Subgraph nomination: query by example subgraph retrieval in networks

Al-Fahad Al-Qadhi · Carey E. Priebe · Hayden S. Helm · Vince Lyzinski

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Abstract
This paper introduces the subgraph nomination inference task, in which example subgraphs of interest are used to query a network for similarly interesting subgraphs. This type of problem appears time and again in real world problems connected to, for example, user recommendation systems and structural retrieval tasks in social and biological/connectomic networks. We formally define the subgraph nomination framework with an emphasis on the notion of a user-in-the-loop in the subgraph nomination pipeline. In this setting, a user can provide additional post-nomination light supervision that can be incorporated into the retrieval task. After introducing and formalizing the retrieval task, we examine the nuanced effect that user-supervision can have on performance, both analytically and across real and simulated data examples.

Keywords Statistical machine learning · Information retrieval · Social and information networks

1 Introduction

Stated succinctly, the subgraph nomination problem is as follows: given a subgraph or subgraphs of interest in a network $G_1$, we seek to find heretofore unknown subgraphs of interest in $G_1$ or in a second network $G_2$. The subgraph nomination problem can be viewed as an amalgam of the problems of noisy subgraph detection (Jin et al. 2019; Caelli and Kosinov 2004; Lladós et al. 2001; Jacob et al. 2018; Sussman et al. 2019) and vertex nomination (Marchette et al. 2011; Coppersmith 2014; Fishkind et al. 2015; Lyzinski et al. 2019), in which our task is to produce a rank-list of candidate subgraphs, with (ideally) unknown subgraphs of interest concentrating at the top of the rank list. While seemingly simple on its surface, subgraph nomination necessarily entails the often non-trivial combination of subgraph detection (i.e., we have to find candidate subgraphs to rank) and multiple graph comparison methodologies (i.e., we have to rank the candidate subgraphs). We note here that the subgraph nomination problem considered herein is closely related to many existing graph query problem formulations (see, for example, the work on query by example in Mottin et al. (2016) and Lisandrini et al. (2020)). Our novel contribution is the novel statistical framework underlying our subgraph nomination formulation, which allows us to frame and rigorously study the problem (and the effect of a user-in-the-loop) in a number of popular statistical network models, with the ultimate goal of establishing results of statistical consistency for subgraph nomination akin to the analogues in the vertex nomination literature (Lyzinski et al. 2016, 2019).

Subgraph detection—here, producing candidate subgraphs of interest from $G_1$ or $G_2$—is an active area of research in machine learning and pattern recognition, and encompasses both the famous subgraph isomorphism problem (see, for example, Alon et al. (1995), Slota and Madduri (2013), Cordella et al. (2004), Ullmann (1976)) as well as numerous noisy subgraph detection routines. In subgraph nomination, subgraph detection is further complicated by the generality of
the (topological) features that define the training subgraphs as “interesting.” In particular, it may be the case that the query set encompasses multiple different interesting templates (Jacob et al. 2018; Jin et al. 2019) or motifs-of-interest (Milo et al. 2002; Lyzinski et al. 2017), each delineating an activity or structure of interest in the network. Moreover, the templates of interesting subgraphs need not be manifestly similar to each other. The notion of similarity across subgraphs will be formalized via a subgraph dissimilarity mapping $\Delta$, and a subgraph nomination scheme can (loosely) be considered as an estimate of the subgraph structure of $g_2$ combined with a dissimilarity $\Delta$ used to rank the subgraphs (see Def. 5). If $\mathcal{S}_G$ represents the collection of all subgraphs of a network $G$, then in the single-graph setting

$$\Delta : \mathcal{S}_{G_1} \times \mathcal{S}_{G_1} \rightarrow [0, 1],$$

and in the pair of graphs setting

$$\Delta : \mathcal{S}_{G_1} \times \mathcal{S}_{G_2} \rightarrow [0, 1],$$

with smaller values indicating more similar subgraphs. We comment here that although the two graph setting will be our focus moving forward, this naturally lifts to the single graph setting by considering the two graphs to nominate across as parts of a partition of a larger network.

Large values of $\Delta$ indicate that the subgraphs are highly dissimilar according to $\Delta$, and candidate subgraphs are defined as interesting if they are sufficiently similar (i.e., insufficiently dissimilar) to any subgraph in the training sample. Choosing an appropriate $\Delta$ from which to construct the subgraph rankings is of primary importance, and adaptive methods (similar to those in the learning-to-rank problem in the information retrieval literature (Liu 2011; Helm et al. 2020)) can be considered to learn $\Delta$ from the training subgraphs of interest. While estimating an optimal $\Delta$ is central in this problem framework, we do not derive formal procedures for estimating $\Delta$ in this paper. Instead, we choose to focus on the effect of user-in-the-loop supervision across a variety of possible $\Delta$ choices, where the user is modeled as additional light supervision that can be used to refine the output of a subgraph nomination routine.

As a motivating example for our focus on studying the effects of a user-in-the-loop, we consider the problem of detecting and nominating particular brain regions of interest across the hemispheres of a human connectome from the BNU1 dataset (Zuo et al. 2014) downloaded from https://neurodata.io/mri/, and finding subreddits across different time periods in the Reddit social network data collected in Baumgartner et al. (2020) and downloaded from https://files.pushshift.io/reddit/comments/.

In the setting of BNU1, subgraph nomination can be used as a tool to better understand (and detect) the structural similarity between brain regions across hemispheres. To this end, we consider a region (or regions) of interest in the left hemisphere, and there might be multiple subgraphs in the right hemisphere that match (according to $\Delta$) well to the training data. Moreover, the most similar region of interest in the right hemisphere (i.e., the $\Delta$-optimal subgraph of the right hemisphere) may vary dramatically depending on the $\Delta$ used (see, for example, Fig. 7 in Sussman et al. (2019)), and the optimal region may not significantly overlap with a true latent region of interest in the left hemisphere. In this case, even a sensible ranking scheme would be potentially unable to correctly retrieve the desired region of interest in the right hemisphere (even if the proper subgraph was identified). In this case, we can naturally employ the help of a user-in-the-loop (Amershi et al. 2014), who, given a vertex, can decide whether it is interesting/part of an interesting subgraph to help refine our ranking.

Meanwhile, in the setting of the Reddit social network, subgraph nomination can be used to track subreddits across time based on their structure. Therefore, we consider a subreddit of interest in a week (based on posting time) and attempt to find it among unlabeled subreddits in the next week. For the $\Delta$, we use the Graph Matching Network (GMN) from Yujia et al. (2019). Here, the graph matching network training starts quick, but once a certain level of accuracy is passed the training slows significantly and oscillates. We investigate adding the user in the loop to improve the accuracy with less training. Accuracy in this setting represents the chance of finding the subreddit among the top $l$ nominated graphs. As can be seen in 11, the user in the loop can achieve better accuracy with less training and at a lower level, even when the user is relatively faulty. For large networks, such as Reddit this could save computational resources, as well as human resources by reducing the amount of subreddits an expert needs to investigate.

The subgraph nomination framework we present in this paper is both general enough to include the operationally significant effect of a user-in-the-loop and principled enough to theoretically analyze nomination schemes. After discussing preliminaries and presenting the problem framework, we demonstrate multiple pathologies to highlight the power of the subsequent theory as a predictive tool of the efficacy of the user in general situations. In particular, we show that a user can improve the performance over a scheme that achieves the Bayes optimal rate sans user-in-the-loop, even in the presence of potential user-error. This result is similar to results in the classification literature when given noisy training labels (Frénay and Verleysen 2013; Natarajan et al. 2013). We further show that, given noisily recovered subgraphs, including information from an oracle user can deteriorate performance due to the noise within the subgraph detection routines.
Notation: For a positive integer $n$, we will denote $[n] := \{1, 2, 3, \ldots, n\}$; $J_n$ to the $n \times n$ matrix of all 1's; $G_n$ to be the set of labeled $n$-vertex graphs; for $G \in G_n$, and $W \subseteq V(G)$, we let $G[W]$ denote the induced subgraph of $G$ on $W$.

2 Introduction and background

In subgraph nomination, our task is as follows: given a subgraph or subgraphs of interest in a network $G_1$, we seek to find heretofore unknown subgraphs of interest in a second network $G_2$. Our approach to subgraph nomination proceeds by

1. Partitioning the vertices in $G_2$ into candidate subgraphs (note that the choice of non-overlapping candidate subgraphs is merely a theoretical/notational convenience, and in practice, the subgraphs to be ranked can have overlap);
2. Ordering the candidate subgraphs in $G_2$ into a rank list with the unknown subgraphs of interest ideally concentrating at the top of the rank list.

Subgraph nomination is a natural extension of vertex nomination (VN) (Marchette et al. 2011; Coppersmith 2014; Suwan et al. 2015; Fishkind et al. 2015; Lyzinski et al. 2019; Agterberg et al. 2020), and we will begin by providing a brief background on recent developments in VN, as this will provide useful context for our later novel formulation of subgraph nomination.

2.1 Vertex nomination

Semi-supervised querying of large graphs and databases is a common graph inference task. For example, in a graph database with multiple features, one may be given a collection of book names all with the common latent feature “Horror Fiction Best Sellers,” and the goal would be to query the database for more titles that fit this description; see, for example, the work in Renzo et al. (2016) and Rastogi et al. (2017). Learning this unifying feature from the query set is a challenging problem unto itself (Rastogi et al. 2019), especially in settings where the features that define interestingness are nuanced or multi-faceted.

While we can interpret such a problem as a vertex classification problem where vertices are either labeled interesting or not, in the presence of very large networks with large class imbalance between interesting and non-interesting vertices, there is often limited training data and limited user resources for verifying returned results. In this setting, an information retrieval (IR) framework may be more appropriate, wherein unlabeled vertices would be ordered in a rank list based on how interesting they are deemed to be. This inference task is synonymous with vertex nomination (or personal recommender systems on graphs).

