alpha, \beta)), \quad (9)$$

because there are h identical false-positive-free filters (each containing α links and not containing β links) confined in the stage filter. Using the above formulae Table II shows a comparison between the expected length of the traditional and multi-stage Bloom filters, when $\alpha_i = 10$, $\beta_i = 30$ and the number of hops increases from 1 to 5. Evidently for a single hop ($h = 1$) Eq. (8) and Eq. (9) results the same expected value 54.31, as the number of α and β links are the same. However, notice that the improvement due to using more stages can be stunning even if the number of stages is relatively small, for trees with larger depth 30-40% improvement can be reached.

Although the expected length of FPF Bloom filter can not be used directly for filter generation (according to Figure 2 the filter with length of the expected value will be false-positive-free with only about 2% probability¹), it plays an important role in finding an appropriate minimal length for obtaining a false-positive-free filter, hence it is worth being analyzed. Based on this analysis, a test region of lengths (δ) around the expected value can be determined, in which the false positive free filter can be found with very high probability. For example, according to the minimal length distribution in Figure 2, the minimal length of the FPF Bloom filter can be found in the region $[80, \dots, 240]$ with probability $1 - 10^{-5}$.

B. Approximations and Analysis of Minimal Filter Length

In this subsection we provide analytical approximations for the probability distribution and the expected value of the minimal length of the false-positive-free Bloom filters. Based on these we analyze how these quantities depend on α and β , and also provide quantification on improvements in multi-stage filters against the single-stage FPF Bloom filter.

¹This also means that the filter length generated by the FPF filter generation process may largely deviate from the expected FPF filter length.

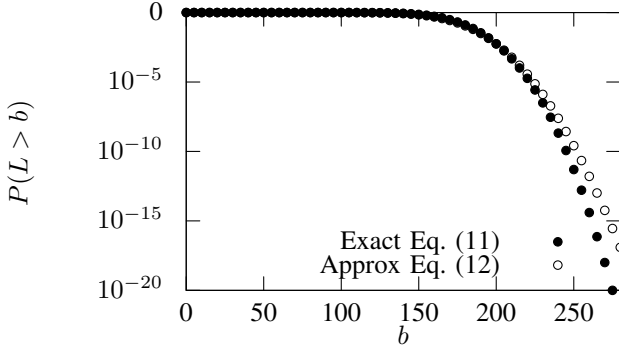


Fig. 3. The tail probability and its approximation for $\alpha = 30$, $\beta = 40$.

As previously presented in Eq. (6)-(7) the distribution of the filter length L can be characterized as

$$P(L = b) = (1 - c^{\frac{b}{\alpha}})^{\beta} \prod_{i=0}^{b-1} (1 - (1 - c^{\frac{i}{\alpha}})^{\beta}). \quad (10)$$

Based on this the tail of the distribution (CCDF) of the filter length L can be written as

$$P(L > b) = \prod_{i=0}^b (1 - (1 - c^{\frac{i}{\alpha}})^{\beta}). \quad (11)$$

First we show a useful approximation for this tail probability.

Lemma 1:

$$P(L > b) \approx \text{Exp} \left(-\frac{\alpha \text{Ei}(-c^{b/\alpha} \beta)}{\ln(c)} \right), \quad (12)$$

where $c = \frac{1}{2^{\frac{1}{\ln 2}}}$ and $\text{Ei}(z) = -\int_{-z}^{\infty} \frac{e^{-t}}{t} dt$ is the so-called exponential integral function.

The proof is relegated to Appendix. Extensive numerical investigation showed that this tail probability approximation is accurate enough for practical size problems $\alpha = (2, \dots, 100)$, $\beta = (10, \dots, 1000)$, and in the region $1, \dots, 10^{-10}$ of $P(L > b)$. In Fig. 3 one can observe good coincidence of the curves, and even in the range under 10^{-10} the "horizontal"² relative difference between the curves are under few percent.

In what follows, an accurate approximation is presented on the expected filter length $E[L]$ (either for $E[L]^{1S}$ or $E[L]^{MS}$). In principle, $E[L]$ can be expressed as the sum of the tail probabilities as $E[L] = \sum_{b=0}^{\infty} P(L > b)$. Unfortunately, it seems that it can not be directly used as an integrate approximation sum, because the integral $\int_{x=0}^{\infty} \text{Exp} \left(-\frac{\alpha \text{Ei}(-c^{x/\alpha} \beta)}{\ln(c)} \right) dx$ turned out being not characterizable by using elementary and/or special functions. Instead, we utilize the property of the CDF $P(L = b)$ that it is close to a symmetric distribution, therefore, the median b^* is an acceptable approximation for

²From practical point of view, the "horizontal" relative difference has higher significance than the "vertical" one. That is for given prescribed tail probability ε the smallest b and \tilde{b} for which $P(L > b) \leq \varepsilon$, $P(L > \tilde{b}) \leq \varepsilon$ are to be find and compared as a "horizontal" relative difference $\frac{\tilde{b}}{b} - 1$.

