# Efficient indexing and querying of annotations in a pangenome graph

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## **Abstract**

The current reference genome is the backbone of diverse and rich annotations. Simple text formats, like VCF or BED, have been widely adopted and helped the critical exchange of genomic information. There is a dire need for tools and formats enabling pangenomic annotation to facilitate such enrichment pangenomic references. The Graph Alignment Format (GAF) is a text format, tab-delimitted like BED/VCF files, which was proposed to represent alignments. While it could represent any annotation in a pangenome graph, there are no tools to index and query them efficiently.

Here, we present extensions to vg and HTSlib that provide efficient sorting, indexing, and querying for GAF files. With this approach, annotations overlapping a subgraph can be extracted quickly. Paths are sorted based on the IDs of traversed nodes, compressed with BGZIP, and indexed with HTSlib/tabix that we extended to work with the GAF format. Compared to the GAM format, GAF files are easier to write text files, and we show that they are twice as fast to sort and twice as small on disk. In addition, we updated vg annotate to better produce graph annotations, as paths, starting from annotation files relative to linear references. More precisely, it can to take annotations in BED or GFF3 files, written relative to reference paths or haplotypes, and produce GAF files representing their paths through the pangenome.

We showcased these new commands on several applications. We projected annotations for all haplotypes in the latest draft human pangenome (HPRC v1.1 GRCh38-based Minigraph-Cactus pangenome), including genes, segmental duplications, tandem repeats and repeats annotations. We also projected known variants from the GWAS catalog and expression QTLs from the GTEx project to the pangenome. Finally, we reanalyzed ATAC-seq data from ENCODE to highlight how a coverage track could look like in a pangenome graph. These rich annotations can be quickly queried with vg and visualized using existing tools like the sequenceTubeMap or Bandage.

## Introduction

The current reference genome is the backbone of diverse and rich annotations. It is used as a reference to map sequencing reads. The coordinate system it provides is used by large annotation databases gathering functional elements, known variant information, genomic elements. Over the past decades, the organization and visualization of these annotations have been central to understanding and sharing results from genomic studies. In practice, annotations are typically saved in additional files following a text format that can be compressed and indexed for fast query (e.g. VCF, BED, GFF). Under the hood, the HTSlib library supports the indexing of the most used file formats[1]. These annotation files are easy to write and to load in software like IGV[2] or websites like the UCSC Genome Browser[3].

With the improved genome sequencing technologies, more high-quality genomes can be produced and combined into pangenomes. A pangenome represents multiple genomes, often as a graph describing adjacency (edges) of the sequences (nodes) in the genomes. Tools to work with such pangenomes are still in their infancy, although there are now several options that shows improved performance over traditional approaches. In particular, sequencing data analysis benefits from using pangenomes as a reference for read mapping[4,5], variant calling[6], peak calling[7,8], or transcript expression quantification[9]. Hence, we are now manipulating genomic objects, like sequencing reads, epigenetic regions, or genomic variants, in the pangenome graph space. Currently, those results are typically projected to the linear reference genome. As for linear genome reference, organizing and visualizing genomic annotation in the pangenome will be essential. A lack of user-friendly querying and visualization options would hamper the adoption of the pangenomic paradigm.

Several interactive visualization tools for pangenomes exist but mostly focus on representing the graph topology with specialized integration of additional information layers. Bandage is an interactive assembly graph visualization tool that can scale to pangenomes up to hundreds of thousands of nodes on modern computers[10]. GfaViz is another interactive sequence graph visualization tool that supports the GFA 2 format and its new features[11]. In particular, it can represent the different paths annotated in the GFA file. GfaViz cannot load additional annotation files: the annotations, written for example as GFA paths, must be included in the main pangenome file to be accessible during exploration. The sequenceTubeMap visualizes the pangenome and sequencing reads aligned to it, allowing the user to query specific regions[12]. MoMI-G, or Modular Multi-scale Integrated Genome Graph Browser, focuses on visualizing structural variants[13]. Variants are interactively selected and represented with a pangenome view as a custom sequence Tube Map representation. Supporting reads and genome annotations can also be included in that representation. Only the reference genome path can be annotated from files in BED/GFF/WIG format. Panache is another pangenomic tool specializing in gene-centric visualization of a linearized view of the pangenome, where blocks of homologous sequences are represented side by side[14]. The blocks are interactively explored in a web-based application to analyze their gene content and their absence/presence in the different genomes. Panache is notably used in the Banana Genome Hub[15]. In summary, some visualization tools exist, but the best way to provide additional annotation layers to their graph representation needs to be clarified.