In Patsolic et al. (2020), Lyzinski et al. (2019), Agterberg et al. (2020) and Levin et al. (2020), the vertex nomination problem is defined as follows. Given vertices of interest $V^* \subseteq V(G_1)$ in a network $G_1$, and corresponding unknown vertices of interest $U^* \subseteq V(G_2)$ in a second network $G_2$ (whose identities are hidden to the user), use $G_1, G_2, V^*$ to rank the vertices in $G_2$ into a nomination list, with vertices in $U^*$ concentrating at the top of the nomination list. In its initial formulation (Coppersmith 2014; Marchette et al. 2011; Fishkind et al. 2015; Yoder et al. 2020), the feature that defined vertices as interesting was membership in a community of interest, and the community memberships of vertices in $V^*$ were used to nominate the vertices in $G_1$ (no second network was introduced, or $G_2 = G_1 \setminus V^*$) with unknown community memberships. Subsequent work (Rastogi et al. 2017, 2019; Patsolic et al. 2020; Lyzinski et al. 2019) sought to generalize the features that defined vertices as interesting beyond simple community membership, and lifted the vertex nomination problem to the two graph setting.

In order to allow for a broadly general class of networks to be considered, the general vertex nomination problem framework of Agterberg et al. (2020) and Lyzinski et al. (2019) referenced above is situated in the context of nominatable distributions, a broad class of random graph distributions defined in Agterberg et al. (2020). Within this broad class of models, the concepts of Bayes optimality and consistency were developed in Lyzinski et al. (2019) and Agterberg et al. (2020) and the important result that universally consistent VN schemes do not exist is proven in Lyzinski et al. (2019). These results were leveraged in Agterberg et al. (2020) to develop an probabilistic adversarial contamination model in the context of VN as well as regularization schemes to counter the adversary. The results of Lyzinski et al. (2019) and Agterberg et al. (2020) were further extended to the richly featured network setting in Levin et al. (2020), in which the (potentially) complementary roles of features and network structure are explored in the VN task.

Beyond the development of novel VN algorithms (Yoder et al. 2020), one of the key aspects of the recent theoretical developments in VN is the notion that vertex labels in $g_2$ are uninformative in the ranking scheme, which is sensible if we are mirroring setting in which the vertex labels do not aid in the delineation between interesting and uninteresting vertices. In subgraph nomination, the uninformative nature of the labels can be accounted for in the dissimilarity $\Delta$ (see Eq. 1). Note that while similar consistency results to those in VN can be derived in the setting of subgraph nomination, we do not focus on that here. Rather we will focus on the role of users-in-the-loop in the subgraph nomination framework.
Remark 1 We note that it is possible to view the subgraph nomination methods as naturally being built upon existing VN methods via uncovering vertices of interest (from VN) and building out from them to uncover subgraph structure of interest. One immediate hurdle to this approach would be if the subgraph structure is the interesting trait rather than the individual vertex identities, as then the nominated vertices in VN would need to look at local structure to be correctly retrieved, and essentially subgraphs would end up being the nominated structures. It is also natural to use subgraph nomination for VN, by nominating structures first and then vertices within the structures.

2.2 Hierarchical subgraph models

We will use \( G_n \) to denote the set of \( n \)-vertex labeled graphs on a common vertex set \( V = V_n \). Given the (informal) explanation of the motivating task in subgraph nomination—given light supervision, partition a graph into subgraphs and rank the subgraphs based on how interesting they are judged to be—hierarchical network models are a natural setting for initially formalizing the subgraph nomination inference task.

We note here that the subgraph nomination inference task can be formulated in the context of any graph division mechanism that yields partition subgraph structure (e.g., graph cuts (Kolmogorov and Rother 2007; Delling et al. 2011), graph clustering methods (Newman 2006; Blondel et al. 2008; Traag et al. 2019), etc.); the Hierarchical blockmodel considered below is one such setting, and it also provides a convenient model setting in which to formulate subgraph nomination formally.

Hierarchical models have seen a surge in popularity in the network literature (see, for example, Sales-Pardo et al. (2007), Clauset et al. (2008), Park et al. (2010), Peixoto (2014), Lyzinski et al. (2017), Tianxi et al. (2018)) and naturally allow for multiple layers of structure to be simultaneously modeled in a network, allowing us to imbue a graph \( G \) with subgraph structures of a desired form or motif-type. We begin with our definition of a hierarchical network as a network together with a Network Hierarchical Function.

Definition 1 Let \( g = (V, E) \in G_n \), and for each \( i \in \mathbb{Z} \geq 0 \) let \( [i] = \{1, 2, \ldots, i\} \). Let \( k \leq n \). A function

\[
H = H_k : V \times [k] \rightarrow [n]
\]

is a \( k \)-level Network Hierarchical Function of \([n]\) if

(i) For each \( i \in [k] \), \( H(\cdot, i) \) represents a partition of the vertices of \( V \) into \( n_i := n(\mathcal{H}) \) nonempty parts (so that \( \max_n H(v, i) = n_i \)). These \( n_i \) satisfy \( 1 \leq n_1 \leq n_2 \leq \cdots \leq n_k \leq n \). The signature of \( H \) is defined to be the vector \( n = (n_1, n_2, \ldots, n_k) \). Note that by considering appending 0’s onto the end of \( n \) as needed to make it length \( n \), we can consider the signature of an \( n \) vertex hierarchical graph to be a length \( n \) vector.

(ii) \( H(v_1, j) = H(v_2, j) \) only if \( H(v_1, i) = H(v_2, i) \) for all \( i \leq j \); i.e., the partition is nested.

For an example of network hierarchical functions, see Fig. 1. A graph \( g \in G_n \) together with a Network Hierarchical Function \( H_k \) of \([n]\) is a \( k \)-level Hierarchical Graph. For \( k \leq n \), we denote

\[
\mathcal{H}_{k,n} = \{H_k \mid H_k \text{ is a } k \text{-level Network Hierarchical Function of } [n]\},
\]

and we define

\[
\mathcal{H}G_n := \{(g, H) \mid g \in G_n, \text{ and } H \in \cup_{k=1}^n \mathcal{H}_{k,n}\}
\]

to be the set of all Hierarchical graphs of order \( n \).

Letting \( k \leq n \), let \( H \in \mathcal{H}_{k,n} \). For each \( i \in [k] \) and \( j \in [n_i] \), we define the sets

\[
B_j^i = \{v \in V(g) \mid H(v, i) = j\}
\]

to be the set of vertices in the \( j \)-th part of the \( i \)-th level of the hierarchy;

\[
B_{(j)}^{-1} = \{v \in V(G) \text{ s.t. for all } v' \in B_j^i, H(v, i - 1) = H(v', i - 1)\}
\]

to be the one-step upward merge of \( B_j^i \) in the hierarchy;

\[
B_i^j = \{B_j^i \mid 1 < j < n_i\}
\]

to be the set of all parts at level \( i \) in the hierarchy; and

\[
B = \{B_i^j \mid i \in [k]\}
\]

to be the set of blocks.

As an example, consider the hierarchical network in Fig. 1, which shows a 3-level hierarchical network, where we have (for example):

\[
n_1 = 1; n_2 = 3; n_3 = 9;
\]

\[
B_1^1 = V = B_2^1 \cup B_2^2 \cup B_2^3;
\]

\[
B_2^1 = B_3^1 \cup B_3^2 \cup B_3^3;
\]

\[
B_3^1 = \{B_4^3, B_4^4, B_4^5, B_5^3, B_5^4, B_5^5, B_6^3, B_6^4, B_6^5, B_7^3, B_7^4, B_7^5\};
\]

\[
B = \{B_1^1, B_1^2, B_1^3, B_2^1, B_2^2, B_2^3, B_2^4, B_2^5, B_3^1, B_3^2, B_3^3, B_3^4, B_3^5\}.
\]
2.2.1 Hierarchical stochastic block models

One important example of a network distribution with hierarchical structure is the hierarchical stochastic blockmodel (HSBM) (Peixoto 2014; Lyzinski et al. 2017). Before defining the HSBM, we first define the standard stochastic blockmodel (Holland et al. 1983).

**Definition 2** We say that an $n$-vertex random graph $G$ is an instantiation of a stochastic blockmodel with parameters $(n, K, \Lambda, \pi)$ (abbreviated $A \sim \text{SBM}(n, K, \Lambda, \pi)$) if

(i) The block membership vector $\pi \in \mathbb{R}^K$ satisfies $\pi(i) \geq 0$ for all $i \in [K]$, and $\sum_i \pi(i) = 1$;
(ii) The vertex set $V = V(G)$ is the disjoint union of $K$ blocks $V = B_1 \sqcup B_2 \sqcup \cdots \sqcup B_K$, where each vertex $v \in V$ is independently assigned to a block according to a Multinomial$(1, \pi)$ distribution. For each vertex $v \in V(G)$, let $b_v$ be the block that $v$ is assigned to.
(iii) The block probability matrix $\Lambda \in [0, 1]^{K \times K}$ is a symmetric matrix. Conditional on the block assignment vector $b = (b_v)$, for each pair of vertices $\{u, v\} \in (V)^2$, $\{1|u \sim_G v\} \overset{\text{ind.}}{\sim} \text{Bernoulli}(\Lambda_{b_u, b_v})$.

As in Peixoto (2014), we will define the HSBM recursively from the top down. In essence, the 2-level HSBM is an SBM where each block itself has an SBM structure. If further, every one of the blocks of the second level is an SBM then we have a three level HSBM, and so on. Formally, we have the following recursive definition.