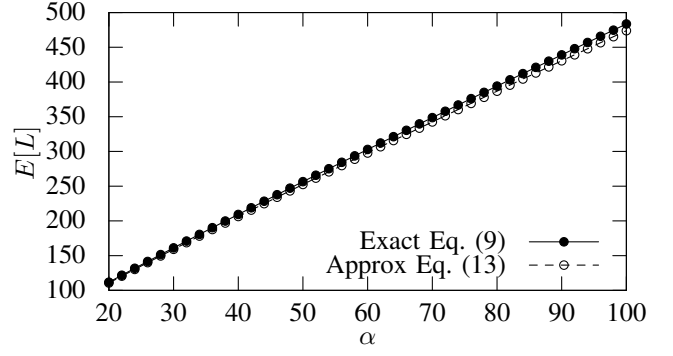


Fig. 4. The expected filter length and its approximation for $\alpha = (20, \dots, 100)$, $\beta = 40$.

the expected value, for which $P(L > b^*) = 1/2$. The next theorem presents the main result of this subsection:

Theorem 1: The expected false-positive filter length can be approximated by the following formula

$$E[L] \approx \frac{\alpha \ln \left(-\frac{\beta}{\text{Ei}^{-1} \left(-\frac{\ln^3(2)}{\alpha} \right)} \right)}{\ln^2(2)} =: \tilde{L}(\alpha, \beta), \quad (13)$$

where $\text{Ei}^{-1}(\cdot)$ is the inverse function of the left branch³ of $\text{Ei}(\cdot)$.

The statement of the theorem immediately follows from Lemma 1 and the symbolic solution of the equation with respect to b

$$\text{Exp} \left(-\frac{\alpha \text{Ei} \left(-\left(\frac{1}{2^{\frac{1}{\ln 2}}} \right)^{b/\alpha} \beta \right)}{\ln^2(2)} \right) = \frac{1}{2}. \quad (14)$$

The approximation $\tilde{L}(\alpha, \beta)$ is worth rephrasing as

$$\tilde{L}(\alpha, \beta) = \frac{\alpha}{\ln^2(2)} (\ln \beta - s(\alpha)), \quad (15)$$

where $s(\alpha) = \ln \left(-\text{Ei}^{-1} \left(-\frac{\ln^3(2)}{\alpha} \right) \right)$ is a very slowly increasing function of α ($s(\alpha)$ is approximately proportional to $\ln(\ln \alpha)$). This mean filter length approximation is in very good agreement with the "exact" average length numerically computable by Eq. (9) or $\sum_{b=0}^{\infty} P(L > b)$. Two figures are presented for illustration, one is on the α dependence with fixed β and the other is on the β dependence with fixed α (Fig. 4 and Fig. 5)⁴.

Now, the main message of Theorem 1 is that the expected false-positive-free filter length is approximately proportional to the number of elements included (α), and to the **logarithm** of the number of elements excluded (β). This is referred to as the property of **Scalability**.

³ $\text{Ei}(\cdot)$ has a branch cut discontinuity at 0, the left branch (with negative arguments) is a negative valued function.

⁴Note that in Fig. 5 the horizontal axe with β has logarithmic scale

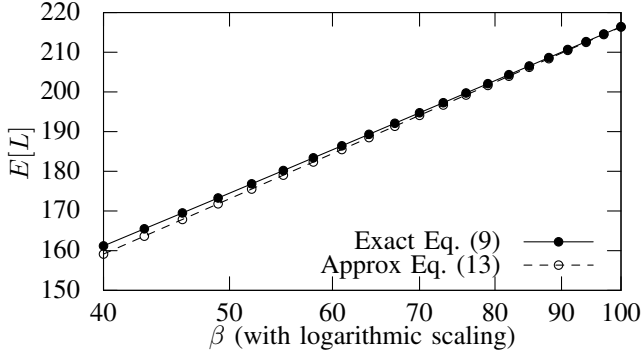


Fig. 5. The expected filter length and its approximation for $\alpha = 30$, $\beta = (40, \dots, 100)$ (log-linear plot).

The formula of the expected filter length can also be used to quantify the expected gain in filter length using multi-stage filters against a single one-stage filter as

$$\tilde{L} \left(\sum_{i=1}^h \alpha_i, \sum_{i=1}^h \beta_i \right) - \sum_{i=1}^h L(\alpha_i, \beta_i). \quad (16)$$

Taking the simple example of the previous section ($h = 5$ and for each stage $\alpha = 10, \beta = 30$) this formula gives 121.9 bits, while the previous numerical calculation (see the last row of Table II) provided 118.06 bits as expected gains.

C. Approximation of The Required Test Range

From a practical point of view, it is important to find the optimal length FPF Bloom filter as fast as we can. It is straightforward to search for a minimal length FPF Bloom filter in $[1, E[L]^{MS}]$ with increasing filter length. However, it requires a lot of unnecessary computations as from Figure 2 one can observe that the most probable range to find an FPF filter is around the expected value $E[L]^{MS}$. In this section, we are interested in the size of this *test range* δ , i.e., the test interval $[E[L]^{MS} - \delta, E[L]^{MS} + \delta]$, where it is practical to seek an FPF filter, as a simple and accurate enough approximation can also be given for the required test range based on the CCDF (tail probability) approximation in Lemma 1. If the prescribed success rate for the test range is $1 - \varepsilon$, then we find b_1 and b_2 such that the probability of the lower tail ($1 - P(L > b_1)$) and the upper tail ($P(L > b_2)$) of the distribution should be at most $\varepsilon/2$. Using our tail probability approximation, the required test range is about

$$b_2 - b_1 \approx \frac{\alpha}{\ln^2(2)} (s(\alpha, \varepsilon/2) - s(\alpha, 1 - \varepsilon/2)), \quad (17)$$

where $s(\alpha, x) = \ln \left(-\text{Ei}^{-1} \left(\frac{\ln^2(2) \ln x}{\alpha} \right) \right)$.

