

# AUTO-TUNE: FINDING AN OPTIMAL DISTANCE THRESHOLD FOR INFERRING HIV TRANSMISSION CLUSTERS

Steven Weaver<sup>1\*</sup>, Vanessa Davila Conn<sup>2</sup>, Hannah Verdonk<sup>1</sup>, Joel O. Wertheim<sup>3</sup>, and Sergei L. Kosakovsky Pond<sup>1</sup>

<sup>1</sup> Center for Viral Evolution, Temple University, Philadelphia, PA, USA

<sup>2</sup> Center for Research in Infectious Diseases, National Institute of Respiratory Diseases, Mexico City, Mexico

<sup>3</sup> Department of Medicine, University of California San Diego, La Jolla, CA, USA

Correspondence\*:  
Steven Weaver  
sweaver@temple.edu

## 2 ABSTRACT

3 Molecular surveillance of viral pathogens and inference of transmission networks and clusters  
4 from genomic data play an increasingly important role in public health applications and transmis-  
5 sion mitigation efforts, especially for HIV-1. For many methods, the genetic distance threshold  
6 used to connect sequences in the transmission network is a key parameter informing the proper-  
7 ties of inferred networks. Using a distance threshold that is too high can result in a network with  
8 many spurious links, making it difficult to interpret and analyze. **HV: I'd clarify that a high distance**  
9 **threshold results in a larger network as well, not just a more densely connected one..** On the  
10 other hand, using a distance threshold that is too low can result in a network with too few links,  
11 which may not capture key insights into rapidly growing clusters among patients with shared  
12 attributes that could benefit from public health intervention measures. Published research using  
13 the HIV-TRACE software package frequently uses the default threshold of 1.5% for HIV pol gene  
14 sequences, but in many cases, investigators select other threshold parameters (heuristically) to  
15 better capture the underlying dynamics of the epidemic they are studying.

16 Here, we present a heuristic scoring approach for tuning a distance threshold adaptively, which  
17 seeks to prevent the formation of giant clusters (measured as the ratio  $R_{12}$  of the sizes of the  
18 largest and the second largest cluster), and to maximize the number of clusters present in the  
19 network.

20 We apply our scoring heuristic to outbreaks with different characteristics, such as regional  
21 or temporal variability, and demonstrate the utility of using the scoring mechanism's suggested  
22 distance threshold to identify clusters exhibiting risk factors that would have otherwise been  
23 more difficult to identify. For example, while we found that a 1.5% distance threshold is typical  
24 for US-like epidemics, recent outbreaks like the CRF07\_BC subtype among men who have sex  
25 with men (MSM) in China have been found to have a lower optimal threshold of 0.5% to better  
26 capture the transition from injected drug use (IDU) to MSM as the primary risk factor. Alterna-  
27 tively, in communities surrounding Lake Victoria, where there has been sustained transmission

for several years, we found that a larger distance threshold is suitable to capture a more risk factor-diverse populace with sparse sampling over a longer period of time. Such identification may allow for more informed intervention action by respective public health officials.

**Keywords:** molecular epidemiology, HIV, network, transmission cluster, surveillance

## 1 INTRODUCTION

The use of genomic data to infer and characterize transmission networks of various pathogens has taken off in the past two decades, with applications to **make a list of citations to cover a broad list of organisms**. Choosing an appropriate genetic distance threshold is an important part of using a molecular transmission network to track the spread of rapidly evolving pathogens Liu et al. (2020); Rose et al. (2020). This distance threshold determines how close pathogen sequences isolated from two individuals must be in to link them with as putative transmission partners in the network **HV: the grammar in the previous sentence is confusing**. Using a distance threshold that is too small can result in a network with many spurious, i.e. epidemiologically uninformative, links, making it difficult to interpret and analyze. On the other hand, using a distance threshold that is too large can result in a network with too few links, underestimating connections between individuals and making it difficult to accurately track the spread of the disease Gore et al. (2022).

To optimize the utility of inferred transmission networks, it is important to carefully consider the appropriate distance threshold,  $d$ . This threshold may vary depending on the specific disease and the context in which it is spreading. For example, a highly contagious respiratory illness may require a smaller  $d$  than a less contagious illness that is primarily spread through direct contact. Viruses are more amenable to molecular studies compared to bacteria due to their high genetic divergence and compact genomes. Given the relatively high evolutionary rate of RNA viruses (compared to chromosomal DNA in humans) detectable genetic fingerprints can be targeted for epidemiological studies over short time periods Paraskevis et al. (2016). **HV: It's not immediately obvious to me why it matters that viruses are more amenable to molecular studies than bacteria, that RNA viruses have detectable genetic fingerprints, and how either of those facts relates to estimating distance thresholds.**

For chronic infections such as HIV, the most appropriate genetic distance threshold should be determined according to the characteristics of the epidemic such as the speed of transmission, and the evolutionary rate of the gene region analyzed Liu et al. (2020). Sampling density and possible delays between infection and diagnosis should be considered, since samples close to the time of seroconversion are more likely to cluster than samples from well after infection. Lower thresholds will capture the most closely related sequences, while higher thresholds will capture long-term epidemics and chronically infected individuals Junqueira et al. (2019).

Cluster analysis, i.e., identification and analysis of connected network components, in public health has been used for early identification of increased transmission Oster et al. (2021, 2018), monitoring response to an HIV outbreak Tumpney et al. (2020); Sizemore et al. (2020); Tookes et al. (2020), evaluating the effectiveness of interventions Peters et al. (2016); Wang et al. (2015); Liu et al. (2020) or predicting clusters that are most likely to grow in the near future Erly et al. (2021); Ragonnet-Cronin et al. (2022).

In general, the goal of informative network inference is to strike a balance between having enough links to accurately track the spread of the disease, while not having so many links that spurious connections are made and hinder the public health response. This can be achieved through careful analysis and consideration of the specific disease and context.

**HV: Add a final short paragraph here that conveys "We're addressing the problem of inferring the correct distance threshold with AutoTune. AutoTune is an improvement over other ways people guess the proper distance threshold for these reasons. AutoTune is a useful tool."**

## 2 METHODS

Assume that there are  $S$  aligned genomic sequences (full or partial, e.g. the HIV-1 pol gene) for a pathogen of interest, each representing the circulating viral diversity at the time of sampling in a single infected individual. We shall infer a putative transmission network comprising  $S$  nodes, and  $E$  links (edges), where an edge is drawn between a pair of sequences if the genetic distance between them is below a threshold  $d$ . In such a network, there will be  $1 \leq C \leq S$  connected components (clusters), which are the primary object of inference. This network inference strategy is used by HIV-TRACE **add reference**, where the genetic distance is computed using the Tamura-Nei (TN93) **add reference** approach, with a variety of options controlling how to deal with ambiguous nucleotide bases; for HIV-1 such bases are informative since they often represent variants co-circulating in the infected individual at the time of sampling.