**Definition 3** We say that an $n$-vertex random graph $G$ is an instantiation of a 2-level hierarchical stochastic blockmodel with parameters

$$(n, K_1, \Lambda_1, \pi_1, [K_2^{(j)}, \Lambda_2^{(j)}, \pi_2^{(j)}]_{j=1}^{K_1})$$

if

(i) The block membership vector $\pi_1 \in \mathbb{R}^{K_1}$ satisfies $\pi_1(i) \geq 0$ for all $i \in [K_1]$, and $\sum_i \pi_1(i) = 1$;
(ii) The vertex set $V = V(G)$ is the disjoint union of $K_1$ blocks $V = B_1^1 \sqcup B_2^1 \sqcup \cdots \sqcup B_{K_1}^1$, where each vertex $v \in V$ is independently assigned to a block according to a Multinomial$(1, \pi_1)$ distribution. For each vertex $v \in V(G)$, let $b_v$ be the block that $v$ is assigned to.
(iii) The block probability matrix $\Lambda_1 \in [0, 1]^{K_1 \times K_1}$ is a hollow symmetric matrix. Conditional on the block assignment vector $b^{(1)} = (b_v^{(1)})$, for each pair of vertices $\{u, v\} \in (V)^2$, $\{1|u \sim_G v\} \overset{\text{ind.}}{\sim} \text{Bernoulli}(\Lambda_1_{b_u^{(1)}, b_v^{(1)}})$.
(iv) For each $j \in [K_1]$, conditional on the block assignment vector $b^{(1)} = (b_v^{(1)})$ and $|B_j^{(1)}| > 0$, we have that $G[B_j^{(1)}] \sim \text{SBM}([B_j^{(1)}], K_2^{(j)}, \Lambda_2^{(j)}, \pi_2^{(j)})$.

Moreover, conditional on the block assignment vector $b^{(1)} = (b_v^{(1)})$ the collection $\{G[B_j^{(1)}]\}_{j=1}^{K_1}$ are mutually independent.

Formally defining the HSBM beyond the second level of the hierarchy is notationally complex, but the main idea is that a $k$-level HSBM has the same set-up as the 2-level HSBM except that at part iv of the definition, we have

(iv') For each $j \in [K_1]$, conditional on the block assignment vector $b^{(1)} = (b_v^{(1)})$ and $|B_j^{(1)}| > 0$, we have that $G[B_j^{(1)}] \sim k-1$ level HSBM.

Moreover, conditional on the block assignment vector $b^{(1)} = (b_v^{(1)})$, we have that the collection $\{G[B_j^{(1)}]\}_{j=1}^{K_1}$ are mutually independent.
Recursively applying \( iv' \) together with the definition of a 2-level HSBM allows us to define HSBMs of arbitrary depth (\( \leq n \) of course).

**Remark 2** For a \( k \)-level HSBM, we shall adopt the notation from the definition of hierarchical graphs already established. Namely, we will denote the \( j \)-th block at level \( i \) via \( B^i_j \), where the blocks are labeled in order beginning with those of \( B^{(1)}_{(1)} \) and ending with those of \( B^{(i-1)}_{(i-1)} \). We will denote the block-membership function at level \( i \) via \( b^i \).

While the model complexity of the HSBM can grow quite rapidly as we allow for more structure in each block of the higher level SBMs, the top-down structure does offer complexity savings versus an SBM with the same number of blocks at the bottom-level of the hierarchy. For example, the 1-level HSBM (i.e., the SBM) with \( K \) blocks requires

\[
\binom{K}{2} + K + K - 1 = O(K^2)
\]

parameters to define, while the 2-level HSBM requires

\[
\binom{K_1}{2} + K_1 + \binom{K_2(j)}{2} + K_2 + K_2 - 1 = O\left(K_1^2 + \sum_j (K_2(j))^2\right)
\]

parameters. As a simple example, if \( K_1 = 3 \) and each \( K_2(j) = 3 \), then the HSBM has 9 blocks at its bottom level and requires 29 parameters while a 9 block SBM requires 53 parameters. The complexity of an HSBM can be further reduced by enforcing repeated motif structure (Lyzinski et al. 2017), though we do not explore this further herein.

**Remark 3** Consider a \( k \)-level HSBM \( G \). Letting \((n_i)_{i=1}^k\) denote the number of blocks at level \( i \) in the hierarchy of \( G \), so that (for example)

\[
n_1 = K_1 \quad n_2 = \sum_{i=1}^{K_1} K_2(i) \quad n_3 = \sum_{i=1}^{K_1} \sum_{j=1}^{K_2(i)} K_3(i,j)
\]

\( n_4 = \sum_{i=1}^{K_1} \sum_{j=1}^{K_2(i)} \sum_{\ell=1}^{K_3(i,j)} K_4(\ell,j,i) \)

(\( K_3(i,j) \) is the number of blocks in the \( j \)-th block of the \( i \)-th block of \( G \); \( K_4(\ell,j,i) \) is the number of blocks in the \( \ell \)-th block of the \( j \)-th block of the \( i \)-th block of \( G \); etc...) and conditioning on all blocks being non-empty at each level of the hierarchy, \( G \) defines a distribution over the subset of \( k \)-level hierarchical graphs

\[
\mathcal{H}_n^m := \{(g, H) \in \mathcal{H}_n \mid \text{the signature of } H \text{ is } n\}.
\]

We then have that (for example), \( H(v, i) = b^i_v \) for all \( v \in [n] \) and \( i \in [k] \).

### 2.2.2 Hierarchical subgraph nomination

A key component to the hierarchical subgraph nomination (HSN) inference task is the notion of subgraphs of interest within and across the network pair. This is easily accomplished in non-random settings, where a fixed set of indices can be used to define the subgraphs of interest within and across networks. In order to expedite this in random networks, for a fixed pair of signatures \( n \) and \( m \), we will consider distributions restricted to \( \mathcal{H}_n^m \times \mathcal{H}_m^m \); this will allow us to define a consistent set of indices for subgraphs of interest across the random network pair.

In hierarchical subgraph nomination, we consider a pair of hierarchical graphs \((g_1, H_1) \in \mathcal{H}_n^m \) and \((g_2, H_2) \in \mathcal{H}_m^m \) where \( H_1 \) and \( H_2 \) are unobserved and only the graphs \( g_1 \) and \( g_2 \) are observed. We are additionally given a set of “training” subgraphs of interest in \( g_1 \), denoted

\[
T_1 = \{B^s_{i_1}\}_{i_1 \in \{s_1, s_2\}} \in S_1
\]

where \( S_1 \) is a set of ordered pair indices \( S_1 = \{s = (s_1, s_2)\} \) of the given subgraphs of interest, with \( s_1 \) denoting the level and \( s_2 \) the index of the subgraph in \( H_1 \). Note that the second index in the subscript of \( B^s_{i_1} \) is used to denote the subgraph in \( g_1 \), whereas \( B^s_{i_2} \) will be used to indicate subgraphs in the hierarchy of \( g_2 \). Note that the possible indices of the seed graphs depends on the structure of \( H_1 \), and so we will explicitly tether these two in the sequel, writing \((g_1, H_1), S_1\), and writing \( \mathcal{H}_n^m \) for the collection of all such feasible triples.

The aim then is to

1. First estimate the latent hierarchical structure \( H_2 \) of \( g_2 \); we will denote this estimate via \( \hat{H}_2 \), and we will let

\[
\hat{B}^i_{j,2} = \{v \in V(g_2) \mid \hat{H}_2(v, i) = j\}.
\]
Let the collection of all subgraphs in the partition defined by \( \tilde{H}_2 \) be denoted \( \tilde{B} \).

(2) Compute dissimilarity measurements

\[ \Delta : T_1 \times \tilde{B} \mapsto [0, 1] \]

between each subgraph of interest in \((g_1, H_1)\) and each subgraph defined by \( \tilde{H}_2 \). Note that \( \Delta \) ideally is a function that can compute the dissimilarity between any graph of order up to \( n \) and any graph of order up to \( m \); and the larger the value of \( \Delta \) between two graphs, the more dissimilar the networks. As \( \Delta \) needs to compute dissimilarities across graphs of different sizes and orders, it may encompass a collection of dissimilarity measures.

After a few preliminary definitions, we will be ready to define a HSN scheme.

**Definition 4** For a \( k \)-level hierarchical graph \((g, H)\) and \( i \in [k] \), let \( T^H_{g,i} \) denote the set of all total orderings of \( B^i \). Let \( T^H_g = \bigotimes_{i=1}^k T^H_{g,i} \).

For example, in the hierarchical subgraph depicted in Fig. 1, we have

\[
T^H_{g,2} = \left\{ \begin{bmatrix} B^2_2 \\ B^2_1 \\ B^2_3 \end{bmatrix}, \begin{bmatrix} B^2_3 \\ B^2_1 \\ B^2_2 \end{bmatrix}, \begin{bmatrix} B^2_1 \\ B^2_2 \\ B^2_3 \end{bmatrix}, \begin{bmatrix} B^2_3 \\ B^2_2 \\ B^2_1 \end{bmatrix}, \begin{bmatrix} B^2_2 \\ B^2_3 \\ B^2_1 \end{bmatrix}, \begin{bmatrix} B^2_1 \\ B^2_3 \\ B^2_2 \end{bmatrix} \right\}
\]

where

\[
\begin{bmatrix} B^2_2 \\ B^2_1 \\ B^2_3 \end{bmatrix}
\]

is shorthand for the ordering

\( B^2_1 > B^2_2 > B^2_3 \).