With $\alpha = 30$ and $\beta = 40$ the required test interval is 160 with $1 - 10^{-5}$ (99.999%) success ratio. The formula in Eq. (17) gives us 159. On Fig. 6 one can observe the small differences between (half of the) test interval (previously noted by δ) and its approximation vs. the success rate, with the same α and β . Based on extensive numerical investigations, we observed that the analytical formula of the test range provides

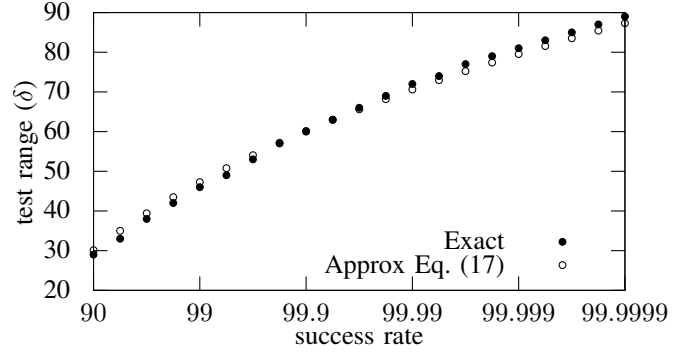


Fig. 6. The required test range and its approximation with success rate $10^{-1}, \dots, 10^{-6}$.

good approximation when $\beta > \alpha$ (i.e., the multicast tree includes less links than it excludes, which is true in most practical scenarios). Moreover, for these cases the numerical computation reveals that the test range hardly depends on β . For example, if β is increased from 40 to 4000, in the previous example the test range increase from 160 to 174. Note that this is also reflected by Equation (17) in which the approximation of the test range $b_2 - b_1$ does not depend on β .

*The main message of this subsection is that the required test range for finding the minimal false-positive-free filter length is approximately **independent** from the number of elements excluded. This is referred to as the property of **Insensitivity**.*

D. Approximation of the Number of Hash Functions

Up until this point in the optimized Bloom filter and hence in the analysis of the false-positive-free Bloom filter the number of hash functions is implicitly treated as a positive real-valued number (Eq. (3)), nevertheless, in implementations it should be positive integer number. Recall that the optimal number of hash functions (in case of an b length filter with α number of elements to be included) is

$$k^{opt}(\alpha, b) = \ln(0.5) \frac{1}{\alpha \ln(1 - \frac{1}{b})} \approx \ln(2) \frac{b}{\alpha}. \quad (18)$$

With given parameters, this value should be rounded to a positive integer number. In finding the one with the minimal length among the false-positive-free Bloom filters, the rounding means that for every length to be tested a rounded value of k^{opt} should be used. Based on thorough numerical assessment of the effect of this rounding procedure, we can state that all the numerically computable formulae and their analytical approximations for the required test range and the expected filter length change negligibly. Moreover, it can also be demonstrated that it is enough to use a *single* integer number of hash functions, and a suitable choice for this is

$$k^* = \text{Round} \left(\ln 2 \frac{E[L]}{\alpha} \right) \approx \text{Round} \left(\frac{1}{\ln 2} (\ln \beta - s(\alpha)) \right), \quad (19)$$

due to the approximation $\tilde{L}(\alpha, \beta)$ in Eq. (15). As an illustrative example for the robustness of the filter length calculation

against the rounding effect is as follows: The expected minimum filter length is 161.2 for $\alpha = 30$, $\beta = 40$ based on Eq. (9) and its approximation is 159.1 based on Eq. (15) (using real-valued numbers of hash functions). The rounded number of hash functions for this length is $k^* = 4$. Recalculating the expected filter length with this fixed number of hash functions is 162.4. Moreover, if we use 3 instead of 4 for the number of hash functions, the expected length is 162.1. Intuitively, this may be due to the fact that around the mean value there are the most probable minimal filter lengths having principle impact on the mean-value calculation, and in this region the rounded k^{opt} may be constant or change its value only once. The very similar observations can also be performed for the required test range.