A few other options exist to display static representations of a pangenome graph (or subgraph). The vg toolkit can output a pangenome and alignments in the DOT format. Paths can be displayed too, although the image can become hard to read when many paths are included or in large graphs. The odgi toolkit offers visualizations that scale better to larger graphs and annotations[16]. The pangenome is, for example, linearized using a 1D layout[17], and annotations are displayed on top of the graph as an array/heatmap. Of note, odgi implemented two options to add annotations from external BED files: one to convert annotations to a CSV file to help color nodes in Bandage, another by injecting new paths into a graph before visualization. One limitation to injecting new paths into a pangenome is that paths must start at the beginning of a node as we cannot provide an *offset* value to specify the base at which a path must start.

There is a dire need for a format supporting annotations in the pangenome that is easy to create, index, and query. As demonstrated by the success of the BED, GFF, or VCF formats, it could help the critical exchange of (pan)genomic information and allow pangenomic tools to access additional information in the pangenomic space. The Graph Alignment Format (GAF) text format, which was proposed to represent alignments, could describe any type of annotation in a pangenome graph. The lack of techniques to compress, index, and query it limits its adoption at a larger scale. Here, we present new features of the vg and HTSlib ecosystem that provide efficient indexing and querying for pangenomic annotations represented as paths in the GAF format. We illustrate its values in several applications: projecting and visualizing gene and repeat annotations, summarizing open-chromatin from epigenomics sequencing datasets, or positioning known variants in the pangenome.

## **Methods**

## Indexing paths in GAF files

The sorting and indexing algorithm is most efficient and makes sense when node IDs are integer sorted based on the topology of the pangenome graph. This is the case for pangenomes constructed by minigraph-cactus[18], PGGB[19], or vg construct [4]. Otherwise, the pangenome graph can be *sorted*, i.e. changing the node IDs, using vg or odgi [16]. If the node IDs are sorted integers, a short path through the graph should only traverse nodes with IDs contained in a small interval (see Figure 1A). The main approach of the GAF sorting and indexing approach is to work with those intervals.

Hence, to sort a GAF file, each path is first parsed to extract the minimum and maximum node IDs (see Figure 1B). The paths are then sorted first by their minimum ID, and then by their maximum ID. This is similar to the approach used to sort BED or VCF files on linear genomes: they are sorted by sequence name, then start position, then end position.

Figure 1: Path sorting and indexing using vg and HTSlib/tabix A. A region of pangenome is represented with nodes (containing sequences) in yellow and edges in black. The node IDs are topologically sorted integers, ranging here from 23 to 36. Three paths are highlighted in red, blue and green. B. The three paths are written with the GAF syntax, by specifying the orientations (</>
/>) and IDs of the traversed nodes. For each path, the node range n-N, between the minimum and maximum node IDs, is used for sorting the path. C. Overview of the workflow to sort a GAF file using vg gamsort, compress it with bgzip and index using tabix. D. The small .tbi index file helps query slices of the GAF file quickly. For example using tabix, or vg subcommands like find or chunk.

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A GAF sorting feature has been added to the vg toolkit, within the gamsort subcommand (see Figure 1C). It first sorts chunks of GAF records (e.g. reads), to avoid having to hold the entire set in memory. The sorted chunks are then merged into the final sorted GAF file. This GAF sorting implementation was introduced in vg version 1.56. A sorted GAF file can be compressed in the BGZF format using bgzip. The BGZF format is an implementation of the standard gzip format that compresses a file by block to facilitate their random access. It is used to compress among others, VCF, BED, or BAM files with HTSlib[1].