An example element of \( T^H_g \) is given by

\[
\left( \begin{bmatrix} B^1_3 \\ B^2_3 \\ B^3_3 \end{bmatrix}, \begin{bmatrix} B^2_3 \\ B^2_1 \\ B^2_2 \end{bmatrix}, \begin{bmatrix} B^2_2 \\ B^2_1 \\ B^2_3 \end{bmatrix} \right) = \left( \begin{bmatrix} B^1_3 \\ B^2_3 \\ B^3_3 \end{bmatrix}, \begin{bmatrix} B^2_3 \\ B^2_1 \\ B^2_2 \end{bmatrix}, \begin{bmatrix} B^2_2 \\ B^2_1 \\ B^2_3 \end{bmatrix} \right).
\]

As in the case of vertex nomination, if vertex labels in \((g_2, H_2)\) are uninformative, then an HSN must account for the indistinguishability of subgraphs with isomorphic structure. To wit, we have the following definition: For a \( k \)-level hierarchical network \((g, H)\) and \( i \in [k], j \in [n_1] \), we define

\[ \mathcal{I}(B^i_j; g) \]

\[
\{ = \ell \in [n_i] | B^i_{\ell} \subset B^i_{(j)} \} \] and there exists an automorphism \( \sigma \) of \( g \) with \( \sigma(B^i_{\ell}) = B^i_{j} \).

These are indices of the elements of the partition at level \( i \) that are indistinguishable from \( B^i_{j} \) without further information/supervision, and it is sensible to require that a HSN rank all these subgraphs as equally interesting. In vertex nomination, accounting for label uncertainty was achieved by means of an obfuscation function; in HSN, the uncertainty is not at the level of labels so much as at the level of subgraphs and this can be achieved by further requiring the dissimilarity \( \Delta \) satisfies the following condition:

\[ \Delta(B^i_{s_1}, \cdot) \text{ is constant over graphs indexed by } \mathcal{I}(\hat{B}^i_{s_2}; g_2) \text{ for each } s = (s_1, s_2) \in S_1. \]

We are now ready to define hierarchical subgraph nomination schemes formally.

**Definition 5** (Hierarchical Subgraph Nomination Scheme (HSN)) Let \((g_1, H_1), S_1 \in \mathcal{H}G\mathcal{S}_m^m\) and \((g_2, H_2), S_2 \in \mathcal{H}G\mathcal{S}_m^m\) and let \( T_1 \) be the parts of \( H_1 \) indexed by \( S_1 \). A Hierarchical subgraph nomination scheme is composed of two parts:

(i) An estimator \( \tilde{H}_2 = \tilde{H}_2(g_1, H_2, T_1) \) of the hierarchical clustering of \( g_2 \) provided by \( H_2 \); let the signature of \( \tilde{H}_2 \) be denoted \( \tilde{m}_2 = (\tilde{m}_{k,2}) \);

(ii) A dissimilarity \( \Delta \) satisfying Eq. (1) which produces a ranking scheme \( \Phi_{n,m} \), where

\[ \Phi_{n,m} (g_1, g_2, T_1, \tilde{H}_2) \in T^H_{g_2}. \]

For each level \( i \) of \( \tilde{H}_2 \), the subgraphs are ordered in \( \Phi_{n,m} \) via increasing value of

\[ \min_{s = (s_1, s_2) \in S_1} \Delta(B^i_{s_1}, \tilde{B}^i_{s_2}) \]

with ties broken in a fixed but arbitrarily manner.

### 2.3 Loss in HSN schemes

The goal of HSN is to effectively query \( g_2 \) given limited training resources from \( g_1 \). Given the resources required for a user to verify the interestingness of the returned subgraphs, we seek to maximize the probability that a priori unknown subgraphs of interest in \( g_2 \) are close to the top of the returned rank list.

For evaluation purposes, it is necessary (at least in theory) to be able to compare the true-but-unknown subgraphs of interest in \((g_2, H_2), S_2 \in \mathcal{H}G\mathcal{S}_m^m\) with elements of the HSN.
ranked list. In order to account for the fact that the dissimilarity used to construct the HSN scheme may be mis-specified (i.e., not captured exactly), the dissimilarity for verification will be denoted \( \Delta_E \) (the dissimilarity used for evaluation that defines the true but unknown subgraphs of interest), though we do not discount the possibility that \( \Delta_E = \Delta \) for any given scheme.

A logical loss function for HSN is motivated by the concept of precision in information retrieval. As the goal of HSN is to efficiently query large networks for structures of interest, at a given level \( k \), considering precision at \( i \) for \( 1 < i < \hat{m}_{k,2} \) enables us to model the practical loss associated with using a HSN scheme to search for \( \{B_{g2,2}^{g1}\}_{(i, s2) \in S2} \) (in \( g2, H_2 \)) given limited resources. There are two levels to the loss function for a given scheme \( \Phi_{n,m} = (\hat{H}_2, \Delta) \):

(i) The error in approximating \( H_2 \) via \( \hat{H}_2 \);

(ii) Given \( \hat{H}_2 \), the potential mismatch between \( \Delta \) and \( \Delta_E \).

Our loss function will account for these error sources as follows. Let \( F \) be a distribution supported on \( H(GS_n \times H(GS_m) \), and let

\[
((G_1, H_1), (G_2, H_2)) \sim F.
\]

Consider fixed subgraph of interest index sets \( S1 \) for \( G1 \) and \( S2 \) for \( G2 \), and let \( T1 \) (resp., \( T2 \)) be those subgraphs in \( (G1, H1) \) (resp., \( (G2, H2) \)) indexed by \( S1 \) (resp., \( S2 \)). Let \( \Phi_{n,m} = (\hat{H}_2, \Delta) \) be an HSN scheme and let the ranking provided at level \( k \) of \( \Phi_{n,m} \) be denoted via \( \Phi_{k,n,m}^k \) (with the implicit assumption that this is an empty list if \( \hat{m}_{k,2} = 0 \)).

**Definition 6** (HSN loss function, level-\((i, k)\) error at hierarchical level) With setup as above, for \( i, k < n \) we define the level-\((i, k)\) nomination loss with threshold \( t > 0 \) under dissimilarity \( \Delta_E \) via (where to ease notation, we will use \( \Phi_{k,n,m}^x \) for \( \Phi_{n,m}^x (g1, g2, T1, \hat{H}_2) \) and \( \Phi_{k,n,m}^x \) \( j \)) the \( j \)-th ranked subgraph in this list)

\[
\ell_{i,k,t} := \ell_{i,k,t}(\Phi_{n,m}, g1, H1, S1, g2, H2, S2, \Delta_E)
\]

where if we let \( J_{i,k} = \min(i, \hat{m}_{k,2}) \) and \( \mathbb{M}_{\Phi_{n,m}[j]} = \left\{ \Delta_E(\Phi_{n,m}[j], B_{g2,2}^{g1}) > t \right\} \)

\[
\ell_{i,k,t} := \left\{ \begin{array}{ll}
\frac{1}{J_{i,k}} \sum_{j=1}^{J_{i,k}} 1 \left\{ \mathbb{M}_{\Phi_{n,m}[j]} \right\} & \text{if } \hat{m}_{k,2} > 0 \\
1 & \text{if } \hat{m}_{k,2} = 0
\end{array} \right.
\]

For distribution \( F \) as above, the level-\((i, k)\) error with threshold \( t > 0 \) under dissimilarity \( \Delta_E \) of \( \Phi_{n,m} \) for recovering \( S2 \) is defined to be \( E_{F}(\ell_{i,k,t}) \). Letting \( S_{n,m} \) be the collection of all HSN schemes (i.e., the set of all ordered pairs of estimators and dissimilarities \( (\hat{H}_2, \Delta) \)), the Bayes optimal level-\((i, k)\) scheme with threshold \( t > 0 \) under dissimilarity \( \Delta_E \) is defined to be any scheme that achieves the Bayes’ error, which here is defined to be

\[
\min_{\Phi_{n,m} \in S_{n,m}} \mathbb{E}_F(\ell_{i,k,t}(\Phi_{n,m}))
\]

While the number of possible dissimilarities is indeed uncountably infinite, there are nonetheless a finite number of possible rankings that can be achieved via a combination of \( \hat{H}_2 \) and \( \Delta \) (hence min rather than inf in the Bayes error definition), and so the Bayes error is indeed achieved by at least one such pair.

### 2.4 User-in-the-loop supervision

Interactive machine learning, via incorporating user-in-the-loop supervision, can lead to an enhanced user experience and better downstream inference performance (Amershi et al. 2014). In subgraph nomination, the need for user-in-the-loop supervision can be understood as follows. Consider nominating in \( ((g2, H2), S2) \) at level \( k \), where \( S2 = \{ (k, \ell) \}, \) and \( \left| \left( B_{g2,2}^k ; g2 \right) \right| > 1 \). Even if \( \hat{H}_2 \) agrees with \( H2 \) at level \( k \), there are multiple subgraphs in \( g2, H2 \) that are isomorphic to the unknown subgraph of interest, and the HSN scheme has no information available to distinguish these. In this setting, it is reasonable to model the relative order of the elements in \( \left( B_{g2,2}^k ; g2 \right) \) in an HSN scheme as (effectively) uniformly random. For example, if \( \left| \left( B_{g2,2}^k ; g2 \right) \right| = c \), then considering top \( h \) of our nomination list \( (h < c) \), a scheme that identifies the correct structure for \( B_{g2,2}^k \) would still have probability

\[
\left( \begin{array}{c}
\frac{c-1}{h} \\
\frac{h}{c}
\end{array} \right) = 1 - \frac{h}{c}
\]

of not finding \( B_{g2,2}^k \). This can be mitigated by the following user-in-the-loop system:

(i) The user can take a limited number of single vertex inputs and can output whether that vertex is interesting or not (i.e., part of an interesting subgraph). This output can be modeled as error-free (oracle-user) or errorful.

(ii) The supervision can then be used to re-rank the subgraphs in the nomination scheme.