In Fig. 7a the contour plot of $k^*(\alpha, \beta)$ are presented on the α, β plane. It can be observed that for a given β , k^* remains intact through a wide range of α . It is also confirmed by the slow increase of the $s(\alpha)$ function. The contours represent the implicit function $\beta(\alpha)$ with fixed k^* , which is approximately logarithmic due to the observation that $s(\alpha) \sim \ln(\ln \alpha)$. In Fig. 7b the contour plot of the expected filter length $E[L]$ is presented on the same plane. It can be observed that for fixed expected length β is quickly decreasing when α is increasing, that is, a filter with given length could exclude fewer elements if the number of elements to be included is larger. It is also justified by the $\tilde{L}(\alpha, \beta)$ approximation formula, because from this β is approximately $\sim \text{Exp}(-\alpha)$ if \tilde{L} is fixed. In Fig. 7c the ratio of the number of hash function k^* and the expected filter length $E[L]$ is plotted. From the previous figures and considerations one may expect that this quantity does not significantly depend on β for a given α . This is acknowledged by the contours of the figure, which are almost "vertical".

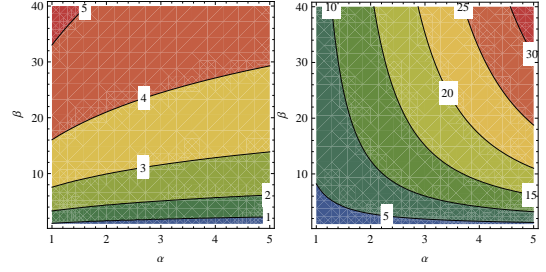
*The main message of this subsection is that the expected filter length and the required test range remain intact when the unimplementable real (continuous-valued) numbers of hash functions is replaced by a single integer number (therefore implementable) of hash functions. This is referred to as the property of **Implementability**.*

V. IMPLEMENTATION ISSUES ON BLOOM FILTER BASED FORWARDING

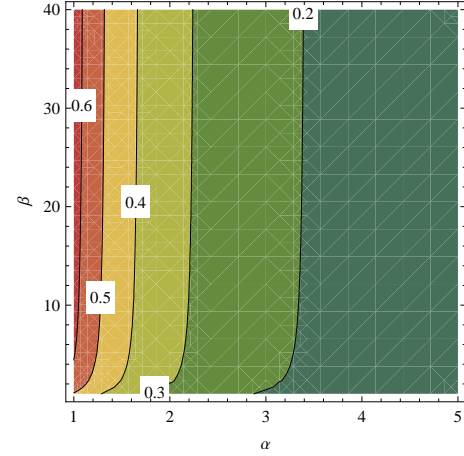
In Section III-C an efficient multicast addressing mechanism in information centric networks, called multi-stage Bloom filters, was introduced. The optimal length MSBF header was calculated at the topology manager based on publisher/subscriber information of the rendezvous function using our results in Section IV. In this section, the implementation issues of Bloom filter based forwarding are discussed in several architectures and environments.

A. Implementation Details in Software Defined Networks

In this section we discuss the implementation details of single-stage Bloom filter based forwarding in OpenFlow. We note here, that to the best of our knowledge our implementation [8] is the only one which is fully supported by



(a) Contour plot of $k^*(\alpha, \beta)$. (b) Contour plot of $E[L]$.



(c) Contour plot of $\frac{k^*}{E[L]}$.

Fig. 7. The optimal number of hash functions.

TABLE III
FLOW ENTRIES IN THE i^{th} FLOW TABLE IN OUR OPENFLOW BLOOM FILTER BASED FORWARDING APPROACH.

match condition	associated actions
(in-packet filter AND Bloom ID _i) CMP Bloom ID _i	output(port _i) goto-table(i+1)
(always match)	goto-table(i+1)

standard OpenFlow v1.0 protocol⁵. Although the extension of this implementation for multi-stage filters can not be done in a standard OpenFlow v1.0 conform manner, the basics of Bloom filter based forwarding can be easily demonstrated through it. Thus, we shortly summarize this approach.

In the model of [8] Bloom IDs are assigned to switch ports. These Bloom IDs are used to generate the in-packet Bloom filter, i.e., port IDs, which the packet needs to pass through are bitwise OR-ed and put into the packet's destination Ethernet address field. In our implementation we need to slightly deviate from the regular usage of flow entries. Normally, flow entries correspond to a specific set of *flows*, whereas we proactively configure a separate flow table for each *port* as shown in Table III. The first entry in the table matches if the packet's Bloom filter contains the port's Bloom ID. In case of a match the switch forwards the packet via the corresponding port and independently of the outcome, the packet is matched

⁵Our implementation is open source and can be downloaded at <http://sb.tmit.bme.hu/mediawiki/index.php/Sigcomm2012>

against the next port's flow table hereby providing the stateless multicast switching capability. Hence, the packet is matched against the port ID of each outgoing interfaces, and forwarded on all necessary ports.

B. Implementation Issues in Intermediary Forwarding Elements

Next, we discuss the implementation issues of the proposed MSBF framework [17]. The forwarding process of the multi-stage filter runs in each relay node along the multicast tree and consists of the following steps.

- (1) The header of the incoming multicast packet is loaded into a fast access type of memory.
- (2) The starting number of 0's is counted, and the *length* field is loaded accordingly (Elias-gamma function).
- (3) The following *length* number of bits is identified as the current stage filter and prepared for Bloom filter membership testing.
- (4) The Bloom filter membership testing is executed in parallel at each link interface card.
- (5) If the test is positive then at the current bit position the header is truncated, that is the current stage filter with the length field is removed from the header.
- (6) If the header size is greater than zero the packet is forwarded using the new header.