HTSlib[1] was then modified to index bgzipped GAF files. Similar to other tab-separated file like VCF or BED, a <code>gaf</code> preset was added to <code>tabix</code>. For BED or VCF, <code>tabix</code> extract the interval information from the columns with the sequence names and <code>genomic</code> position[20]. In the new <code>gaf</code> preset, it instead parses the path information in the GAF file to extract the minimum and maximum node IDs. The indexing is then based on this interval, the same as used for the sorting described above.

We tested the GAF sorting and indexing performance on 30X coverage Illumina short reads from HG002. The reads were downloaded from https://s3-us-west-2.amazonaws.com/human-pangenomics/NHGRI\_UCSC\_panel/HG002/hpp\_HG002\_NA24385\_son\_v1/ILMN/downsampled/. We mapped them using giraffe[5] on a personnalized pangenome[21] from the HPRC v1.1 Minigraph-Cactus pangenome. The reads were outputted in GAM first to compare the file size and sorting runtimes. The GAM file was sorted with vg gamsort. It was then converted to a GAF file using vg convert and sorted using vg gamsort with the new GAF sorting mode described above. We compared the file sizes and sorting runtimes between both approach. The query time was also measured on ten thousand regions of the pangenome defined as node ranges. Those node ranges were defined by picking a random starting node position the reference path and walking 50 steps along that path. The commands and details for this benchmarking is available in the analysis/readsorting folder of this paper's repository[22].

## **Querying GAF files**

Instead of indexing on a sequence name and genomic position, we can query on a node interval (see Figure 1D). In HTSlib, tabix was modified to disregard the sequence name when querying intervals for a GAF file. The interval is defined by the values typically used for the *start* and *end* position of genomic coordinates.

Commands to query slices of the pangenome in vg were also updated. The find and chunk subcommands use the updated HTSlib library to extract the appropriate paths from the nodes selected. Internally, those commands identify a first subgraph, for example corresponding to a genomic interval on the reference path provided by the user. This subgraph is then extended with more *context* to include non-reference regions of the graph. The amount of context is also controlled by the user. Finally, the subgraph or the paths (usually reads) overlapping these nodes are extracted. This last step was updated to be able to extract paths in an indexed GAF file using HTSlib. As for sorted GAM files, it is now possible to extract a slice of a indexed GAF file based on node intervals, coordinates on a reference path, multiple coordinates in a BED file, a provided subgraph (see User Guide in the analysis folder of this paper's repository[22]).

## Projecting annotations onto a pangenome

A pangenome represents multiple genomes for which annotations might be available. We describe an approach to project annotations relative to a genome onto a pangenome. We recently updated the annotate subcommand from the vg toolkit to project regions represented in the BED or GFF files onto the latest pangenomes from the HPRC. Currently, the genome to annotate must be a *reference* path in the pangenome, but any path can be set as reference using the vg gbwt subcommand (~1 minute). Then, vg annotate looks for the location of a path in pangenome graph for each input region. The path, represented as an *alignment* record, is then written either in GAM or GAF formats. Of note, a path can be broken in multiple disjointed parts. It happens in the recent human draft pangenome when some regions are clipped out, for example across centromeres or when creating suspiciously large structural variants. Projected annotations are hence also broken up if needed when they overlap with breakpoints. The name of the annotated path in the output GAF is picked from the BED file's 4th column, or the *Name* and *ID* GFF fields.

We test this approach by projecting the gene annotation, repeat annotations, and segmental duplication annotation for each of the 88 assembled haplotypes in the draft human pangenome from the HPRC (v1.1). These annotations were downloaded from the HPRC public repository at <a href="https://raw.githubusercontent.com/human-">https://raw.githubusercontent.com/human-</a>

pangenomics/HPP\_Year1\_Assemblies/main/annotation\_index/. The gene annotations from CAT[23] (GFF files) were downloaded using the URLs in the

Year1\_assemblies\_v2\_genbank\_CAT\_genes.index file, the repeat annotations from RepeatMasker (BED files) from Year1\_assemblies\_v2\_genbank\_Repeat\_Masker.index, segmental duplications (BED files) from Year1\_assemblies\_v2\_genbank\_Seg\_Dups.index, and simple repeats from Year1\_assemblies\_v2\_genbank\_TRF.index. A helper script was implemented to prepare the BED files with informative names. For example, for repeats from the RepeatMasker annotation, we name each element by their repeat class and repeat family. The projection of those annotations for each haplotype was automated with a Snakemake workflow available in the analysis/annotate folder of this paper's repository[22].