We begin by describing an approach to assign a score to each of the choices of  $d$  in a plausible/informative range of distances. Note that while such a range is continuous, it is sufficient to only consider distance cutoffs that are in the array of pairwise distances between the sequences, as those are the cut-points where one or more additional edges will be added to the network as  $d$  is increased.

### 2.1 Scoring Heuristic Procedure

Network threshold selection procedure proceeds as follows:

1. For each candidate threshold  $d_L$ , in increasing order, ranging from the smallest genetic distance in the dataset, up to either the largest distance or a predetermined maximal threshold, we compute two network statistics:  $R_{12}$ , the ratio of the size of the largest cluster to the size of the second largest cluster, and  $C$  – the number of clusters in the network at this threshold.
2. A priority score is assigned to each  $d_L$ . This score measures two properties of the threshold: Does  $R_{12}$  jump at  $d_L$ ? How far is the number of clusters  $C$  at  $d_L$  from the maximal number of clusters over all threshold values? Let there be  $N$  overall  $d_L$  candidate values, and assume we are examining the  $i$ th candidate,  $d_L^i$  with  $W < i \leq N - W$  ( $W$  is a positive integer defined below).
  - a. The  $R_{12}$  jump is computed by looking at the normalized ratio of the mean  $R_{12}$  values computed over the leading window  $d_L^{i+1} \dots d_L^{i+W}$  and the trailing window  $d_L^{i-W} \dots d_L^{i-1}$ . The width of the window,  $W$ , is defined as  $((\lceil \frac{N}{100} \rceil, 3), 30)$ . **HV: I'm not entirely clear on how the width of the window is calculated. If N=50, then W becomes ((0.5, 3), 30). How do you get a single integer W from that?** The distribution of ratios is converted to  $Z$  scores, and normalized relative to the largest positive  $Z$  score across all candidate distances, yielding the jump component of the score.
  - b. The number of clusters,  $C_i$  at threshold  $d_L^i$  is first normalized to  $[0, 1]$  through  $\frac{C_{max} - C_i}{C_{max} - C_{min}}$  and next gated via a Gompertz function transform  $1 - e^{-e^{-25x+3}}$ . This function provides an *ad hoc* means for penalizing having too few clusters relative to the maximum over all ranges. For example, a

threshold that yields 95% of the maximal number of clusters receives a score of 0.996, while a threshold that yields 85% - a score of 0.376.

c. The priority score for  $d_L^i$  is the sum of the two components defined in (a) and (b).

3. The threshold with the highest priority score will be selected as the suggested automatic distance threshold, if the score is high enough (1.9 or more), and either of the two conditions hold.

a. No other thresholds have priority scores of 1.9 or higher

b. If other thresholds have priority scores of 1.9 or higher, then the range of thresholds represented by these options is small (no more than  $\log N$  times the mean step between successive  $d_L^i$ ).

4. If no single threshold can be selected in step 3, then the one with the highest priority score is suggested, and an inspection of the plot like the one on the analyze page is recommended to ensure that the threshold is sensible. **HV: What's the analyze page? Define it here, or mention that it's defined below.**

## 2.2 Assortativity

Degree-weighted homophily (DWH) is a measure of similarity between nodes in a network based on their attributes (such as demographic characteristics or behaviors) and their degree (i.e., the number of connections they have to other nodes in the network). It is used to quantify the extent to which nodes with similar attributes tend to be connected to each other more frequently than would be expected by chance **add citation. HV: I like this explanation. Simple, clear, easy to see the biological/public health relevance.** DWH is calculated as the ratio of the observed number of connections between nodes with similar attributes to the expected number of connections between such nodes, based on their network degree.

In mathematical terms, it is defined as:

$$DWH = \frac{W_M + W_C - 2W_X}{\frac{d_{in}^2}{nodes_{in}^2} + \frac{d_{out}^2}{nodes_{out}^2}} \quad (1)$$

Where

- $W_M$  : Weight of in-group connections
- $W_C$  : Weight of out-group connections
- $W_X$  : Weight of cross-group connections
- $d_{in}$  : In-group degree
- $d_{out}$  : Out-group degree
- $nodes_{in}$  : number of in-group nodes
- $nodes_{out}$  : number of out-group nodes

DWH ranges from  $-1$  to  $1$ . A DWH value of  $0$  indicates that there is no more homophily than expected with chance, while a value of  $1$  indicates that there is perfect homophily (e.g. birds always link to birds, and only birds). A value of  $-1$  is achieved for perfectly disassortative networks (e.g. a bird never linking with another bird).

DWH has been used extensively in social network analysis and in the study of how different attributes are related to the formation of connections between individuals. It is used as a way to measure the similarity of attributes between individuals in a network. Additionally, randomization is performed by shuffling

attribute labels among nodes, then performing DWH computation. This is useful in creating a null distribution of DWH scores under random mixing. A panmictic range is reported by shuffling attributes multiple times and reporting the minimum and maximum score.

## 2.3 Implementation

The software implementation involves a step-by-step process that utilizes the HIV-TRACE suite of packages. It starts with calculating pairwise distances with the `tn93` tool and a supplied multiple sequence alignment. Thus generated pairwise distances are supplied to the `hivnetworkcsv` script while providing the `-A` keyword argument. A brief outline of the software's implementation is as follows

1. Calculate pairwise distances: The user first calculates the pairwise distances using the `tn93` fast pairwise distance calculator, providing the maximum threshold value to consider (0.03 in this case) and the input FASTA file. The command for this step is

```
1 tn93 -t 0.030 pol.fasta > pairwise_distances.15.tn93.csv
```

Please note that the threshold should include the maximal range one is intending to test.

2. Compute distance threshold scores: The `hivnetworkcsv` script is then executed with the required input file, format, and autotune option to generate a tab-separated output file, as shown below

```
1 hivnetworkcsv -i pairwise_distances.15.tn93.csv -f plain -A 0 > autotune_report.tsv
```

3. Visualize the report: Users can upload the generated `autotune_report.tsv` file to <http://autotune.datamonkey.org/analyze> for visualization and further analysis of the data. This web-based site extends the Datamonkey platform Weaver et al. (2018) to provide an interactive environment to explore scores and other metrics across the range of tested outputs.