The membership testing requires a bitwise logical AND operation of the Bloom filter and the link address. The result is then tested (bitwise CMP) if equals to the link address itself. Recall, that the filters at different stages can have different sizes, which requires link addresses of varying sizes. To store the addresses of all possible lengths for each link can be overwhelmingly memory consuming, prompt generation can provide a better solution instead. For this purpose [22] proposes a lightweight mechanism, where two uniformly distributed random hash functions (even two can be enough) are used to generate varying size hash codes to implement Bloom filters without any loss in the asymptotic false-positive probability. In this case two hash functions $h_1(x)$ and $h_2(x)$ are stored at line cards, where x is the link address. The i^{th} hash function $g_i(x)$ is generated by the formula $g_i(x) = h_1(x) + i \cdot h_2(x) \bmod b$, where b is the length of the hash code of the link to be established, and $i = 1, \dots, k$. Finally, it is tested whether the $g_i(x)^{th}$ bit in the stage filter is 1 for every i . Note that in such implementation membership testing depends mainly on the number of hash functions and less on the size of Bloom filters, allowing fast forwarding even for large Bloom filters.

With respect to the processing power required by the proposed method only $h_1(x)$ and $h_2(x)$ should be stored for each port, which can be stored in the local Level 1 (L1) cache of the CPU. L1 cache can be accessed in just a few CPU cycles and its typical size is tens of kilobytes. Recall that in the proposed method instead of address lookup we need to decode the length of the first stage, performing 2 – 3 modulo divisions with remainder using the two hash functions of $h_1(x)$ and $h_2(x)$, and test the related bit-positions in the header. These operations should not require more than a few tens of

CPU cycles⁶, which is not considered to be overwhelmingly expensive.

In comparison e.g. forwarding an IPv4 packet requires 6 – 8 memory access operations (and up to 16 for IPv6) taking up most of the forwarding time. Note that a 3 GHz processor has a CPU cycle of 0.3 nanoseconds, while a single DDR SDRAM memory access operation requires ~ 20 nanoseconds (that is about 66 CPU cycles) for each memory access, summing up to 396 – 528 CPU cycles for the whole IPv4 address lookup strongly depending on the applied advanced cache technology [24]. In traditional label switching technologies no extra processing is needed besides changing the label in the header; however, this comes for the price of keeping states in the switches (clearly not satisfying Goal 2).

C. Implementation Details in an ICN Prototype

We realize our solution in an available ICN prototype [7] for evaluation and demonstration. The *rendezvous* component implements the respective network function, as outlined in Section III-A. All publish/subscribe requests finally reach this element, which matches publishers with subscribers and triggers the formation of a forwarding path/tree. The *topology manager* manages the network topology and, upon request by a rendezvous component, creates forwarding paths from one or more publishers to one or more subscribers. These paths are sent to the respective publishers that use them when publishing information for a specific information identifier, utilizing the *forwarding* component for the delivery across the network.

We integrate our solution into the platform in [7] by adapting two modules [18]. First, we replace the forwarding component, which is currently based on [2], with a realization of the forwarding operation described in Section V-B. The platform design allows for forwarding functions to co-exist, delimited by the dissemination strategy for a particular part of the information structure. Second, we also adapt the topology manager by extending the current minimal spanning tree mechanism with the header encoding, i.e., the topology management function splits the tree into stages and calculates stage Bloom filters by executing the following steps (initializing the filter length to $b := b_{min}$):

- Step 1** Create the stage filter BF as the bitwise OR of the first b bits of the α in-tree links (if $b \in [b_{min}, b_{max}]$ calculated in Section IV-C).
- Step 2** Check whether BF contains any of the β out-tree link identifiers.
- Step 3** If BF is false-positive-free, return BF , else go to Step 1 with $b := b + 1$.

Note that the test interval $[b_{min}, b_{max}]$ can be efficiently computed based on the *Scalability* and *Insensitivity* properties (presented in Section IV-B and Section IV-C), that is given α

⁶Note that as $h_1(x), h_2(x) \leq b$ computing the remainder requires at most k subtractions and k comparisons in the worst case even without applying any software or hardware optimized algorithm. In general case a native and single (i.e., not considering pipeline execution) 32-bit unsigned integer division operation costs around 26-38 cycles in IA-32 or 64 bit Intel processor architectures [23].

and β , the expected filter length can be approximated by (15) and the required test range (given a prescribed success ratio $1 - \varepsilon$) around this mean value can be determined by (17).

Observe also that the three-step loop above succeeds to find the false-positive-free Bloom Filter in $[b_{min}, b_{max}]$ with probability $1 - \varepsilon$.

In the stage filter creation *BF* in **Step 1** a single integer number of hash functions is used (according to (19)) and its applicability is credited with the *Implementability* property discussed in Section IV-D.

The length of the FPF Bloom filter (b) is included into the stage filter using an Elias gamma encoding with $\gamma = 2 \cdot \lceil \log_2(b) \rceil + 1$ bits used for the length encoding per stage.