## Coverage track from mapped reads

Functional genomics datasets are often visualized as a coverage track. High coverage in a region might suggest a strong transcription factor binding site or regulatory region.

We implemented an approach to summarize the coverage of reads across the pangenome into paths with similar coverage (see Figure 2). The coverage in every node is first binned into a few coverage classes, for example representing low, medium and high coverage. By default we use 1, 5, and 30 reads as coverage breakpoints to save three bins: 1-5, 5-30, 30+. Regions with no coverage are not saved. Once the coverage is binned, we extend greedily to connected nodes and bins if in the same coverage class. This extension step produces path through the pangenome with consistent coverage.

The paths are written in the GAF format, recording the coverage class and the average coverage across the path.

Figure 2: Read coverage bin-and-extend approach to produce coarse-grained coverage tracks In each node (grey rectangles), the read coverage is first binned using user-defined coverage bins (red blocks). Each bin is then extended, one at a time (green flags), until reaching a different coverage bin (red flags).

**Figure 2:** Read coverage bin-and-extend approach to produce coarse-grained coverage tracks In each node (*grey rectangles*), the read coverage is first binned using user-defined coverage bins (*red blocks*). Each bin is then extended, one at a time (*green flags*), until reaching a different coverage bin (*red flag*).

This algorithm was implemented in Python and uses the *libbdsg* module[24] to query the pangenome. It is made available in the public repository of this study in the analysis/encode folder of this paper's repository[22].

## **Annotating known variants**

Variants from public databases were annotated when matching a variant site in the pangenome. We implemented an approach to look for input variants in the pangenome and write a GAF file with the path followed by those variants in the pangenome. The variants are matched with the VCF representation of the HPRC Minigraph-Cactus v1.1 pangenome, produced by running vg deconstruct on the pangenome file. This VCF file contains the all variants and their paths through the pangenome in the *AT INFO* field. We look for the variants of interest in this VCF and extract the appropriate path using the *AT* field. The annotation path follows either the alternate allele of the variant if known and present in the pangenome, or the reference allele path if the alternate allele is unknown but there is a variant at this position in the pangenome. Variant that are not matched in the pangenome are skipped. This algorithm was implemented in Python and uses the *libbdsg* module[24] internally to get node size information necessary to write proper GAF records.

We test this approach on the GWAS catalog[25] and GTEX expression QTLs[26]. The GWAS catalog was downloaded from the UCSC Genome Browser gwasCatalog table[3]. The eQTLs from GTEX v8 were downloaded from https://storage.googleapis.com/adult-gtex/bulk-qtl/v8/single-tissue-cis-qtl/GTEx\_Analysis\_v8\_eQTL.tar. The files defining associations between variant/gene pairs were parsed for each tissue. As before, the output annotation in GAF were bgzipped, sorted and indexed.

We implemented another, more straightforward approach, to annotate variants that were genotyped using the pangenome. Here we simply convert a VCF that contain allele traversal information (*AT* field) to a GAF file representing the alternate allele. We test this approach by genotyping HG002 from short-reads Illumina Novaseq data. more details

The scripts and pipeline to annotate variants is available in the public repository of this study in the analysis/variants folder of this paper's repository[22].

## Visualization in the sequenceTubeMap

The sequenceTubeMap was developed to interactively explore a pangenome graph, haplotypes traversing it, and reads mapping to it[12]. It internally calls vg to extract the relevant slice of the pangenome graph and reads. Only accepting reads in the GAM format, sorted and indexed vg, were originally accepted. We updated sequenceTubeMap to accept GAF files that have been sorted, compressed and indexed as explained above.