While practically, we do not suspect that there will be many perfect matches to the subgraphs of interest in the hierarchy provided by \( H2 \), it may be the case that there are many subgraphs “close” to the subgraph of interest in the (possibly lossy) estimate \( \hat{H}_2 \), in which case this supervision is equally necessary.
What follows is a formalization of the above heuristic. We first define the concept of a user-in-the-loop; noting that our definition is not the most general, as we focus in our analysis on binary users; i.e., they can only output 1 (yes) or 0 (no) for the interestingness of a vertex. In general, one might allow for the use to rating system with more levels, or even add a continuous space for responses.

**Definition 7 (VNUser)** Let \((g_1, H_1) \in \mathcal{HG}S_n\) with indices of interest \(S_1\) and \((g_2, H_2) \in \mathcal{HG}S_m\) with indices of interest \(S_2\). Let \((\theta, \gamma) \in [0, 1] \times [0, 1]\) and \(t \in \mathbb{Z} > 0\) with \(t < m\). We define the capacity of the user (i.e., only limited supervision is available) in the setting in which user-in-the-loop resources are costly and users by \(\eta\) acts on \(\mathcal{HG}S_n\) for \((g_2, H_2), S_2\) with parameters \((\theta, \gamma)\) (denoted \(U_t\)) as follows:

(i) Letting \(\Omega\) be the underlying sample space, the user is a function

\[
U_t : T_{V_2, t} \times \Omega \mapsto \{0, 1\}^t
\]

where \(T_{V_2, t}\) is the set of ordered \(t\)-tuples of distinct elements from \(V_2 = V(g_2)\). Note that dependence on \(\Omega\) will be suppressed when appropriate.

(ii) For each \(\eta \in T_{V_2, t}\) and \(x \in \{0, 1\}^t\), let \(\mathcal{F}_{\eta, i} = \{\eta_i \in \bigcup_{(s_1, s_2) \in S_2} B_{s_2, 2}^1\}, \mathcal{F}_{\eta, i} = \{\eta_i \notin \bigcup_{(s_1, s_2) \in S_2} B_{s_2, 2}^1\}\), define

\[
\mathbb{P}(U_t(\eta) = x) = \prod_{i=1}^t \left[ \left( \mathbb{I}\{\mathcal{F}_{\eta, i}\} \theta^x_i (1 - \theta)^{1-x_i} \right) + \left( \mathbb{I}\{\mathcal{F}_{\eta, i}\} \gamma^x_i (1 - \gamma)^{1-x_i} \right) \right]
\]

In essence, the user has independent binary components where the Bernoulli success probabilities depend only on the membership (or lack thereof) in a subgraph of interest.

For a given \((g_2, H_2), S_2\) and \(t\), we define the set of all such users by \(\mathcal{U}_t = \mathcal{U}_{(g_2, H_2), S_2}^t\).

Effectively, the HSN user-in-the-loop outputs a sequence of \(\{0, 1\}\) values, one for each vertex in a training set \(\eta\). An output of 1 is considered the positive answer, i.e., an interesting vertex; for vertices in \(\bigcup_{(s_1, s_2) \in S_2} B_{s_2, 2}^1\), this is a correct answer, and an error otherwise. An output of 0 is the negative answer meaning that this vertex is not of interest; for vertices not in \(\bigcup_{(s_1, s_2) \in S_2} B_{s_2, 2}^1\), this is a correct answer and is an error otherwise. This allows us to simultaneously model oracle users (\(\theta = 1\), and \(\gamma = 0\) and errorful users (\(\theta < 1\), and \(\gamma > 0\). The capacity of the user (i.e., \(t\)) allows us to model the practical setting in which user-in-the-loop resources are costly and only limited supervision is available.

Users can be incorporated into HSN schemes as follows.

**Definition 8 (User Aided HSN Scheme (UHSN))** Consider the setting in Definition (5). Let \(t \leq n_{(2)}\) and \(U_t \in \mathcal{U}_t\). Consider an HSN \(\Phi_{n, m} = (\hat{H}_2, \Delta)\), and let \(\eta\) be an ordered \(t\)-tuple of distinct elements of \(V_2 = V(g_2)\). For each \(k\), \(U_t\) acts on \(\Phi_{n, m}\) as follows:

(i) Given \(\eta, U_t\) is distributed according to Eq. (3). Let the output of the user process be denoted \(x \in \{0, 1\}^t\).

(ii) Let

\[
I_k := \{\ell \in [\hat{m}_k]\text{ s.t. more than half of the vertices in } \eta \cap \hat{B}_k^{1, 2}\}
\]

be the underlying sample space, the user is a function, were labeled interesting (i.e., as 1) by the user

\[
N_k := \{\ell \in [\hat{m}_k] \text{ s.t. at least half of the vertices in } \eta \cap \hat{B}_k^{1, 2}\}
\]

were labeled as not interesting (i.e., as 0) by the user

\[
\text{If } \eta \cap \hat{B}_k^{1, 2} = \emptyset, \text{ then } \ell \notin I_t \cup N_t.
\]

(iii) Let the indices of \(I_t\) be indexed via \((i_1, \ldots, i_t)\) where (according to \(\Phi_{n, m}\))

\[
\hat{b}_{i_1, 2} > \hat{b}_{i_2, 2} > \cdots > \hat{b}_{i_t, 2}.
\]

Similarly index \(N_t\) via \((n_1, \ldots, n_{|N_t|})\), and \(M_t := [\hat{m}_k] \setminus [I_t, N_t]\) (the unsupervised subgraph indices) via \((m_1, \ldots, m_{|M_t|})\) The user improved ranking is then given by

\[
\Phi_{n, m} = \left( \hat{b}_{i_1, 2} > \hat{b}_{i_2, 2} > \hat{b}_{i_3, 2} > \cdots > \hat{b}_{i_t, 2} \right) \cdot \left( \hat{b}_{m_1, 2} > \hat{b}_{m_2, 2} > \hat{b}_{m_3, 2} > \cdots \right).
\]

**Remark 4** We note here the possibility of even an oracle user introducing large errors into a ranking scheme based on an approximate \(\hat{H}_2\). Indeed, even if \(\hat{b}_{i_1, 2}^k\) is very similar an interesting subgraph in \(g_1\) according to 1, it is still possible that uninteresting vertices are chosen to provide to the user and \(\ell \in N_t\). This can be mitigated by choosing multiple vertices from each of some of the top ranked subgraphs for the user to evaluate.

**Remark 5** For iterative applications of the user-in-the-loop, we can sequentially run the user-in-the-loop, first on the output of \(\Phi_{n, m}\), then on \(\Phi_{n, m}^t\) (with \(t\) new user training points), and so on.
2.4.1 The (theoretical) benefit of the user-in-the-loop

We turn our attention now to take a look at the theoretical benefit of the user-in-the-loop in the setting of HSN. As in the motivating example for the user, the setting for this section will be as follows. Letting \((g_1, H_1) \in \mathcal{HGS}_n\) and \((g_2, H_2) \in \mathcal{HGS}_m\) with respective interesting index sets \(S_1\) and \(S_2\), consider a HSN scheme \(\Phi_{n,m}\) with \(\hat{H}_2 = H_2\). Consider the nomination provided by \(\Phi_{n,m}\) and the simple setting in which \(S_2 = \{(k, \ell)\} \in \mathcal{I}(B_{k,2}^r; g_2) = c > 1\). In this setting, it is reasonable to model the relative order of the elements in \(\mathcal{I}(B_{k,2}^r; g_2)\) in an \(\Phi_{n,m}\) (effectively) uniformly at random (though in practice, they are a fixed arbitrary order).

Theorem 1  Given the setup above where the scheme \(\Phi_{n,m}\) effectively ranks the elements of \(\mathcal{I}(B_{k,2}^r; g_2)\) uniformly at random at the top of the rank list, let \(t \leq c < m_k\) be the capacity of the user-in-the-loop. Consider any training set for the user where \(\eta\) contains exactly one element of each \(\Phi_{n,m}^k\) with \(1 \leq h \leq t\). Let \(E_h\) denote the event that \(\Phi_{n,m}^h\) ranks \(B_{k,2}^r\) not in the top \(h\). Considering \(c > t, h\), we have the following.

(i) Let \(U\) be an oracle user, then \(\mathbb{P}(E_h) = \max(1 - \frac{h+1}{c}, 0)\).

(ii) Let \((\theta, \gamma) = (1 - p, 0)\) for \(p < 1\), then

- If \(t + h \leq c\), then \(\mathbb{P}(E_h) = 1 - \frac{h}{c} - (1 - p)\frac{h}{c}^2\); note that this is strictly less than \(1 - h/c\) for all \(p \in (0, 1)\).

- If \(t + h > c\), then \(\mathbb{P}(E_h) = \frac{1 - p}{c}\); note that if \(p < (c - h)/t\), this is strictly less than \(1 - h/c\).

(iii) Let \((\theta, \gamma) = (1 - p, q)\) for \(p < 1\) and \(q > 0\). Then, letting \(F(i; n, p)\) be the CDF of a Binomial\((n, p)\) random variable evaluated at \(i\), we have

- If \(h + t \leq c\), and \(t \leq h\), then \(\mathbb{P}(E_h) = 1 - \frac{h}{c} - (1 - p - q)\frac{h}{c}^2\); note that if \(p + q < 1\), this is less than \(1 - \frac{h}{c}\).

- If \(h + t > c\), and \(t \leq h\), then \(\mathbb{P}(E_h) = \frac{pt}{c} + \frac{1}{c} \sum_{i=1}^{c-h} F(i - 1; t, 1 - q)\); this is upper bounded by \((p + q)t/c\) and if \((p + q) < (c - h)/t\), this is strictly less than \(1 - h/c\).