4. Run HIV-TRACE: Once AUTO-TUNED threshold(s) are settled upon after review, the user runs the HIV-TRACE command with the appropriate input FASTA file, distance threshold, and other required arguments. The output is saved as a JSON file. An example command is

```
1 hivtrace -i ./INPUT.FASTA -a resolve -r HXB2_prnt -t < autotune_threshold > -m 500 -g .05
  ↪> hivtrace.results.json
```

|||||| HEAD

### 2.3.1 Optional : Compute Assortativity Metrics

5. Annotate results: The `hivnetworkannotate` script is used to annotate the results obtained from the HIV-TRACE step with attributes. The script takes the JSON results file, node attributes file, schema file, and a resolve flag as input.

```
1 hivnetworkannotate -n hivtrace.results.json -a node_attributes.json -g schema.json -r
```

For more information, users can refer to the `hivnetworkannotate` documentation.

6. Analyze the results with DWH: After the results file has been annotated, the user can proceed to the assortativity page, <http://autotune.datamonkey.org/assortativity>, for further analysis of the output.

AUTO-TUNE is readily accessible on GitHub as part of the `hivclustering` repository (<https://github.com/veg/hivclustering>). It is integrated into the command-line interface of the software as the `-A` or `--auto-profile` argument. `hivclustering` is a key

component of the HIV-TRACE suite of tools, a resource for the inference, analysis, and visualization of HIV transmission networks.

The Degree Weighted Homophily (DWH) calculation tool, an integral component of the assortativity step, is developed using TypeScript, a statically typed superset of JavaScript that ensures robustness and scalability. In an effort to promote accessibility and ease of integration, the DWH tool is packaged and distributed through the Node Package Manager (npm), enabling researchers and developers to conveniently incorporate this advanced analytical tool into their own projects and workflows. DWH can be used in-browser or as a command-line tool, allowing researchers and developers to employ the tool in an interactive command-line interface or integrate it into larger software applications, thus catering to a diverse array of technical needs and preferences. Instructions for usage and installation are found on Github (<https://github.com/veg/dwh>).

The described workflow offers a systematic approach to analyze potential distance thresholds for one's data with AUTO-TUNE, from calculating pairwise distances to visualizing and annotating results.

## 2.4 Visualization

Visualizations of AUTO-TUNE results are accessible at <http://autotune.datamonkey.org/analyze>. It is a dynamic and interactive web-based platform that offers visualization and analysis of results generated by AUTO-TUNE. The website provides a comprehensive view of the data by generating various plots across candidate distance thresholds. These include a score plot, allowing users to identify trends and anomalies across the full range of thresholds. Additionally, it generates a graph showing the number of clusters across candidate thresholds, one of the components that contribute to the score. The site also includes an R1/R2 plot that displays the ratio of the largest cluster to the second largest cluster across candidate thresholds, which is the other metric that contributes to the scoring heuristic. **this could benefit from referencing a figure showing such plots**

An assortativity tool is available at <http://autotune.datamonkey.org/assortativity>, and is an advanced analytical tool engineered to facilitate the calculation of Degree Weighted Homophily (DWH) values. It utilizes the DWH NPM package to generate a tabular representation of DWH values corresponding to each value for a selected attribute annotation, providing an exhaustive examination of the interrelationships for the field. A notable feature is the computation of the panmictic range, which involves a label permutation test to generate the null distribution of DWH values. This feature establishes a comparative baseline that aids in determining the significance of homophily versus what would be expected by chance. Lastly, the site also provides a plot of the fraction of pairwise connections, normalized by degree, for each value pertinent to the selected field. This visual depiction facilitates an intuitive comprehension of the distribution and interconnections within the dataset.

The site aims to offer a user-friendly interface for data visualization, playing an important role in interpreting and understanding AUTO-TUNE's output data. The visualization code is available on Github (<https://github.com/stevenweaver/autotune-app/>).

## 2.5 Comparisons with previously published analyses

In conducting our comparisons with the established `clustuneR` method **citation**, we procured datasets from Wolf et al. (2017) and Vrancken et al. (2017) utilizing the identical approach delineated in Chato et al. (2020). These datasets, namely Middle Tennessee, Seattle, and Alberta, were processed using the workflow prescribed in Section 2.3. This enabled us to determine an optimal threshold for



each dataset using AUTO-TUNE. We further executed the command as detailed in step 4 of Section 2.3, deploying thresholds previously established as optimal by Chato et al. (2020).

To perform comparisons, we computed the average degree-weighted homophily score over a set of three-year sliding windows. Specifically, the homophily among nodes was calculated for a collection of date ranges as follows:

$$\bar{H} = \frac{1}{N} \sum_{i=1}^N H(w_i) \quad (2)$$

where  $\bar{H}$  represents the average degree-weighted homophily score,  $N$  is the total number of sliding windows,  $H(w_i)$  is the homophily score for the  $i$ -th window, and the windows  $w_i$  correspond to the date ranges, e.g., '2012-2015', '2013-2016', '2014-2017', etc. This methodology allowed us to compare the "best thresholds" derived from our proposed AUTO-TUNE method against those defined as optimal in Chato et al. (2020).

Second, we set out to compare the thresholds obtained in numerous published investigations with those obtained by AUTO-TUNE. To select the data sets for this analysis, we conducted a scientific literature search to identify studies focused on HIV networks for public health purposes. We then filtered the studies that utilized HIV-TRACE to infer genetic networks and had publicly available sequences. Thus, we attempted to include studies from different countries and regions, enabling us to assess the performance of our method across various epidemic contexts, risk groups, and network sizes in real-data sets that used variable clustering thresholds.

In order to evaluate the influence of sampling density on the genetic distance threshold as determined by AUTO-TUNE, we implemented a strategy of random subsampling from the original dataset sourced from Rhee et al. (2019). This study was selected due to its satisfactory AUTO-TUNE score when utilized in its entirety, as well as its inherent design as a Geographically-Stratified set of 716 Pol Subtype/CRF (GSPS) reference sequence dataset. The dataset, which comprises 6034 samples gathered between 1959 and 2016, was subjected to random subsampling ten times at proportions of 25%, 50%, and 75% of the original sample size. For each subsample, the optimal threshold and associated scores were determined via AUTO-TUNE.

## 3 RESULTS

### 3.1 Comparison with clustuneR

We compared results to `clustuneR`, which employs the recency of sample collection or diagnosis as individual-level weights in a predictive model to estimate the growth of HIV clusters. The thresholds deemed optimal by `clustuneR` were found by finding the minimum GAIC (generalized AIC) across candidate distances between 0 and 0.04 in steps of  $8 \times 10^{-4}$ . GAIC is the difference between a null model that is only influenced by cluster size, and a weighted model that includes individual-level attributes among known cases in the cluster. Using the minimum GAIC metric, it was found that 0.016 was the optimal threshold for Tennessee and Seattle, and 0.0104 for Northern Alberta.