The new version of the sequenceTubeMap can also display multiple layers of haplotypes or reads. As *reads*, the user can now add layers of annotations written indexed GAF files. A different color or color palette can be assigned to each layer to facilitate the visualization of different datasets in the same local pangenome region. Moreover, a new option has been added to use an element's opacity to represent an integer score, like a mapping quality or read coverage. For example, one could visualize the coding regions of genes and coverage tracks for different cell types from the ENCODE project, as described below and highlighted in Figure 4.

## **Visualization in Bandage**

A fork of Bandage[10], called BandageNG[27], can read GAF files and visualize its paths through the input graph by coloring the nodes. Of note, the offset positions are not taken into account, which means that nodes are colored entirely even if the path only overlaps part of it. We implemented a wrapper script to facilitate the preprocessing of a subgraph to visualize with BandageNG. It starts by extracting the subgraph of a full pangenome for a region of interest using vg chunk. This graph is converted to GFA, making sure that the path are written last. The script also extracts annotations from indexed files on that same region. Those GAF files are merged and modified if necessary: path names are de-duplicated and paths are trimmed when traversing nodes absent from the extracted subgraph. These modifications ensure that the final GAF file can be loaded in BandageNG. The outputs of this script can be opened with BandageNG for interactive exploration. In particular, the *Find path* feature can find nodes corresponding to specific paths, which are either haplotypes or annotations in this preprocessed GFA file. The user can select a *path* and color its nodes. The helper script and a tutorial are available at the analysis/visualization folder of this paper's repository[22].

## Results

## Sorting and indexing short sequencing reads

Read sorting was tested on Illumina HiSeq paired-end short reads for the HG002 sample, each 150 bp long and a genome coverage of about 30X. The gzipped GAF file with about 682 million reads was sorted in 6h32 using a single thread and about 2Gb of memory (Table 1). Indexing the sorted GAF with tabix took 18 mins. We compared that approach with the current implementation in vg which uses files in the GAM format. GAM is a protobuf alignemnt format introduced in Garrison et al.[4]. Sorting a GAM with the same reads took 11h47 using a single thread and about 6 Gb of memory.

In addition to being about twice as fast to sort, reads written in the GAF format (and bgzipped) also take about half the disk space (52G vs 108G). The main reason for this reduced space is that GAF doesn't save the full read sequence, only the path through the pangenome and edits to reconstruct it. GAM files produced by vg giraffe can also save additional information as annotations, like the mapping time of each read, that are currently not kept when converting to the GAF format. Overall, the bgzipped GAF files are half as small and twice as fast to sort for short sequencing reads.

Once indexed, extracting a slice of the pangenome is as efficient as extracting a slice of an indexed BAM/VCF/BED file for a genomic region. For example, extracting reads for ten thousand random regions in the pangenome took about 0.07 second per region to retrieve an average of 1707 reads. For comparison, the same extraction took on average 0.8 second per region using the GAM format.

**Table 1:** Resources used to sort short sequencing reads for a 30x coverage Illumina human dataset.

Format	Time (H:M:S)	Max. memory used (Kb)	File size (Gb)
GAM	11:46:58	6,236.60	108

Format	Time (H:M:S)	Max. memory used (Kb)	File size (Gb)
GAF	6:50:28	1,904.83	52

## Annotation of a human pangenome

To showcase these commands, we projected annotations for all haplotypes in the latest draft human pangenome (HPRC v1.1 GRCh38-based Minigraph-Cactus pangenome)[28]. This included genes, segmental duplications, tandem repeats and repeats annotations. vg annotate can annotate ~4M gene annotations in ~16 mins, and ~5.5M repeats from RepeatMasker in ~9 mins on a single-threaded machine. Finally, these rich annotations can then be quickly queried with vg and visualized using existing tools like the sequenceTubeMap or Bandage. Using BandageNG, a fork that can import paths in GAF files, paths were searched and colored to illustrate a mobile element insertion.