- If \(h + t \leq c\), and \(t > h\), then

\[
\mathbb{P}(E_h) = 1 - \frac{h}{c} - (1 - p)\frac{h}{c}^2 + \frac{1 - p}{c} \sum_{i=1}^{c-h} F(i - 1; t, 1 - q) + \frac{1}{c} \sum_{i=c-h+1}^{t} F(i - 1; t, 1 - q)
\]

Note that a rough upper bound of this is given by

\[
1 - h/c + qt/c - (1 - p)h/c, \text{ and if } qt < (1 - p)h, \text{ this is strictly less than } 1 - h/c.
\]

- If \(h + t > c\), and \(t > h\), then

\[
\mathbb{P}(E_h) = \frac{pt}{c} + \frac{1 - p}{c} \sum_{i=1}^{c-h} F(i - 1; t, 1 - q)
\]
For $c = 50$ we plot ($h, t \in \{1, 2, \ldots , 49\}$) $\min(R(c,t,h,p,q), 1)$ for the loss function in the setting of Theorem 1 part (iii) and $R$ defined in Eq. (4). Different panels correspond to different values of $p$ and $q + 1$:

$$F(i-1; t, 1 - q)$$

Note that a rough upper bound of this is given by $(1 + q)(1 - (1 - h/p)h)$, and if $(1 + q)t + ph < c$, this is strictly less than $1 - h/c$.

We, perhaps surprisingly, see that there are $p$ and $q$ values that guarantee improvement (over the user-in-the-loop-free setting) in all cases. The conditions in part iii. bear further consideration. The interaction of $p$, $q$, $h$, $t$ here is nuanced, and is most easily analyzed numerically, as is shown in Figs. 2 and 3. Letting $L(c, t, h, p, q)$ denote the loss from Theorem 1 part (iii) (where we can interpret this in the context of Definition 6 by considering $\Delta$ as a 0/1 oracle dissimilarity function), for $c = 50$ we plot the relative loss improvement,

$$R(c, t, h, p, q) = \frac{L(c, t, h, p, q) - (1 - h/c)}{1 - h/c}$$

(4)

for $h$ between 1 and 49 (on the $y$-axis), and $t$ between 1 and 49 (on the $x$-axis). Different panels correspond to different values of $p$ (varies by row) and $q$ (varies by column). Note that in Fig. 2, we plot the relative loss in the left panel $R(c, t, h, p, q)$ and $\min(R(c,t,h,p,q), 1)$ in the right panel; this truncating aids in distinguishing the areas of improvement from those where $R(c,t,h,p,q) > 0$.

In each plot, darker red areas correspond to parameter settings where the loss is improved with user-supervision, and darker blue areas to parameter settings where the loss is.
We consider 2000 Monte Carlo replicates of distributions as in Sect. 3.1.1, and from each distribution, we generate 100 networks. We use Method 1, and performance with an oracle user (varying the replies for the user; i.e., the amount of supervision provided) and the chance algorithm are compared. We plot the density of all trials (y-axis) that ranked the correct block in the nth place (x-axis).

increased with user-supervision (due to the user error). A few trends are clear from the figures. Unsurprisingly, supervision becomes detrimental (i.e., more supervision leads to more loss) as \( p \) and \( q \) increase and \( q \geq 0.5 \), although the loss appears to be more tolerant of higher values of \( p \) than it is of \( q \). In all cases, large values of \( t \) and \( h \) simultaneously lead to training becoming less effective, though \( h < t \) seems to yield resilience to larger user-supervised loss for all \( p, q \) pairs.

In the event that the hierarchical clustering is noisily recovered in \( g_2 \) (i.e., \( \hat{H}_2 \neq H_2 \)), even the supervision provided by an oracle user-in-the-loop may be worse than no extra supervision. Indeed, consider the setting where \( S_2 = \{(k, \ell)\} \) and

\[
\alpha = \frac{\lvert \hat{B}_{k,2} \cap B_{k,2}^* \rvert}{\lvert \hat{B}_{k,2}^* \rvert}; \quad \beta = \min_{i \neq \ell} \frac{\lvert \hat{B}_{i,2} \cap B_{i,2}^* \rvert}{\lvert \hat{B}_{i,2}^* \rvert},
\]

then, adopting the notation and setting above, the probability that \( \hat{B}_{k,2} \) is not ranked in the top \( h \) after oracle \( U_t \) supervision is bounded below by the probabilities in part (iii) of Theorem 1 with \( p = \alpha \) and \( q = \beta \). In particular, there are values of \( \alpha \) and \( \beta \) under which this error is worse than \( 1 - h/c \) (the error sans additional supervision).

Fig. 5

3 Simulations and real data experiments

We now provide empirical evidence for the theory we have outlined. Namely, we will show through simulations and real data examples the impact of a user-in-the-loop. We first con-
consider simulations where graphs are drawn from an HSBM distribution. In this setting we evaluate our methodology on the task of nominating subgraphs with a similar motif to our subgraph of interest. We then consider 57 pairs of analyzed human neural connectomes for which brain regions (i.e., communities) are well defined and the community for each voxel is known. We evaluate our methodology on the task of nominating the same brain region within a given pair of connectomes with varying levels of a user-in-the-loop.

### 3.1 Simulations

In this section we will first provide a complete description of the model and procedures that were employed to test our theory in a simulated data setting. We then discuss results and how they fit within our theory.

#### 3.1.1 Model description

For our first example, we consider nominating within the Hierarchical Stochastic Blockmodel setting of Lyzinski et al. (2017). We consider sampling from a 2-level HSBM that has 16 blocks in the first level and 3 sub-blocks in each of the first level blocks (so that this model can also be realized as a 48 block standard SBM). Further, the blocks of the first level belong to one of three motifs $B_1$, $B_2$, or $B_3$, with constant cross-community connection probability $p = 0.01$ for the first level. Formally, we are sampling from the following 2-level HSBM:

(i) There are $K_1 = 16$ blocks in the first level of the hierarchy, and the size of the blocks is i.i.d. $10 \times [20 + 50 \times \text{Unif}(0, 1)]$; note that this differs slightly from the random block assignment HSBM defined previously.

(ii) The motifs $B_i \in \mathbb{R}^{3 \times 3}$ for $i = 1, 2, 3$ are i.i.d. samples satisfying $B_i = X_i^T X_i$ where $X_i \in \mathbb{R}^{3 \times 3}$ satisfies $X_i[\cdot, j]^i.i.d. \sim \text{Dirichlet}(1, 1, 1)$; of course, this distribution is artificial, but it provides a maximum entropy sampling of the $X_i$’s which provides a difficult testing setting for our algorithm.

(iii) For each $j \in [16] \setminus [9]$, we have $B_j^i = B_i$ independently with probability $1/3$ for $i = 1, 2, 3$. $B_2^3$ is set to equal $B_1^2$;

(iv) In the second level of the hierarchy ($K_2^j = 3$ for all $j$), the block sizes are defined via (where $\lceil \cdot \rceil_1$ rounds the number to the nearest tenth)

$$|b^{(2)}_{j,i}| = |\{ v \in V | b^{(1)}(v) = j, b^{(2)}(v) = i \}|$$

where the entries of $\omega = (\omega_1, \omega_2, \omega_3)$ are

$$\omega_i \overset{i.i.d.}{\sim} \text{Unif}(2, 10).$$

Lastly, we set

$$|b^{(2)}_{j,3}| = |\{ v \in V | b^{(1)}(v) = j, b^{(2)}(v) = 3 \} = |b^{(1)}_j| - |b^{(2)}_{j,i}|- |b^{(2)}_{j,3}|$$

This particular parameterization is chosen to produce different motifs that stress our HSGN framework by producing motifs that are very similar, and motifs where the block structure is extremely subtle (i.e., close to flat), and cases where both these problems happen, as it draws from a distribution in which it is not uncommon that the motifs turn out to be extremely similar and/or very flat.

The block structure was inferred by clustering via Gaussian mixture modeling after embedding, utilizing the R package Mclust (Fraley and Raftery 1999) to cluster the graph we are nominating from into 8 sub-graphs. After that, we use two different methods to infer similarity. The first approximates $\Delta$ via the value of the non-parametric test statistic of Tang et al. (2017) computed between re-embeddings of each inferred community (as in Lyzinski et al. (2017)); simulation results are shown in Figs. 4 and 5. We will call this Method 1 in the sequel. The second approximates $\Delta$ between communities $i$ and $j$ via the scaled graph matching pseudo-distance (Fishkind et al. 2019),

$$1 - \frac{\min_{P \in \Omega(n)} \| A_i - P A_j P^T \|_F^2}{\frac{1}{n^2} \sum_{P \in \Omega(n)} \| A_i - P A_j P^T \|_F^2}$$

where either $A_i$ (the induced subgraph of community $i$) or $A_j$ has been appropriately padded as in Fishkind et al. (2019)); results are shown in Figs. 6 and 7. We will call this Method 2 in the sequel.

The user replies are those of an oracle. However, as discussed previously since we do perfectly recover the true network partition (rather than using an estimate of the true network partition), our user could still end up being errorful. In particular, the user could be asked to rank vertices that are not in the subgraph of interest. This exemplifies the behavior characterized by $iii$ of Theorem 1, albeit with a different $q$ for each of the blocks. Another significant difference in the scheme of the simulation is that the nomination process takes the first affirmative reply as the correct community and returns the others in their order as is.
Fig. 7 We consider 2000 Monte Carlo replicates of distributions as in Sect. 3.1.1, and from each distribution, we generate 100 networks. We use Method 2 and performance with an oracle user (varying the replies for the user; i.e., the amount of supervision provided) and the chance algorithm are compared. We plot the density of all trials (y-axis) that ranked the correct block in the nth place (x-axis).