In contrast, AUTO-TUNE does not incorporate any attribute data in its scoring heuristic. Instead, it relies on clustering metrics constructed purely from pairwise distances between sequences. Using the same datasets analyzed by `clustuneR` Chato et al. (2020), AUTO-TUNE found the thresholds with the

highest scores to be 0.01431 for Middle Tennessee, 0.01354 for Seattle, and 0.01099 for Northern Alberta. Table 1. These values are in broad agreement with the original results.

discuss how our thresholds perform vs the published thresholds; look at the GAIC plots

Because there is no "ground truth" of what constitutes the best threshold, we chose to use assortativity as a proxy measure. this may need more justification

Performance of the inferred optimal thresholds were performed using an average degree-weighted homophily (DWH) score across 3-year collection date windows starting from the oldest collection year for each respective dataset, as that is the metadata that was consistently available across all three datasets and was the attribute of focus used by `clustuneR`. DWH in this case measures the affinity for nodes within the network to link with other nodes in the same collection date window. For example, samples collected between 2012-2015 linking more often with other samples within the same time window would result in a higher DWH score. It was found that, across all three datasets, using the threshold with the highest score reported by AUTO-TUNE resulted in a higher average DWH score across all three datasets. compared to what?

When reviewing scores across all candidate thresholds with AUTO-TUNE, none of the three datasets reached one confident score over any other. The most confident score was achieved by the Seattle dataset at 1.53, then Tennessee with a high score of 1.26, and lastly Canada at 1.02. When finding peaks, a second peak in Seattle denotes that 0.01166 may also be an optimal threshold to consider, as its score, 1.52976, is only 0.003 less than the highest score. The optimal score for Tennessee and Canada are a bit more dubious, as there are multiple peaks within close scores of each other. Indeed, after applying a 0.75 minimum score threshold after visual inspection for peak ranges, standard deviation among scores were 0.0475, 0.0089, and 0.0084 for Seattle, northern Alberta, and Tennessee, respectively. This implies there may be multiple thresholds that would be considered reasonable, and downstream homophily metrics with attributes may aid in coming to a decision.

### 3.2 Comparison with Prior Publications Citing HIV-TRACE

Next, we curated publications citing HIV-TRACE that also had publicly accessible data associated with the study Rhee et al. (2019); Brenner et al. (2021); H et al. (2021); Liu et al. (2020); Bbosa et al. (2020); Yan et al. (2020); Dalai et al. (2018); Sivay et al. (2018). We found that a variety of different ways has been used to determine distance thresholds: using the precedent set by the CDC for detecting recent and rapid clusters Yan et al. (2020); using thresholds from other studies Sivay et al. (2018); visual inspection of the number of clusters and nodes across candidate distance thresholds Liu et al. (2020). Because these thresholds were determined largely qualitatively, they tended to be round numbers and only somewhat tuned to their respective geographic region of research. Table 2 When AUTO-TUNE was applied to these datasets, a range of dataset-specific thresholds was inferred. This analysis demonstrated that network inference can be highly sensitive to relatively small changes in the distance threshold. For example, when reviewing the distribution of scores for the dataset used by Dalai et al. (2018), the score is exactly 2 at 0.01848, but at thresholds tested at just 0.00002 difference, scores drop precipitously to 1.638 and 0.826 for candidates 0.01846 and 0.0185, respectively. Another example of a seemingly close threshold yet perhaps not optimal is found with Bbosa et al. (2020). While no score across candidate thresholds was found to be above 1.9, a high score was found at distance 1.707, with 1.2415. Contrast this with the threshold used, 1.5%, with a score of 0.0124. I would add some specifics here: what is  $R_{12}$  and  $C$  for those thresholds, for example? The composite score is difficult to appreciate intuitively



### 3.3 Evaluating Performance using DWH

need to better articulate WHY we think that higher DWH is \*desirable\*

To assess the performance of an AUTO-TUNED optimized threshold using degree-weighted homophily, we first evaluated a CRF07\_BC network with data from China. We used 8178 HIV-1 CRF07\_BC pol sequences collected by a national survey in China to construct longitudinal transmission networks, each pol sequence was annotated with risk factor detailing whether the patient was heterosexual (Hetero), Person With Injected Drug Use (PWID), or Men who have Sex with Men (MSM), among other attributes. Using AUTO-TUNE, no distance threshold receives a score above 1.9, but using the default 1.5% threshold is clearly suboptimal. Using a 1.5% threshold, the network captures 5923 nodes, of which 559 are PWID, 3371 MSM, 1993 Hetero, and has an AUTO-TUNE score of 0.029. When evaluating DWH among risk factors, MSM, Hetero, and PWID had scores of 0.211, 0.133, and 0.168, respectively.

When using AUTO-TUNE, two separate ranges appear. The highest score (1.1369) is obtained at a distance threshold 0.76%. The second-highest score is 1.0303 at distance threshold 0.0019. Networks at both thresholds were also evaluated with DWH based on risk factors. The network at 0.76% captured 3537 nodes, of which 236 are PWID, 2271 MSM, 1030 Hetero. When evaluating DWH among risk factors, MSM and Hetero both had slightly increased scores of 0.237 and 0.185, respectively. PWID DWH dramatically increased to 0.401. The network at 0.19% captures 1654 nodes, of which 151 are PWID, 1075 MSM, 428 Hetero. When evaluating DWH among risk factors, MSM, Hetero, and PWID had slightly increased scores over 0.76% of 0.292, 0.25, and 0.445, respectively.

The data in the above 2 paragraphs need to be in a table; the text should interpret it. You should also mention whether or not DWH estimates were significantly different from 0

We next evaluated Rhee et al. (2019) with DWH. The dataset received a clear optimal threshold of 0.01699 with an AUTO-TUNE score of 1.9998. At the default threshold of 1.5%, the AUTO-TUNE score was 0.9782. At threshold 1.5%, the network captured 1351 nodes, compared with 1592 nodes out of 6034 captured at threshold 1.699%. When evaluating DWH with country, substantial change were only found for China and Thailand, improving from no better than random linkage at threshold 1.5% with DWH  $-0.166$  and  $-0.051$ , to  $0.116$  and  $0.132$  at threshold 1.699%, respectively. Notably, no country scored worse with the optimal AUTO-TUNE threshold, despite it being larger than the default 1.5%.

Same here

### 3.4 The Effect of Subsampling on Optimal Thresholds and AUTO-TUNE Scores

Also should include some motivation

Next, we evaluated the performance of AUTO-TUNE when subsampling a dataset. Since the Rhee et al. (2019) dataset exhibited a clear optimal peak, we used the dataset for analysis, and randomly sampled 10 times from the entire dataset at 25%, 50%, and 75% each. The original full dataset confidently determined 0.01699 (AUTO-TUNE score 1.9998).