Figure 3: Visualization examples with BandageNG Example of a AluYa5 transposon insertion (red) within the coding sequence of the PRAMEF4 gene (blue). Both annotations were originally produced at the haplotype level by the Human Pangenome Reference Consoritium. We projected them to the pangenome, indexed them, and queried a small region to visualize with BandageNG. The nodes were colored based on those annotations, loaded as paths by BandageNG.

**Figure 3: Visualization examples with BandageNG** Example of a AluYa5 transposon insertion (*red*) within the coding sequence of the *PRAMEF4* gene (*blue*). Both annotations were originally produced at the haplotype level by the Human Pangenome Reference Consoritium. We projected them to the pangenome, indexed them, and queried a small region to visualize with BandageNG. The nodes were colored based on those annotations, loaded as paths by BandageNG.

We also matched and annotated more than 660 thousand variants from the GWAS catalog[25] and expression QTLs from the GTEx catalog[26] across 49 tissues (on average 1.45 million variants per tissue). On average, 94% variants were found in the HPRC pangenome. The variants files in GAF take only 907Mb of space and can be queried fast for visualization in the sequenceTubeMap or Bandage. This annotation is showcased in Figure 4B. Of note, it is also straightforward to convert genotypes for variants from the pangenome to annotated paths. This could be the case when the pangenome is used as the backbone of the genotyping of new samples, for example with Pangenie or *vg call*. To illustrate this use case, we genotyped HG002 using short Illumina reads aligned on the draft human pangenome with vg giraffe and vg call. The predicted genotypes were converted to GAF and indexed. They could be visualized using the sequenceTubeMap, for example, along with the aligned reads. This type of annotation can help a developer explore variant calls and aligned reads. A example is shown in Figure 4C. On the traditional reference genome, the equivalent would be to load both reads and variants in IGV[2].

Figure 4: Visualization examples with the sequenceTubeMap A Projection of the HPRC gene annotation, here visualizing a coding sequence (CDS) of the CFD gene. There is an insertion (see insertion node in the middle), that the reference path (violet on top) skips. Most CDS annotations are similar (blue paths), but the CSD from the haplotype with the insertion seems to be split in two parts, highlighted in green. B. Projecting known variants in the pangenome. black: Reference path (GRCh38), pale colors: other human haplotypes, reds: GWAS catalog, blues: eQTLs across tissues (GTEx). C. Aligned reads and genotype calls. yellow/green: annotation paths from vg call genotypes. reds/blues: short sequencing reads. D. ATAC-seq coverage tracks for 7 tissues from the ENCODE project. The reference path (violet) and other HPRC haplotypes (greys) are shown on top and traverse two variation sites. ATAC-seq coverage track for the different tissues are shown in different colors with different opacity representing coverage level. The thyroid track (blue) is more opaque than others suggesting that this region is more open in this tissue. The red tracks at the bottom show the position of exons from the TPO gene, a thyroid-specific gene.

**Figure 4: Visualization examples with the sequenceTubeMap A** Projection of the HPRC gene annotation, here visualizing a coding sequence (CDS) of the *CFD* gene. There is an insertion (see *insertion node* in the middle), that the reference path (*violet* on top) skips. Most CDS annotations are similar (*blue* paths), but the CSD from the haplotype with the insertion seems to be split in two parts, highlighted in *green.* **B.** Projecting known variants in the pangenome. *black*: Reference path (GRCh38), *pale colors*: other human haplotypes, *reds*: GWAS catalog, *blues*: eQTLs across tissues (GTEx). **C.** Aligned reads and genotype calls. *yellowl green*: annotation paths from vg call genotypes. *redsl blues*: short sequencing reads. **D.** ATAC-seq coverage tracks for 7 tissues from the ENCODE project. The reference path (*violet*) and other HPRC haplotypes (*greys*) are shown on top and traverse two variation sites. ATAC-seq coverage track for the

different tissues are shown in different *colors* with different opacity representing coverage level. The thyroid track (*blue*) is more opaque than others suggesting that this region is more open in this tissue. The *red* tracks at the bottom show the position of exons from the *TPO* gene, a thyroid-specific gene.

Using sequenceTubeMap, haplotypes, read alignments and paths can be visualized interactively.