VI. SIMULATION AND MEASUREMENT RESULTS

A. Measures on Filter Length

In order to ensure the fair comparison of different Bloom filter based approaches, in this section we define performance measures to compare the header length of different forwarding schemes. Remember that the filter length m_l at link l in the proposed architecture stands for:

$$m_l = \gamma_l + b_l, \quad (20)$$

where γ_l is the total length of the Elias gamma codes of the stage filters, while b_l is the total bits consumed by (stage) Bloom filters in the header at link l .

Definition 1: The *filter compactness* of a multicast tree \mathbb{L} is denoted by $\eta(\mathbb{L})$, which is the sum of the header overhead along each link in the multicast tree divided by the square of the number of links $|\mathbb{L}|$ in the multicast tree, formally

$$\eta(\mathbb{L}) = \frac{\sum_{l \in \mathbb{L}} m_l}{|\mathbb{L}|^2} = \frac{m_{avg}}{|\mathbb{L}|}.$$

Obviously, a lower $\eta(\mathbb{L})$ refers to a more compact filter. To further explain, filter compactness refers to the average filter length divided by the number of tree links stored in the filter. Using this definition the performance of an architecture does not depend on the tree size, and the average filter length can be obtained by $\eta(\mathbb{L}) \cdot |\mathbb{L}|$.

To compare the overall performance of different filters, in the simulations a set of multicast tree is generated for a given network topology $G = (V, E)$, denoted by \mathcal{D} , and the *average filter compactness*

$$\eta_{avg} = \frac{\sum_{\mathbb{L} \in \mathcal{D}} \eta(\mathbb{L})}{|\mathcal{D}|}$$

of these trees is evaluated in the investigated architecture.

Recall that in our multicast architecture the headers are truncated at forwarding (i.e., the route information already traversed by the packet is erased), which leads to a better filter compactness η_{avg} . Therefore as a reference average filter compactness is also evaluated without abbreviating the headers at forwarding, denoted by μ_{avg} . In order to obtain a fair comparison of the traditional and our stage filters, similarly to the filter compactness we define the following metric.

TABLE IV
IN-TREE AND OUT-TREE LINKS PER STAGE IN THE COST 266
PAN-EUROPEAN BACKBONE NETWORK WITH 37 NODES AND 57 LINKS
($\mathcal{D} = 2000$).

	Hop-distance (stage h) from source							ISBF Total
	1	2	3	4	5	6	7	
α	1.85	2.66	2.67	2.90	2.28	1.87	1.32	15.55
β	1.08	3.90	7.01	6.86	8.26	6.17	6.24	39.52

Definition 2: The *Bloom filter compactness* of a multicast tree \mathbb{L} is denoted by $\lambda(\mathbb{L})$, which is the sum of the lengths of the in-packet Bloom-filters along each link in the multicast tree divided by the square of the number of links $|\mathbb{L}|$ in the multicast tree, formally

$$\lambda(\mathbb{L}) = \frac{\sum_{l \in \mathbb{L}} b_l}{|\mathbb{L}|^2} = \frac{b_{avg}}{|\mathbb{L}|}.$$

Similarly, *average Bloom filter compactness* without abbreviating the headers is defined as λ_{avg} for a set of \mathcal{D} multicast trees.

B. False-Positive-Free Bloom Filter Length Analysis

In the simulation we compare the two Bloom filter based forwarding approaches, namely the traditional in-packet Bloom filters [2] which was modified to handle varying size headers (drawn with filled marks on the charts), and the proposed MSBF approach (drawn with empty marks on the charts). We compare their performance in terms of the metrics proposed in Section VI-A, i.e., η_{avg} and μ_{avg} to investigate the effect of abbreviating the header, and μ_{avg} and λ_{avg} to investigate the effect of varying size headers.

Fig. 8 shows the result on the COST 266 pan-European backbone network with 37 nodes and 57 links. The demands were classified according to the maximum hop distance in the multicast tree. $\mathcal{D} = 2000$ random demands were generated with unicast traffic only on Fig. 8(a), and multicast traffic with up to 10 terminal nodes on Fig. 8(b). Coinciding with our analytical evaluation the multi-stage Bloom filter has significantly shorter filter sizes. Surprisingly this is true even when the multi-stage Bloom filter consists of one Bloom filter in one hop trees. This is because in multi-stage Bloom filters the number of links on which false-positive can occur is smaller, since none of the links, which are two hops away from the source node can generate a false-positive, due to the header truncation mechanism, i.e., zero sized header prevents the forwarding of a packet in the multi-stage case. Note that the size of the multicast trees increases as the number of hops increases and shows the great scalability multi-stage Bloom filter can achieve. This is validated by the number of in-tree and out-tree links per stage as well in Table IV.