Sampling at 25% yielded a mean top threshold of 0.021509, median at 0.019765, and standard deviation of 0.004388. 50% yielded 0.018581 and 0.01871 mean and median, respectively with a standard deviation of 0.001629. Finally, 75% calculated mean is approximately 0.017403, with a median of approximately 0.01699. The standard deviation was 0.000924.

explain why the trend of HIGHER thresholds for the SPARSER sampling makes sense

As the proportion increased from 25% to 50% and 75%, observable shifts were also noted in the mean, median, and standard deviation of the AUTO-TUNE scores. At 25%, the mean and median scores were 1.5585 and 1.5014 respectively, with a standard deviation of 0.3568. At 50%, both mean and median scores significantly increased to 1.8171 and 1.9191 respectively, and the standard deviation dropped to 0.2482. Upon reaching an AUTO-TUNE of 75%, the mean and median scores rose further to 1.9870 and 1.9997 respectively, while the standard deviation shrank substantially to 0.0364, indicating higher consistency in scores.

As the sample proportion increased, an upward trend was noted in average AUTO-TUNE scores. Additionally, the standard deviation reduced significantly with sample proportion. This implies that as sampling becomes denser, AUTO-TUNE will become more confident in determining the optimal threshold for a particular dataset.

I still think that being able to show what fraction of the nodes connected in the full network are connected in the subsampled networks; if you have network JSONs, I can check that pretty quickly

## 4 DISCUSSION

AUTO-TUNE addresses the challenge of selecting setting an appropriate genetic distance threshold in HIV transmission network analysis by implementing a heuristic scoring system that is based on two desirable properties of networks generated by candidate genetic distance thresholds: a large number of clusters, and the absence of a giant component. Too few small clusters are a hallmark of an excessively low threshold, and a giant cluster that includes too many sequences is a hallmark of an excessively high threshold. In the application of AUTO-TUNE to various datasets, the results demonstrated its efficacy across different epidemic contexts, risk groups, and network sizes. Not sure what efficacy means AUTO-TUNE consistently selected thresholds that were comparable comparable how?, if not better better how?, to those manually chosen in prior studies using the same data, illustrating the value of a more systematic, automated, and data-adaptive approach.

For example, the results of our study suggest that AUTO-TUNE, which relies solely on clustering metrics from pairwise distances, could be an effective alternative to other distance-based methods, such as `clustuneR` while less time-consuming and possessing a gentle learning curve, which makes it easy to use by personnel not specialized in bioinformatics and computer science. Furthermore, the simplicity of the method represents an advantage over phylogenetic methods where, in addition to the calculation of genetic distances, it must also determine a support/distance threshold where a rationale for the selection of these thresholds is rarely provided Junqueira et al. (2019).

AUTO-TUNE generated thresholds for all three examined datasets (Middle Tennessee, Seattle, and Northern Alberta) that outperformed `clustuneR` using DWH on 3-year collection date windows across all three datasets again, need to be clear about why higher DWH is better. This indicates that even without incorporating attribute data, AUTO-TUNE's scoring heuristic could provide reliable thresholds for HIV clusters. However, for the determination of the optimal genetic distance threshold, time-related and context-specific factors might need to be considered if there is no significant score for any one candidate threshold, especially if there are multiple peaks. For example, during HIV outbreaks in injection drug users (that usually occur over several months), it may be more appropriate to use the shorter genetic distance threshold Peters et al. (2016); Campbell et al. (2017) between multiple high-scoring thresholds. On the contrary, larger and more extended epidemics over time exhibit a tendency toward larger genetic

distance thresholds in order to capture transmission than younger epidemics and less densely sampled epidemic investigations Patil et al. (2022); Leung et al. (2019); Di Giallonardo et al. (2021).

Our evaluation of publications citing HIV-TRACE revealed the largely qualitative determination of distance thresholds. This approach may result in less accurate or suboptimal thresholds due to a lack of systematic analysis. In contrast, AUTO-TUNE offers a more systematic and granular approach to threshold selection, with our findings demonstrating that even minor adjustments to the distance can drastically change the score. Therefore, using AUTO-TUNE could potentially improve the quality of HIV clustering and transmission network studies.

The Degree-Weighted Homophily (DWH) evaluation showed that AUTO-TUNE could improve network quality based on specific attributes, such as risk factor, which is an important part of HIV studies and informing prevention measures Potterat et al. (2002); Fujimoto et al. (2021). For example, the use of AUTO-TUNE resulted in an increased DWH among the MSM, Hetero, and PWID groups when analyzing a CRF07\_BC network. Additionally, the results from the Rhee et al. dataset also demonstrated AUTO-TUNE's ability to improve DWH geographically, enhancing the network's ability to accurately reflect transmission dynamics.

Our analysis of AUTO-TUNE's performance on subsamples of a dataset revealed its sensitivity to sample size. The results indicated a correlation between increased sample size and higher average AUTO-TUNE scores, as well as lower score variability. This suggests that denser sampling could enhance AUTO-TUNE's ability to determine the optimal threshold for a dataset. Further studies might be needed to establish the minimum sample size required for reliable threshold determination.

#### 4.1 When a Score is Below 1.9

add some text to explain that multiple scores at different thresholds could be indicative of inherently different scales in the network, e.g. global AND local combined into one

The use of AUTO-TUNE, while offering a method for automated threshold selection, may not always provide a single, decisive score that unambiguously determines the optimal threshold. In certain situations, several candidate thresholds may yield similar AUTO-TUNE scores, making it difficult to single out one as the clear-cut 'optimal' threshold. In these scenarios, the process of threshold selection becomes more nuanced and requires a deeper analysis. The plot of AUTO-TUNE scores across candidate thresholds can serve as a valuable tool in these cases. For instance, researchers could identify a range of thresholds that all produce similar scores, suggesting that the specific choice of threshold within this range may not significantly impact the resulting network. Moreover, combining AUTO-TUNE with the DWH measure can enhance the interpretation of such plots. By considering how assortativity changes across the range of candidates, researchers can make more informed decisions about the appropriate choice. If there is a certain threshold at which the DWH measure noticeably changes for an attribute of interest, this could suggest a meaningful shift in the network structure that would be worth considering when selecting a threshold. The symbiotic approach of combining AUTO-TUNE scores, DWH measure, and visual analysis of score plots provides a more nuanced method for threshold selection when no clear optimal threshold emerges from the AUTO-TUNE scores alone.