## **Coverage of seven functional datasets from ENCODE**

We aligned ATAC-seq datasets from 7 cell types to the draft human pangenome to produce coverage tracks as indexed GAF files. On average, there were about 475 thousand paths representing high read coverage which were 2.6 nodes (104.8 bases) long. On average, 63 thousand paths with high ATAC-seq read coverage traversed more three or more nodes, i.e. regions of the pangenome with variation (see Table 2).

**Table 2:** High coverage tracks from seven functional datasets on the HPRC pangenome. For each sample, the table shows how many paths had a mean coverage of at least 10 reads, and how long they were.

Dataset	Paths	Average bases	Average nodes	Traversing >2 nodes
Breast epithelium	570,155	109.80	2.68	65,095 (11.4%)
Gastrocnemius medialis	342,830	94.47	2.43	53,595 (15.6%)
Gastroesophageal sphincter	555,618	115.69	2.73	72,407 (13%)
Peyer's patch	270,094	97.88	2.62	43,591 (16.1%)
Sigmoid colon	678,380	115.10	2.69	82,906 (12.2%)
Spleen	531,910	104.97	2.66	67,153 (12.6%)
Thyroid gland	377,157	95.63	2.41	56,441 (15%)

It took on average 7 cpu.hours to map the reads to the pangenome using VG Giraffe, and 2.8 cpu.hours to produce the coverage tracks. Sorting, compressing and indexing them took only 0.23 cpu.hours, on average. Table 3 compiles the runtimes and memory used for each step across all samples.

Figure 4D shows an example of a region, visualized with a tweaked version of the sequenceTubeMap (see Methods), near an exon of the *TPO* gene. The exon annotations came from the projection of the HPRC gene annotations described above. The coverage track for the ATAC-seq dataset shows that this region is only opened in the thyroid, consistent with the thyroid-specific role of *TPO*. By integrating these two external sources of annotations (gene annotation and ATAC-seq coverage), we can visualize them in the context of the genomic variation in the pangenome.

## **Discussion**

The tools and applications described here present an option to streamline the production of annotations in the GAF format, their indexing and efficient querying. First, these techniques will help integrate additional information into existing or future pangenomic tools. They will simplify the integration of information like gene or repeat annotations, known variants, or more generally other functional annotations, in pangenomic analysis. These analyses will be able to work with those annotations in light of underlying genomic variation recapitulated in the pangenome. Visualization is a specific example where the users typically want to include several layers of annotations, including some custom-made. Tools to visualize pangenomes will greatly benefit from a simple pipeline to create or load pangenomic annotations. Second, this work is critical to allow the community to make, share and reuse annotations in the pangenome space. The GAF format is already used by multiple

independent tools, although mostly read mappers. Thanks to our work, GAF files can now be indexed and queried efficiently, like BED, VCF, or GFF on a linear reference genome. We also showcase how annotations on a linear genome can be converted to a path in the pangenome in the GAF format. For these reasons, we believe it could become the de facto format to represent annotations in the pangenome and accelerate the adoption of the new pangenomic paradigm by the broader genomics field. Although informative annotations could be analyzed already, the current approach has some limitations.

The indexing scheme relies on integer node IDs, compacted relative to their position in the pangenome. If node IDs are not integer, it adds a layer of complexity for the user as they will need to convert them to integers and keep track of the translation between original pangenome and new pangenome. Both vg and odgi offer ways to convert pangenome to a new pangenome with compact integer node IDs.

The current implementation is designed with short paths in mind, where the user wants to extract the full annotated paths in a region. It is convenient when working with short reads, gene annotations, and most genomic repeats, for example. Larger genomic regions, like chromosome bands, large assembled contigs, or large segmental duplications, can still be represented and manipulated, but might be less practical to work with. With the current implementation, the full annotated region will be extracted when querying an overlapping region. The extracted annotations are not clipped to the specified range which could be inconvenient to a user who zooming into a small region but would still like to know about the much larger annotated path traversing this subgraph. To address this use case, we plan to implement an extraction mode where output annotations are trimmed to keep only the queried subgraph.