Fig. 8 With perfect knowledge of the block structure of the brain (the division into 71 regions provided by the data) and using Method 1, (left) we plot on the y-axis the proportion of subjects (averaged across all blocks considered as the block-of-interest in the left hemisphere) that ranked the correct block in the right hemisphere the n-th place (x-axis); (right) we plot on the y-axis the the proportion of subjects (averaged across all blocks considered as the block-of-interest in the left hemisphere) for which we find the total proportion of the block of interest in the top 25 vertices nominated (x-axis).
Fig. 9 Clustering the graph using Gaussian Mixture Modeling on the embedded network, in the left two panels, we use Method 1 to estimate $\Delta$, while the right two panels we use Method 2. In the top row, on the $y$-axis we plot the proportion of subjects (averaged across all blocks considered as the block-of-interest in the left hemisphere) that ranked the correct block in the right hemisphere the at worst $x$-th place ($x$-axis). In the bottom row, on the $y$-axis we plot the proportion of subjects for which we find the total proportion of the block of interest in the top $25$ vertices nominated ($x$-axis).

3.1.2 Results and discussion

In this section we will go over the results of the experiments we described. The aim is to bolster the claims in the theoretical results with experimental results, as such we will have a special focus on the features that are iconic from the theory. First we start with Method 1, the experiment that uses the test statistic form Tang et al. (2017) to estimate the dissimilarity function.

In Fig. 4 we can see that the user training provides an improvement over the original algorithm (“No user”), as suggested by Theorem 1, with more training often yielding superior performance. Moreover, especially for small $x$-values, the algorithm outperforms a chance ranking of the obtained blocks. Interestingly, we see that more user-supervision is not always better. Indeed, as mentioned previously, in our regime of lossy block recovery additional supervision can be deleterious. Figure 5 elucidates this finding further from the perspective of the distribution density of the correct block in each position. We notice that as user-supervision increases the distribution are shifted towards putting the correct block in the first position. We also notice the trend predicted by Fig. 3, as we increase $h$ (the nomination error level) the ideal value of $t$ (user-provided replies) is decreasing, and less supervision is preferred. We see this, as the curves with less supervision overtake those of higher supervision as we plot $1 - L_n(\phi, B^*)$. However we note that here the value of $q$ is not constant and hence the minimizers of the loss function do not not adhere to the figure exactly. In Figs. 6 and 7 we see similar trends using Method 2. Here the main contribution to the error of the methods is the misclustering of the vertices in the initial GMM step, and both dissimilarity estimates provide good estimates of the latent $\Delta$. In the real data example considered below, there is a more pronounced differentiation across methods.
3.2 Subgraph nomination in BNU1 connectomes

Data acquisition resources and limitations can be eased through automation. For example, in the study of human connectomes, one of the more labour intensive steps, is classifying different parts of the brain. One such classification task is finding corresponding regions of interest across hemispheres in a human connectome. This task is time and human resource intensive when done manually, and automation is essential for achieving a high throughput (Gray et al. 2012). To illustrate our HSN methodology further, we will consider then the task of nominating brain regions across hemispheres in the DT-MRI derived brain networks in the BNU1 database (Zuo et al. 2014). The spatial image of the brain includes information about its structure; in other words, the neural fibers that run through different areas of the brain. A graph of the subject’s brain is then created with the following definitions: nodes or vertices correspond to different spatial units (usually the brain is divided into cubic sections called volumetric pixels, or voxels), edges are weighted based on the amount of neural fibers that run through two regions. One may hypothesize that there is a sense of structural symmetry between paired regions across hemispheres, and discovery of this pairing is the inference task for this HSN application.

Here, we apply the user aided subgraph nomination scheme to a connectomic dataset. The data we apply this to is a set of 114 neuronal networks—two repeated scans for each of 57 human subjects—derived from the BNU1 connectome dataset (Zuo et al. 2014). Each brain scan sections the brain into one thousand voxels—or volumetric pixels—with weighted edges between them indicating the amount of neural batches that are shared, indicating connectedness in a structural but not necessarily functional sense. Additionally each voxels contains information on its membership to the left or right hemispheres, membership to one of the 71 neuronal regions (Desikan et al. 2006), whether it is grey or white matter, and its XYZ coordinates in the partially registered scan. The relative size of the networks is then ≈ 1000 vertices, with pre-defined neuronal regions varying in size between 2 and 100 vertices. As in the simulation, although the distributional properties in paired regions across hemispheres is often similar (i.e. similar motifs), there are no guarantees that the numbers of vertices that make up each region are equal. Errors from this (and misclustering) are magnified in smaller subgraphs, especially since there is less information about the motif in the small subgraph setting. The aforementioned sources of errors are mitigated by only considering paired regions that have at least 10 vertices in each hemisphere and using similarity metrics (Methods 1 and 2) that do not depend on the size of a region.

Our procedure can then be described as follows, considering each pair of sufficiently large matched regions (≥ 10 vertices in each hemisphere) separately, first we use the information given by spectrally embedding the adjacency matrix of the connectome (and the collected spatial coordinate data in Fig. 10) to infer the subgraph structure in the right hemisphere via clustering by GMM (Fraley and Raftery 1999), using the region-of-interest in the left hemisphere as our training, next we use Methods 1 and 2 to rank above to provide different estimates of Δ. We then supply a user-in-the-loop with a vertex (or a few vertices) from each of the top k communities to re-rank the nomination list. We plot the performance of our algorithms with perfect knowledge of the block structure of the brain (the division into 71 regions provided by the data) and using Method 1 in Fig. 8. We plot:

- (Left) on the y-axis the proportion of subjects (averaged across all blocks considered as the block-of-interest in the left hemisphere) that ranked the correct block in the right hemisphere in the at worst x-th place (x-axis).
- (Right) on the y-axis the the proportion of subjects for which we find the total proportion of the block of interest in the top 25 vertices nominated (x-axis).

Note that an artifact of the metric in panel (b) is that we may never find a complete block that is of a size larger than 25 this is why we see a drop near 90%. As ideal performance in the left and right panels corresponds to the function f(x) ≡ 1, this figure enforces the intuition that, given high fidelity clusters, our ranking procedure is significantly better than chance and benefits strongly from user-in-the-loop supervision.

In Fig. 9, in the setting of potentially errorful clusters (obtained via McIust applied to the adjacency spectral embeddings of the brain networks, we plot (similar to in Fig. 8), in the top panels the proportion whose match is found in the top x (similar to the left panel of Fig. 9) and in the bottom panels the percent who had the proportion of the top 25 vertices come from the block of interest (similar to the right panel of Fig. 9). The left two panels use Method 1 to estimate Δ, while the right two panels use Method 2. In light of perfect performance corresponding the the constant 1 function in all cases, we see here that Method 2 achieves better performance (especially for large values on the x-axis in each figure) than Method 1 both when making use of the user-supervision and in the no-user setting. Here, we see the general trend that the methods perform poorly without a user-in-the-loop (due to the error in the clustering; see Fig. 10), but that the method can efficiently make use of the user to achieve better performance. As was the case previously, the clustering is errorful, and on average this can have a deleterious effect on the user-supervision (top row). However, in the bottom row we see that sample-wise the supervision is monotonically beneficial.

We repeat the above experiment using Method 2 and incorporating the XYZ coordinates of each voxel into the clustering (by appending the features onto the spectral graph embeddings), and plot the results in Fig. 10. The increased
clustering fidelity achieved by incorporating the vertex features manifests itself here by the performance gain achieved in the no-user setting when compared with Fig. 9. As a result of this, better performance is achieved across all settings here (again when compared with Fig. 9). While the clustering here is still errorful, and on average this can have a deleterious effect on the user-supervision, the right panel again shows that sample-wise the supervision is monotonically beneficial. This means that a user can successfully aid in finding larger portions of the block of interest. Therefore, we can use these vertices in the input to further investigate and find the rest of the block, by searching in the $k$-nearest neighbours of these vertices.

### 3.3 Subgraph nomination on the Reddit social network

Social network moderation is another area where the biggest limitation is the human resources required to analyze the prohibitively large amount of data. One problem that appears in Reddit, is finding subreddits for which the name has changed. This can be needed to track a previously banned subreddit whose users create new accounts and a new subreddit. Further, since this experiment depends only on structure, which is informed by user interaction patterns, it could be used for recommender systems, or finding suspicious behaviour in other subreddits. The first is under the assumptions that people who communicate in the same way may enjoy similar other subreddits. These form our node types, the focus here is on structure so we do not maintain node types. We also dispose of the users, as they will form a leaf coming out of each comment and thread. Each of these comes with a multitude of features, notably parent comment/thread, and containing subreddit. All features are discarded for a structure based approach, however we use the relations above to create the graph. Our edges represent $S \leftrightarrow T$, $T \leftrightarrow C$ if $T$ is not parent of $C$, else $T \leftrightarrow C$, $C_1 \leftrightarrow C_2$ if $C_1$ is parent of $C_2$, and $S \leftrightarrow F$. A graph is constructed for the dataset, then the subgraph for each subreddit is retrieved. Using the post-time date each month can be subset to 4 week time periods. The data is subset to subreddits of size 1000–2000 nodes.

The Graph Matching Network from Yujia et al. (2019), a Graph Neural Network approach for comparison at the graph level, is used as the graph similarity (or dissimilarity) function. The neural network embeds graphs into a space where the closer two graphs are to one another the more similar they are, the normalized embeddings provide a similarity metric. At this point, normally the top $h$ recommendations are passed to an expert to analyse. The expert would analyze the subreddits to make a decision. Here, we can insert a user in the loop, the user would consider a node (or small collection of nodes) in the graph instead of the entire graph. Here, where we are looking to recover the subreddits in time, the node-level user defines a correct match if the node is in the subreddit of interest. Training is done on the first 2 weeks of the reddit data, and the remaining two weeks are used to test the neural network’s performance and implement the user. Note that the user input can be used as a node feature and implemented during the training as well; we do not pursue that here.