The AUTO-TUNE methodology has several limitations. First, even though it provides the advantage of operating without the need for metadata, the size and the subgenomic region analyzed may affect the accuracy of transmission inference Junqueira et al. (2019). Second, our analysis of AUTO-TUNE's performance on subsamples of a dataset revealed its sensitivity to sample size, as the performance of

the method can be affected by sampling density, improving the reliability of the test as the sampling density increases (figure X). However, our results were consistent with previous studies, which have suggested an optimal sampling density of 50–70% for HIV-1 cluster analysis (Novitsky et al. (2014)). Third, even when it provides an insight of the optimal threshold to analyze a network, the supplied information might still need validation by experts, especially when no clear threshold is identified. In this case, it has been recommended to combine genetic data with clinical and sociodemographic information for a better characterization of the network structure. Finally, the performance of the method needs to be assessed in pathogens different from HIV, leading to opportunities for future research.

## 5 CONCLUSION

AUTO-TUNE operates solely utilizing genetic sequence data to ascertain a decisive threshold. It employs a scoring heuristic, which is based on the number of clusters produced by a pairwise distance threshold and the ratio of the largest cluster to the second largest across a range of possible thresholds using sliding windows.

A key advantage of this approach is its autonomy from supplementary data. When a patient tests positive for HIV, data collection protocols can greatly vary, and additional data are not always available or consistent. However, by leveraging only genetic sequence data, AUTO-TUNE eliminates the need for such information.

Consequently, AUTO-TUNE's performance is consistently controlled, irrespective of the fluctuations seen in data collection protocols after a positive HIV diagnosis. This level of adaptability demonstrates its suitability for integration into various contexts related to HIV, and possibly other viral cluster detection and response protocols. This versatility underscores the strong methodological foundation of AUTO-TUNE and its potential utility.

## CONFLICT OF INTEREST STATEMENT

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

## AUTHOR CONTRIBUTIONS

The Author Contributions section is mandatory for all articles, including articles by sole authors. If an appropriate statement is not provided on submission, a standard one will be inserted during the production process. The Author Contributions statement must describe the contributions of individual authors referred to by their initials and, in doing so, all authors agree to be accountable for the content of the work. Please see here for full authorship criteria.

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## SUPPLEMENTAL DATA

Supplementary Material should be uploaded separately on submission, if there are Supplementary Figures, please include the caption in the same file as the figure. LaTeX Supplementary Material templates can be found in the Frontiers LaTeX folder.

## DATA AVAILABILITY STATEMENT

Data are available at GenBank accession numbers JX160108-JX161480, JX498971-JX498972, JX498976-JX498990, JX498992-JX499018, KU190031-KU190839, KY34691-KY37792, KY883695-KY883762, KY888784-KY888875, KY921717-KY921757, MG434786-MG435347, MG435358-MG436769, MH352627-MH355541, MK25548, MN424584-MN427369, MT336755-MT336776, MT368043-MT369927.

## REFERENCES

- Bbosa, N., Ssemwanga, D., and Kaleebu, P. (2020). Short Communication: Choosing the Right Program for the Identification of HIV-1 Transmission Networks from Nucleotide Sequences Sampled from Different Populations. *AIDS research and human retroviruses* 36, 948–951. doi:10.1089/AID.2020.0033
- Brenner, B. G., Ibanescu, R.-I., Osman, N., Cuadra-Foy, E., Oliveira, M., Chaillon, A., et al. (2021). The Role of Phylogenetics in Unravelling Patterns of HIV Transmission towards Epidemic Control: The Quebec Experience (2002-2020). *Viruses* 13, 1643. doi:10.3390/v13081643
- Campbell, E. M., Jia, H., Shankar, A., Hanson, D., Luo, W., Masciotra, S., et al. (2017). Detailed Transmission Network Analysis of a Large Opiate-Driven Outbreak of HIV Infection in the United States. *The Journal of Infectious Diseases* 216, 1053–1062. doi:10.1093/infdis/jix307
- Chato, C., Kalish, M. L., and Poon, A. F. Y. (2020). Public health in genetic spaces: a statistical framework to optimize cluster-based outbreak detection. *Virus Evolution* 6, veaa011. doi:10.1093/ve/veaa011
- Dalai, S. C., Junqueira, D. M., Wilkinson, E., Mehra, R., Kosakovsky Pond, S. L., Levy, V., et al. (2018). Combining Phylogenetic and Network Approaches to Identify HIV-1 Transmission Links in San Mateo County, California. *Frontiers in Microbiology* 9, 2799. doi:10.3389/fmicb.2018.02799
- Di Giallonardo, F., Pinto, A. N., Keen, P., Shaik, A., Carrera, A., Salem, H., et al. (2021). Subtype-specific differences in transmission cluster dynamics of HIV-1 B and CRF01\_ae in New South Wales, Australia. *Journal of the International AIDS Society* 24, e25655. doi:10.1002/jia2.25655
- Ding, X., Chaillon, A., Pan, X., Zhang, J., Zhong, P., He, L., et al. (2022). Characterizing genetic transmission networks among newly diagnosed HIV-1 infected individuals in eastern China: 2012–2016. *PLOS ONE* 17, e0269973. doi:10.1371/journal.pone.0269973. Publisher: Public Library of Science
- Erly, S. J., Naismith, K., Kerani, R., Buskin, S. E., and Reuer, J. R. (2021). Predictive Value of Time-Space Clusters for HIV Transmission in Washington State, 2017-2019. *Journal of Acquired Immune Deficiency Syndromes (1999)* 87, 912–917. doi:10.1097/QAI.0000000000002675
- Fujimoto, K., Bahl, J., Wertheim, J. O., Del Vecchio, N., Hicks, J. T., Damodaran, L., et al. (2021). Methodological synthesis of Bayesian phylodynamics, HIV-TRACE, and GEE: HIV-1 transmission