The naive metadata integration is another limitation. Indeed, the annotation information is currently reduced to a single label. In our applications, those label would contain for example a gene name and haplotype names, or a read coverage bin and average coverage value, all in one label. This was easy to implement and sufficient for now because there are no tools that handle more advanced metadata organization. For many annotations, it would be useful to keep the metadata better organized, so that the user can access/use it within visualization tools. The different metadata could be saved using optional tags that can be added at the end of each GAF record.

Our approach to convert annotations from a linear genome to the pangenome assumes that we have annotations of the different haplotypes in the pangenome. There is still no clear solution to lift annotations from one reference/haplotype to other haplotypes in the pangenome, except through reanalysis/reannotation of each haplotype. The homology information embedded in the pangenome could potentially be used to propagate annotations from one haplotype to others more easily. This strategy can already be used by annotation tools like CAT[23], as an additional source of gene annotation evidence. In the future, these techniques might help propagate other types of annotations across the pangenome more efficiently than by reanalyzing the raw data from scratch on each haplotype. something about odgi paths/untangle maybe

It also highlights the limitations of the existing tools to integrate these files. Some tools, like Bandage or GfaViz, require manual pre-processing, for example extracting a subgraph and integrating the annotations as embedded paths. The sequenceTubeMap can now handle indexed bgzipped GAF files, but the query time for large pangenome remains long in practice. Defining and integrating annotation metadata into its interface will also require a significant amount of development. Overall, we stress the need for visualization tools that can efficiently layout and organize many paths through a pangenome.

We showed that the GAF format, thanks to new tools for their efficient manipulation, could offer a path for the future of annotations in pangenome graphs. While it provides an important building

block, it is clear that more needs to be done to make it a useful solution for the community.

## Code and data availability

vg is available at <a href="https://github.com/vgteam/vg">https://github.com/vgteam/vg</a>. The fork of Bandage that allow for path coloring is available at <a href="https://github.com/vgteam/sequenceTubeMap">https://github.com/vgteam/sequenceTubeMap</a> is hosted at <a href="https://github.com/vgteam/sequenceTubemap">https://github.com/vgteam/sequenceTubemap</a>

The analysis presented in this manuscript is documented in the <a href="https://github.com/jmonlong/manu-vggafannot">https://github.com/jmonlong/manu-vggafannot</a>. It contains scripts used to prepare the annotation files, commands and automated pipelines used to annotate them in the pangenome, and helper scripts to summarize the output files.

The annotations of the HPRC v1.1 pangenome were deposited on Zenodo at ??ZENODO\_LINK??. This includes gene annotations, repeats from RepeatMasker, simple repeats and segmental duplications. Coverage tracks for the seven ENCODE ATAC-seq samples are also available in this repository.

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## **Supplementary material**

**Table 3:** Compute resources used for the analysis of the functional datasets and production of the indexed coverage tracks.

Dataset	reads (M)	read mapping	coverage track	sorting + compressing + indexing
Breast epithelium	193.6	8.9 cpu.h (54 Gb)	3 cpu.h (109 Gb)	0.3 cpu.h (1 Gb)
Gastrocnemius medialis	98.8	4.8 cpu.h (54 Gb)	2.6 cpu.h (99 Gb)	0.3 cpu.h (1 Gb)
Gastroesophageal sphincter	168.5	7.3 cpu.h (54 Gb)	3 cpu.h (108 Gb)	0.2 cpu.h (1 Gb)
Peyer's patch	145.3	8 cpu.h (54 Gb)	3 cpu.h (104 Gb)	0.2 cpu.h (1 Gb)
Sigmoid colon	173.5	8.2 cpu.h (54 Gb)	3 cpu.h (106 Gb)	0.3 cpu.h (1 Gb)
Spleen	157.2	7.6 cpu.h (54 Gb)	3 cpu.h (104 Gb)	0.2 cpu.h (1 Gb)
Thyroid gland	91.4	4.6 cpu.h (54 Gb)	2.5 cpu.h (94 Gb)	0.1 cpu.h (1 Gb)