In the case of traditional in-packet Bloom filters encoding the size adds small overhead, especially for large multicast trees. Since multi-stage Bloom filters consists of several consecutive Bloom filters, one may argue that encoding these boundaries may end up in a larger overhead. Our results support that this is not the case, since due to the truncation

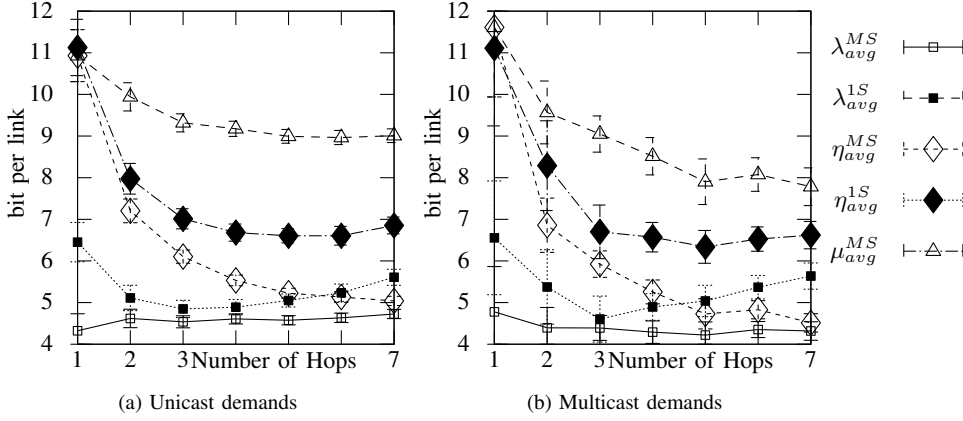


Fig. 8. The header length versus the number of hops in the multicast tree for the 37-node European reference network (legends are shown in Figure 9).

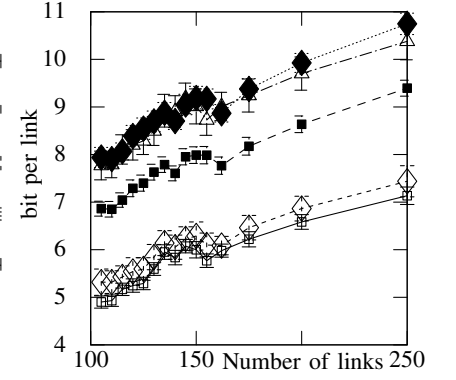


Fig. 9. Results of 2-connected 50-node networks with different number of links.

TABLE V

TOPOLOGIES USED FOR THE NUMERICAL EVALUATIONS WITH FILTER LENGTH MEASURES FOR THE TRADITIONAL IN-PACKET BLOOM FILTERS (1SBF) AND FOR THE MULTI-STAGE (MSBF) USING THE EXACT AND APPROXIMATION FORMULAE.

Topology	Nodes ($ V $)	Edges ($ E $)	Xcast IPv4	MSBF	η_{avg} Approx	1SBF	μ_{avg} MSBF	λ_{avg}		δ	
								MSBF	1SBF	Incremental	$E[L] \pm \delta$
Cost266 [25]	37	57	18.77	4.33	4.33	10.97	6.79	3.90	9.90	52.38	34.28
Germany [25]	50	88	17.88	5.97	5.97	11.79	8.14	5.13	10.42	66.69	40.15
Deltacom [26]	113	161	9.89	4.18	4.07	12.11	6.87	3.92	11.27	105.38	70.53
Random E-R	1001	1997	150.15	5.15	4.97	5.77	7.41	3.15	3.73	13.38	21.21
AS level	34306	71448	10.23	10.88	10.93	19.75	16.39	13.53	18.78	366.27	151.84

of the header at forwarding and the space efficiency of the multi-stage filter, in average header length the multi-stage Bloom filters remarkably outperform the original approach. The advantage of the MSBF approach is even more significant for unicast demands.

Next we investigate the performance respect the network density. 20 random 50-node, two-connected networks are generated with different network density. Fig. 9 shows the results of $\mathcal{D} = 500$ demands where the horizontal axis corresponds to the number of links in the graph. One can observe that the performance gain of multi-stage Bloom filter is always 3 – 4 bit per link and does not depend on the density of the topology.

Finally, we investigate the scalability of the approaches as the network grows. Table V shows the results on topologies with different sizes and with multicast demands of terminal nodes at most 10. As a reference Xcast based solution is also added to the table, in which the header consists of a series of IPv4 addresses with 32 bit for each destination. However, note that IPv4 and Xcast are not source routing solutions and require large routing tables at the routers (i.e., clearly not supporting Goal 1 and Goal 2). We were surprised to see that, although the forwarding decision for an in-packet Bloom filter is significantly simpler and faster compared to traditional IPv4 forwarding, using in-packet Bloom filters even at the AS-level has similar performance than Xcast. Further note that using the approximation formula developed in Section IV-C for the test range δ requires a lower number of tested filter lengths in order to find an FPF Bloom filter, which is important

from a practical point of view. Furthermore, these filters are comparable to those in η_{avg} which were generated with the greedy approach.