- epidemiology in a racially/ethnically diverse Southern U.S. context. *Scientific Reports* 11, 3325. doi:10.1038/s41598-021-82673-8. Number: 1 Publisher: Nature Publishing Group
- Gore, D. J., Schueler, K., Ramani, S., Uvin, A., Phillips, G., McNulty, M., et al. (2022). HIV Response Interventions that Integrate HIV Molecular Cluster and Social Network Analysis: A Systematic Review. *AIDS and behavior* 26, 1750–1792. doi:10.1007/s10461-021-03525-0
- H, Y., H, W., Y, X., L, H., Y, L., Q, L., et al. (2021). Acquisition and transmission of HIV-1 among migrants and Chinese in Guangzhou, China from 2008 to 2012: Phylogenetic analysis of surveillance data. *Infection, genetics and evolution : journal of molecular epidemiology and evolutionary genetics in infectious diseases* 92. doi:10.1016/j.meegid.2021.104870. Publisher: Infect Genet Evol
- Holmes, E. C., Zhang, L. Q., Robertson, P., Cleland, A., Harvey, E., Simmonds, P., et al. (1995). The molecular epidemiology of human immunodeficiency virus type 1 in Edinburgh. *The Journal of Infectious Diseases* 171, 45–53. doi:10.1093/infdis/171.1.45
- Junqueira, D. M., Sibisi, Z., Wilkinson, E., and de Oliveira, T. (2019). Factors influencing HIV-1 phylogenetic clustering. *Current opinion in HIV and AIDS* 14, 161–172. doi:10.1097/COH.0000000000000540
- Kosakovsky Pond, S. L., Weaver, S., Leigh Brown, A. J., and Wertheim, J. O. (2018). HIV-TRACE (TRANsmiSSion Cluster Engine): a Tool for Large Scale Molecular Epidemiology of HIV-1 and Other Rapidly Evolving Pathogens. *Molecular Biology and Evolution* 35, 1812–1819. doi:10.1093/molbev/msy016
- Leung, K. S.-S., To, S. W.-C., Chen, J. H.-K., Siu, G. K.-H., Chan, K. C.-W., and Yam, W.-C. (2019). Molecular Characterization of HIV-1 Minority Subtypes in Hong Kong: A Recent Epidemic of CRF07\_bc among the Men who have Sex with Men Population. *Current HIV research* 17, 53–64. doi:10.2174/1570162X17666190530081355
- Liu, M., Han, X., Zhao, B., An, M., He, W., Wang, Z., et al. (2020). Dynamics of HIV-1 Molecular Networks Reveal Effective Control of Large Transmission Clusters in an Area Affected by an Epidemic of Multiple HIV Subtypes. *Frontiers in Microbiology* 11, 604993. doi:10.3389/fmicb.2020.604993
- Novitsky, V., Moyo, S., Lei, Q., DeGruttola, V., and Essex, M. (2014). Impact of sampling density on the extent of HIV clustering. *AIDS research and human retroviruses* 30, 1226–1235. doi:10.1089/aid.2014.0173
- Oster, A. M., France, A. M., Panneer, N., Bañez Ocfemia, M. C., Campbell, E., Dasgupta, S., et al. (2018). Identifying Clusters of Recent and Rapid HIV Transmission Through Analysis of Molecular Surveillance Data. *Journal of Acquired Immune Deficiency Syndromes (1999)* 79, 543–550. doi:10.1097/QAI.0000000000001856
- Oster, A. M., Lyss, S. B., McClung, R. P., Watson, M., Panneer, N., Hernandez, A. L., et al. (2021). HIV Cluster and Outbreak Detection and Response: The Science and Experience. *American Journal of Preventive Medicine* 61, S130–S142. doi:10.1016/j.amepre.2021.05.029
- Paraskevis, D., Nikolopoulos, G. K., Magiorkinis, G., Hodges-Mameletzis, I., and Hatzakis, A. (2016). The application of HIV molecular epidemiology to public health. *Infection, Genetics and Evolution: Journal of Molecular Epidemiology and Evolutionary Genetics in Infectious Diseases* 46, 159–168. doi:10.1016/j.meegid.2016.06.021
- Patil, A., Patil, S., Rao, A., Gadhe, S., Kurle, S., and Panda, S. (2022). Exploring the Evolutionary History and Phylodynamics of Human Immunodeficiency Virus Type 1 Outbreak From Unnao, India Using Phylogenetic Approach. *Frontiers in Microbiology* 13, 848250. doi:10.3389/fmicb.2022.848250



- Peters, P. J., Pontones, P., Hoover, K. W., Patel, M. R., Galang, R. R., Shields, J., et al. (2016). HIV Infection Linked to Injection Use of Oxymorphone in Indiana, 2014–2015. *The New England Journal of Medicine* 375, 229–239. doi:10.1056/NEJMoa1515195
- Potterat, J. J., Phillips-Plummer, L., Muth, S. Q., Rothenberg, R. B., Woodhouse, D. E., Maldonado-Long, T. S., et al. (2002). Risk network structure in the early epidemic phase of HIV transmission in Colorado Springs. *Sexually Transmitted Infections* 78, i159–i163. doi:10.1136/sti.78.suppl\_1.i159. Publisher: The Medical Society for the Study of Venereal Disease Section: Symposium
- Ragonnet-Cronin, M., Hayford, C., D'Aquila, R., Ma, F., Ward, C., Benbow, N., et al. (2022). Forecasting HIV-1 Genetic Cluster Growth in Illinois, United States. *Journal of Acquired Immune Deficiency Syndromes (1999)* 89, 49–55. doi:10.1097/QAI.0000000000002821
- Rhee, S.-Y., Magalis, B. R., Hurley, L., Silverberg, M. J., Marcus, J. L., Slome, S., et al. (2019). National and International Dimensions of Human Immunodeficiency Virus-1 Sequence Clusters in a Northern California Clinical Cohort. *Open Forum Infectious Diseases* 6, ofz135. doi:10.1093/ofid/ofz135
- Rose, R., Cross, S., Lamers, S. L., Astemborski, J., Kirk, G. D., Mehta, S. H., et al. (2020). Persistence of HIV transmission clusters among people who inject drugs. *AIDS (London, England)* 34, 2037–2044. doi:10.1097/QAD.0000000000002662
- Sivay, M. V., Hudelson, S. E., Wang, J., Agyei, Y., Hamilton, E. L., Selin, A., et al. (2018). HIV-1 diversity among young women in rural South Africa: HPTN 068. *PloS One* 13, e0198999. doi:10.1371/journal.pone.0198999
- Sizemore, L., Fill, M.-M., Mathieson, S. A., Black, J., Brantley, M., Cooper, K., et al. (2020). Using an Established Outbreak Response Plan and Molecular Epidemiology Methods in an HIV Transmission Cluster Investigation, Tennessee, January–June 2017. *Public Health Reports (Washington, D.C.: 1974)* 135, 329–333. doi:10.1177/0033354920915445
- Tookes, H., Bartholomew, T. S., Geary, S., Matthias, J., Poschman, K., Blackmore, C., et al. (2020). Rapid Identification and Investigation of an HIV Risk Network Among People Who Inject Drugs -Miami, FL, 2018. *AIDS and behavior* 24, 246–256. doi:10.1007/s10461-019-02680-9
- Tumpney, M., John, B., Panneer, N., McClung, R. P., Campbell, E. M., Roosevelt, K., et al. (2020). Human Immunodeficiency Virus (HIV) Outbreak Investigation Among Persons Who Inject Drugs in Massachusetts Enhanced by HIV Sequence Data. *The Journal of Infectious Diseases* 222, S259–S267. doi:10.1093/infdis/jiaa053
- Vrancken, B., Adachi, D., Benedet, M., Singh, A., Read, R., Shafran, S., et al. (2017). The multi-faceted dynamics of HIV-1 transmission in Northern Alberta: A combined analysis of virus genetic and public health data. *Infection, Genetics and Evolution: Journal of Molecular Epidemiology and Evolutionary Genetics in Infectious Diseases* 52, 100–105. doi:10.1016/j.meegid.2017.04.005
- Wang, X., Wu, Y., Mao, L., Xia, W., Zhang, W., Dai, L., et al. (2015). Targeting HIV Prevention Based on Molecular Epidemiology Among Deeply Sampled Subnetworks of Men Who Have Sex With Men. *Clinical Infectious Diseases: An Official Publication of the Infectious Diseases Society of America* 61, 1462–1468. doi:10.1093/cid/civ526
- Weaver, S., Shank, S. D., Spielman, S. J., Li, M., Muse, S. V., and Kosakovsky Pond, S. L. (2018). Datamonkey 2.0: A Modern Web Application for Characterizing Selective and Other Evolutionary Processes. *Molecular Biology and Evolution* 35, 773–777. doi:10.1093/molbev/msx335
- Wolf, E., Herbeck, J. T., Van Rompaey, S., Kitahata, M., Thomas, K., Pepper, G., et al. (2017). Short Communication: Phylogenetic Evidence of HIV-1 Transmission Between Adult and Adolescent Men Who Have Sex with Men. *AIDS research and human retroviruses* 33, 318–322. doi:10.1089/AID.2016.0061