After the training is complete and the tests are performed, the top 8 subgraphs are nominated, and we simulate VN users-in-the-loop with different values for $\theta$ and $\gamma$. The result is displayed in Fig. 11, where the x-axis is the number of GMN training steps and the y-axis represents the retrieval accuracy. Accuracy here is the percentage of subgraphs for which the correct community is nominated in the top $\ell$ graphs. Denoting the user $U_\ell$ with parameters $(\theta, \gamma)$ with $VN(\theta, 1 - \gamma)$ (recalling that $\theta$ and $1 - \gamma$ are the probabilities of correct user input for true positive and true negative nodes resp.), the left plot is the accuracy for $\ell = 2$ and on the right is for $\ell = 8$.

Notice in Fig. 11 the addition of the user-in-the-loop helps achieve better results especially at the lower level $\ell = 2$. Also, we can achieve desirable results with less training, even with a faulty VN user. The performance at $\ell = 2$ suggests the user can lower the costs of verification, as an expert user needs to look at less subgraphs to determine whether the nomination is successful. Moreover, this suggests that we can automate the VN user-in-the-loop by using node features to make decisions, and train the GMN for a shorter amount of time to reduce the computational cost. This is critical since the GMN computational cost grows quadratically with the total number of nodes in the graphs.

### 4 Conclusions and future direction

In this paper we have introduced a formal, and versatile framework for the novel subgraph nomination inference task, with special emphasis paid to the utility of users-in-the-loop in the context of subgraph nomination. Subgraph nomination is an important tool for structural queries within and across networks, and we demonstrated its utility in both real and synthetic data examples. In addition, the relatively simple users-in-the-loop formulation outlined herein can easily be lifted to more complex cases including supervision done over several iterations, and can be simply extended to the case of
Fig. 10  Clustering the embedded graphs using Gaussian Mixture Modeling incorporating the XYZ coordinate data and using Method 2 to estimate $\Delta$. In the left panel, on the $y$-axis we plot the proportion of subjects (averaged across all blocks considered as the block-of-interest in the left hemisphere) that ranked the correct block in the right hemisphere the at worst $x$-th place (x-axis). In the right panel, on the $y$-axis we plot the proportion of subjects (averaged across all blocks considered as the block-of-interest in the left hemisphere) for which we find the total proportion of the block of interest in the top 25 vertices nominated (x-axis).

Fig. 11  Clustering the graphs using their subreddit membership and using GMN to estimate $\Delta$. The $y$-axis we plot the proportion of subreddits (averaged across 100 runs of the VN user) that ranked the correct subreddit in the second time period. The $x$-axis is the training time. In the left panel, we look at the top 2 nominations. In the right panel, we look at the top 8 nominations.

Multiple users. The theory and experiments included herein highlight the important role that user supervision can play in effective information retrieval.

Further, this paper shows the importance of analysing the original algorithm and the validity of user input before including them. For example, if the inferred clustering is incorrect in a hierarchical subgraph nomination framework, in the sense that the correct clusters’ vertices are evenly spread among the inferred blocks, then our resources should be focused on collecting relevant data that would improve the inferred blocking structure, as even oracle user supervision in this case could be detrimental to the performance. In our approach to subgraph nomination, inferring high-fidelity clusters is essential, and we have demonstrated that clustering can be improved by incorporating informative vertex features. Exploring this further, understanding the information theoretic gain of features on our nomination performance as in Levin et al. (2020), is an open area of research that we are actively pursuing.

The validity of the user input itself is also important to consider when thinking about adding a user-in-the-loop. While we have theoretically demonstrated the possibility of increased performance even with an errorful user, in practice the situation is significantly more nuanced. The user errors, rather than being uniformly random, may be systematic or deterministic. For example an adversarial attacks could manipulate the results of the search using a user bot attack. More nuanced users-in-the-loop demand more nuanced theory to understand their broad effects. For example, an adversarial attacks could manipulate the results of the search using a user bot attack. More robust
users would be, most likely, more costly and a cost-benefit optimization analysis would be needed in these more nuanced setting to tease out the positive (or negative) impact of incorporating the user.

Another avenue that has not been investigated explicitly but for which the framework is still sufficient is the situation with multiple blocks of interest. For that we need to adjust the definition of the VN-user to accommodate for different values of $p$ for each block of interest. Then, after some necessary adjustment to the loss function, one may obtain results generalized to the case of multiple blocks of interest in the simple user-in-the-loop setting. On the other hand, one may want to extend the results of Theorem 1 to more nuanced users.

We close with a discussion of scalability and computational complexity. For implementing the core subgraph nomination routine, the computational burden is in graph partitioning and subgraph comparison, both of which can be implemented efficiently; e.g., any number of scalable clustering routines for graph division, and fast graph similarity algorithms (e.g., Bai et al. (2019)). For the VN user-in-the-loop it will greatly depend on the setting, essentially we randomly retrieve an element from each of the top $h$ communities. We then apply the user, if the user is an expert then the cost is not only computational (if the expert needs to perform some analysis on the node’s feature for example) but also in human resources, which is beyond the scope of this paper, but intuitively scalability is an issue. If an automated user is used (e.g., one who is trained on the node features alongside the algorithm) then the cost is that of training and $h$ times the cost of retrieving the model’s fit on the selected nodes, here scalability is less of an issue.

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### Appendix I: Proof of Theorem 1

For $i \leq j \leq c$, let the event $A_{i,j}$ be the event that the unknown subgraph of interest in $g^2$ has rank in $[i, j]$. Let $E_h$ be the event that after user-in-the-loop supervision, the subgraph of interest has rank strictly greater than $h$.

**Part I:** With an oracle user-in-the-loop, we have that

$$
P(E_h) = P(E_h | A_{[1,h]} ) P( A_{[1,h]})
$$

$$
+ P(E_h | A_{[h+1,h+t]} ) P( A_{[h+1,h+t]} )
$$

$$
= 0 + P(E_h | A_{[h+t+1,m_k]} ) P( A_{[h+t+1,m_k]} )
$$

$$
= P( A_{[h+t+1,m_k]} )
$$

If $h + t \leq c$, then $P( A_{[h+t+1,k]} ) = 1 - \frac{h+t}{c}$, else it is 0. Hence, $P(E_h) = \max(0, 1 - \frac{h+t}{c})$ as desired.

**Part II:** When the user has probability $p$ of misidentifying the vertex from the subgraph of interest as non-interesting, then when $h + t \leq c$,

$$
P(E_h) = P(E_h | A_{[1,t]} ) P( A_{[1,t]} )
$$

$$
+ P(E_h | A_{[t+1,h+t]} ) P( A_{[t+1,h+t]} )
$$

$$
= 0 + P(E_h | A_{[h+t+1,m_k]} ) P( A_{[h+t+1,m_k]} )
$$

$$
= p P( A_{[1,t]} ) + P( A_{[h+t+1,m_k]} )
$$

$$
= \frac{pt}{c} + 1 - \frac{h+t}{c} = 1 - (1-p) \frac{t}{c} - \frac{h}{c}.
$$

When $m_k > h + t > c$, the above probability is simply $\frac{pt}{c}$.

**Part III:** When the user has probability $p$ of misidentifying the vertex from the subgraph of interest as non-interesting and probability $q$ of misidentifying a non-interesting vertex as interesting, then when $h + t \leq c$ and $t \leq h$,

$$
P(E_h) = P(E_h | A_{[1,t]} ) P( A_{[1,t]} )
$$

$$
+ P(E_h | A_{[t+1,h]} ) P( A_{[t+1,h]} )
$$

$$
+ \sum_{i=1}^{t} \left[ P(E_h | A_{[h+i,h+i]} ) P( A_{[h+i,h+i]} ) \right]
$$

$$
+ P(E_h | A_{[h+t+1,m_k]} ) P( A_{[h+t+1,m_k]} )
$$

$$
= \frac{pt}{c} + \sum_{i=1}^{t} \left[ \frac{1}{c} \sum_{j=0}^{i-1} \left( \frac{t}{j} \right) (1-q)^j q^{t-j} \right]
$$

$$
+ 1 - \frac{h+t}{c}
$$

$$
= \frac{pt}{c} + \sum_{j=0}^{t-1} \left[ \frac{1}{c} \sum_{j=0}^{t-j} \left( \frac{t}{j} \right) (1-q)^j q^{t-j} \right]
$$

$$
+ 1 - \frac{h+t}{c}.$$
\[ \frac{pt}{c} + \sum_{j=0}^{t} \frac{t-j}{c} (1-q)^{j} q^{t-j} \]
\[ + 1 - \frac{h+t}{c} \]
\[ = \frac{pt}{c} + \frac{qt}{c} + 1 - \frac{h+t}{c} \]

When \( h + t > c \) and \( t \leq h \),

\[ \mathbb{P}(E_h) = \mathbb{P}(E_h \mid A_{[1,t]}) \mathbb{P}(A_{[1,t]}) \]
\[ + \sum_{i=1}^{c-h} \mathbb{P}(E_h \mid A_{[h+i,h+i]}) \mathbb{P}(A_{[h+i,h+i]}) \]
\[ = \frac{pt}{c} + \frac{1}{c} \sum_{i=1}^{c-h} F(i - 1; t, 1 - q) \]

When \( t > h \), and \( h + t \leq c \), then (assuming \( c \geq t, h \))

\[ \mathbb{P}(E_h) = \mathbb{P}(E_h \mid A_{[1,h]}) \mathbb{P}(A_{[1,h]}) \]
\[ + \sum_{i=1}^{t-h} \mathbb{P}(E_h \mid A_{[h+i,h+i]}) \mathbb{P}(A_{[h+i,h+i]}) \]
\[ = \frac{pt}{c} + \frac{1}{c} \sum_{i=1}^{t-h} F(i - 1; h + i - 1, 1 - q) \]
\[ + \frac{1}{c} \sum_{i=t-h+1}^{t} F(i - 1; t, 1 - q) \]

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