C. Forwarding Complexity in the ICN Prototype

We compare the complexity of our scalable multicast architecture in each forwarding element with LIPSIN [2], based on the MSBF implementation within our ICN prototype [18]. Referring to Section V-B, the MSBF forwarding complexity not only includes the Bloom filter based membership test but also the on-demand hash creation as well as the extraction of the stage FPF Bloom filter. On the other hand, traditional Bloom filter approach (LIPSIN) only performs a constant-length Bloom filter membership test. For comparison, we created three different multicast trees on the same topology with 3, 4, and 5 stages, respectively. We repeatedly executed the forwarding function in our prototype (processing the LIPSIN header at each stage as well), and determined the average of the execution times at the various stage boundaries. Table VI shows the results of these tests, compared to the constant filter length and forwarding time of the LIPSIN. One can observe that the execution times for stage Bloom filters with length smaller than 50 are faster than comparable LIPSIN times. This is due to the membership test operating on smaller bit sets compared to the 256 bits used in the LIPSIN alternative. Again, these results can only be indicative since our current implementation is not optimized through, e.g. replacing the on-demand hash function through table lookups or utilizing hardware assistance could be further improve the performance

TABLE VI
HEADER LENGTH AND FORWARDING EXECUTION TIMES.

Stage (h)	Header Length at Forwarding				Forwarding time	
	Elias (γ)	Filter (b)	MSBF ($m = \gamma + b$)	LIPSIN ($m = b$)	MSBF (μ s)	LIPSIN (μ s)
1	5	7	12	256	8.0	16.1
2	15	9	24	256	11.8	16.1
3	18	11	29	256	12.4	16.1
Total	38	27	65	256	32.3	48.3
1	2	3	5	256	4.9	16.1
2	21	11	32	256	12.9	16.1
3	31	11	42	256	14.2	16.1
4	18	11	29	256	12.4	16.1
Total	72	36	108	256	44.4	64.4
1	2	3	5	256	4.9	16.1
2	17	11	28	256	12.2	16.1
3	56	13	69	256	16.3	16.1
4	59	13	72	256	16.5	16.1
5	37	13	50	256	14.9	16.1
Total	171	53	224	256	64.8	80.5

of our MSBF approach based on varying length FPF Bloom filters.

D. Discussion on Forwarding Loops and Packet Storms

A well-known problem of Bloom filters is that through a chain of false-positives a packet can loop back to a previously visited node where generates a false-positive again and stuck in an infinite loop. In extreme situations such a behavior may cause even packets storms. In [4] a bit permutation technique was proposed to prevent such anomalies with very high probability. Our proposed multi-stage Bloom filters remedy these illnesses in a fairly natural way (even without using FPF filters), since due to the truncation of the filter after every stage, the packets cannot go further than a few hops. In such a way our MSBF approach certainly prevents the formation of infinite loops by encoding the stage decomposition into the header. Note that the flow duplication is indirectly prevented by limiting the multi-cast routing on tree without false positive forwarding.

VII. CONCLUSIONS

In this paper we addressed the scalable multicast forwarding problem, and introduced a novel multi-stage Bloom filter based architecture, which tackles Goals 1-3. Although in ICN architectures and with the application of in-packet Bloom filters Goals 1 and Goal 2 have been addressed, the scalability in terms of supported multicast tree size was lost in these concepts owing to neglecting topology information when creating the in-packet filter. Furthermore, false-positive forwarding was a serious issue in these environments. We gave a thorough analysis on the design of minimal length false-positive-free Bloom filter both for the traditional Bloom filter approach with varying size filters, and also for our novel multi-stage architecture in order to avoid such anomalies. Approximation formulae were proposed, which can be directly used in a practical implementation of e.g. an ICN prototype. Our simulation results suggest that encoding topology related information into varying length multi-stage filters results in

much better efficiency in several topologies than its previous counterparts.

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APPENDIX

A. Proof of Lemma 1

Proof: First we rewrite and approximate $(1 - c^{\frac{i}{\alpha}})^\beta$ as

$$(1 - c^{\frac{i}{\alpha}})^\beta = \text{Exp} \left(\beta \ln(1 - c^{\frac{i}{\alpha}}) \right) \approx \text{Exp} \left(-\beta c^{\frac{i}{\alpha}} \right) \quad (21)$$

due to $\ln(1 - x) \approx -x$ for small x . Based on this, consider the following approximation of the logarithm of $P(L > b)$:

$$\begin{aligned} \ln P(L > b) &= \sum_{i=0}^b \ln(1 - (1 - c^{\frac{i}{\alpha}})^\beta) \approx \\ &\approx \sum_{i=0}^b \ln(1 - \text{Exp}(-\beta c^{\frac{i}{\alpha}})) \approx \sum_{i=0}^b \left(-\text{Exp}(-\beta c^{\frac{i}{\alpha}}) \right), \quad (22) \end{aligned}$$

where the last approximation again due to $\ln(1 - x) \approx -x$. This sum above can also be viewed as an integrate approximation sum, hence

$$\sum_{i=0}^b \left(-\text{Exp}(-\beta c^{\frac{i}{\alpha}}) \right) \approx \int_{x=0}^b \left(-\text{Exp}(-\beta c^{\frac{x}{\alpha}}) \right) dx \quad (23)$$

The integrate on the right hand side can be expressed by the exponential integral $\text{Ei}(z)$ function as

$$\frac{\alpha \text{Ei}(-c^{1/\alpha} \beta)}{\ln c} - \frac{\alpha \text{Ei}(-c^{b/\alpha} \beta)}{\ln c} \approx -\frac{\alpha \text{Ei}(-c^{b/\alpha} \beta)}{\ln c}, \quad (24)$$

for larger b , from which the statement follows. ■