- 580 Yan, H., He, W., Huang, L., Wu, H., Liang, Y., Li, Q., et al. (2020). The Central Role of Nondisclosed Men  
 581 Who Have Sex With Men in Human Immunodeficiency Virus-1 Transmission Networks in Guangzhou,  
 582 China. *Open Forum Infectious Diseases* 7, ofaa154. doi:10.1093/ofid/ofaa154

## TABLES

### 583 Add p-values for DWH

**Table 1.** clustuneR Comparison

Dataset	clustuneR		AUTO-TUNE		
	Threshold	Avg. Homophily	Threshold	Avg. Homophily	Max Score
Middle Tennessee	0.0160	0.0079	0.01431	0.0147	1.25807
Seattle	0.0160	0.0259	0.01354	0.0348	1.53325
Northern Alberta	0.0104	-0.0536	0.01099	-0.0448	1.01678

**Table 2.** Threshold Comparison with Prior Publications Citing HIV-TRACE

PMID	Country	Collection Date	Threshold Used	AUTO-TUNE
29975689	South Africa	2011-2015	2.5%	2.584%
30574123	USA	1997-2008	2%	1.848%
32500089	China	2008-2015	0.5%	0.675%
32693608	Uganda	2009-2016	1.5%	1.707%
33281803	China	2000-2016	0.5%/0.7%	0.676%
33901684	China	2008-2012	1.5%	1.215%
34452506	Canada	1996-2017	1.5%/2.5%	0.547%
31041344	USA	1997-2017	1.5%	0.927%

**Table 3.** CRF07\_BC DWH and Panmictic Range at Different Thresholds

Record	Threshold 1.5%		Threshold 0.76%		Threshold 0.19%	
	DWH	Panmictic Range	DWH	Panmictic Range	DWH	Panmictic Range
MSM	0.211	[-0.105, -0.205]	0.237	[-0.120, -0.240]	0.292	[-0.146, -0.280]
Hetero	0.133	[-0.092, -0.190]	0.185	[-0.100, -0.211]	0.25	[-0.093, -0.256]
PWID	0.168	[-0.001, -0.089]	0.401	[-0.005, -0.081]	0.445	[-0.012, -0.129]

## 6 FIGURE CAPTIONS

**Figure 1.** AUTO-TUNE scoring across candidate thresholds ranging from 0% to 2.5% genetic distance. The plots represent the datasets from Seattle, Middle Tennessee, and Northern Alberta as described by clustuneR. The y-axis represents the AUTO-TUNE score, with higher scores suggesting more optimal thresholds. The x-axis represents the candidate thresholds. None of the three datasets exhibited an extreme peak of over 1.9, implying multiple thresholds could serve well and that a more complex decision-making process that includes downstream metrics such as DWH or careful inspection may be necessary.

**Figure 2.** AUTO-TUNE scoring across candidate thresholds from 0% to 2.5% genetic distance for eight datasets from various studies that have previously employed HIV-TRACE with qualitatively defined thresholds. Each plot represents one dataset, and the y-axis shows the AUTO-TUNE score. Higher scores indicate more optimal thresholds for clustering. The x-axis represents the range of candidate thresholds. Each plot is labeled by the respective studies' PubMed ID. Cases such as Dalai et al. and Bbosa et al., exemplify the potential for substantial score variation even within very narrow distance intervals, underscoring the value of a more granular and systematic approach to threshold selection.

**Figure 3.** Comparative visualizations of HIV-1 CRF07\_BC networks at different thresholds, colored by risk factor: heterosexual (green), person who injects drugs (light blue), and men who have sex with men (dark blue). Panel A represents the network at a 1.5% threshold, encompassing 5923 nodes. Panel B illustrates the network at the 0.76% threshold, as indicated by the highest AUTO-TUNE score, capturing 3537 nodes. Panel C displays the network at a 0.19% threshold, corresponding to the second highest AUTO-TUNE score, comprising 1654 nodes. Panel D features the AUTO-TUNE score plot, spanning from 0% to 0.5% thresholds, with significant peaks at 0.76% and 0.19%

**Figure 4.** Comparison of HIV networks before and after AUTO-TUNE optimization. The left panel shows the network constructed using HIV-TRACE's default threshold of 1.5%, while the right panel displays the network after applying AUTO-TUNE's optimal threshold of 1.699%. Notably, the largest cluster, primarily consisting of samples from China, Thailand, and Vietnam, is reinforced with more nodes in the AUTO-TUNED network. Despite the increase in threshold, overall homophily among other countries remains consistent.

**Figure 5.** (A) Box plot representing the AUTO-TUNE scores across ten random samples at 25%, 50%, and 75% of the Rhee et al. (2019) dataset, showing a trend of increasing confidence in score estimates with denser sampling. (B) Box plot of the selected distance thresholds across the same random samples at 25%, 50%, and 75% proportions, demonstrating improved consistency in threshold selection with increased sample size. (C) Scatterplot of the chosen thresholds (Y-axis) against their corresponding AUTO-TUNE scores (X-axis) for the three subsample proportions.