An international virtual hackathon to build tools for the analysis of structural variants within species ranging from coronaviruses to vertebrates [version 1; peer review: 4 approved with reservations]

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Abstract
In October 2020, 62 scientists from nine nations worked together remotely in the Second Baylor College of Medicine & DNAnexus hackathon, focusing on different related topics on Structural Variation, Pan-genomes, and SARS-CoV-2 related research. The overarching focus was to assess the current status of the field and identify the remaining challenges. Furthermore, how to combine the strengths of the different interests to drive research and method development forward. Over the four days, eight groups each designed and developed new open-source methods to improve the identification and analysis of variations among species, including humans and SARS-CoV-2. These included improvements in SV calling, genotyping, annotations and filtering. Together with advancements in benchmarking existing methods. Furthermore, groups focused on the diversity of SARS-CoV-2. Daily discussion summary and methods are available publicly at https://github.com/collaborativebioinformatics provides valuable insights for both participants and the research community.

Keywords
Structural variant, CNV, SARS-CoV-2, NextGeneration Sequencing

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Introduction

Structural variants (SVs) comprise a number of genomic imbalances including copy number variations (CNVs), insertions (INS), deletions (DELS), inversions (INVs), duplications (DUPs), and inter-chromosomal translocations.1–3 SVs have been implicated as clinically significant mutations with proven associations to multiple diseases.4,5 Despite next-generation sequencing becoming increasingly common within the field of biomedical research, several practical challenges exist for comprehensively detecting and evaluating SVs particularly in regard to the high false positive or negative rate along with the accuracy of breakpoint prediction.6,7 While SV detection with genotyping arrays remains the most commonly used method, the toolbox for SV detection is expanding to incorporate the advancements in third generation sequencing technologies provided by Pacific BioSciences,9 Oxford Nanopore Technologies,9,10 optical mapping and NanoString11 to name a few. These advancements offer potential for solving previously unresolved structural variants.

In October 2020, 62 scientists from nine nations worked together remotely in the Second Baylor College of Medicine & DNAnexus hackathon, focusing on different related topics on SV, Pan-genomes, and SARS-CoV-2 related research. Consequently, this international structural variation hackathon meeting focused on eight themes: 1.) efficiently genotyping vast quantities of SVs; 2.) mapping CNVs to SV types; 3.) detecting and validating SVs for SARS-CoV-2; 4.) filtering high-confidence SV calls for clinical genomics; 5.) SV read-based phasing for haplotype analysis; 6.) genome graph generation without a reference; 7.) machine learning approaches to predict lab-of-origin of a sample; and 8.) gene-centric data browsing for SV analysis.

Overall, this manuscript details our tools’ objectives, value-add, implementations, and applications and sets the foundation for further concept development beyond. In this article we present 10 software tools that were the results of this hackathon.

NibbleSV: efficient genotyping of SVs from short read datasets. Detection of SVs longer than a short-read (<500bp) DNA trace is very challenging as the SV allele becomes split across multiple reads. To this end, long read sequencing technology is preferential for overcoming this challenge however, although long read sequencing has proven more accurate in SV identification, obtaining accurate allele frequencies across a population is important in order to rank and identify potential pathogenic variations.

Thus, it is still important to genotype SV events in pre-existing short read datasets such as those provided by the 1000 genomes project, Topmed, CCDG, etc. Recently, two main approaches, Paragraph12 and VG,13 have achieved this with high accuracy even for insertion SV events. However, these methods are computationally expensive particularly when the number of SVs to be genotyped per sample increases. Furthermore, and maybe more crucially, both methods rely on precise breakpoints that do not change in other samples, an assumption that is potentially flawed particularly over repetitive regions. NibbleSV is a software package able to efficiently genotype vast quantities of SVs whilst also using a kmer catalogue of SVs in order to circumvent the need for re-mapping the same dataset to different versions of the same reference genome (e.g. hg19 vs. GRCH38 vs. CHM13), again aiding computational efficiency (Figure 1).

CNV2SV: supplement CNV calling in SV detection. CNVs are a subset of SV consisting of deletions and duplications. One common challenge is mapping CNVs to specific SV types. Automatically linking these events together will improve our understanding and interpretation of regions where large CNVs occur and potentially also lead to improvements in breakpoint accuracy and thus resolution if the breakpoints are more complex (i.e. including other SV events, e.g. inversions). Here, we demonstrate CNV2SV using the haploid human genome of T2T CHM13 and will showcase CNV2SV on additional diploid, publicly available samples (Figure 2).

CoronaSV: SV pipeline for SARS-CoV-2. While deletions have been reported in several SARS-CoV-2 genomes at consensus level,14–17 the confidence in how these deletions are detected has not yet thoroughly been evaluated. Existing methods for detecting SV at the individual read level often suffer from false positive calls.16,17 Additionally, analyses with different variant calling pipelines often result in inconsistent calls.16,19 To examine the landscape and extent of SV across SARS-CoV-2 genomes, a method for generating accurate and trustworthy SV calls is needed. With this in mind, we developed the CoronaSV bioinformatics pipeline.

CoronaSV is a SV detection and SV validation pipeline for SARS-CoV-2 that combines an ensemble SV calling approach that relies on both long read and short read sequencing technologies (Figure 3). Both assembly-based and read based SV detection methods are used by CoronaSV. By combining different sequencing technologies and variant detection approaches, we can identify both a) confident SV call set and b) artifacts that may result from specific technologies + computational approaches.
CleanSV: Filtering High-quality SVs. Short-read sequencing is performed within clinical genomics to both inform and directly guide patient care. This has been immensely successful for various Mendelian disorders, where patients now routinely have their genomes sequenced to detect high-quality variants. Indeed, this approach has been utilized within clinical genomics for close to a decade, often to correct misdiagnoses (see for a recent example).

Within precision oncology, short-read sequencing (normally targeted sequencing or whole exome sequencing (WES)) has proved successful not only illuminating the nature of specific cancers, but also guiding novel drug development. Today routine sequencing is used to apply therapies for specific cancer subtypes, and influence the treatment of individual patients. For clinical work, samples from tumors are sequenced for somatic variants in well-studied oncogenes/tumor suppressor genes. Bioinformaticians will then manually investigate the variant calls within IGV in order to validate how accurate they are, and finally send reports summarizing these data to clinicians.

However, SV calling using short-read data is marred by high false positive (FP) call rates, sometimes up to 90% with modern callers. As a result, manual curation for each patient proves oppressively time-consuming for the needs of modern precision care and is prone to human error. Even though aneuploidy has been long studied for its role in tumor progression (see for a recent review), due to algorithmic uncertainties, routine inclusion of high-confidence SVs within clinical reports is often infeasible today.

Therefore, there exists a pressing need within bioinformatics to develop methods to remove false positives from the outputs commonly used SV callers, and benchmark their performance across a variety of assays (including a range of sequencing depths and tumor purities). Individual SV callers rely upon specific strategies to detect SVs, which makes the nature of the false positives algorithm-specific. Having access to a call set with a lower false positive rate would certainly not eliminate the requirements of manual curation, but it would make the problem more tractable.

The goal of this project was to develop a set of publicly available filters tailored for cancer genomics which have been measured to perform reliably across popular SV callers, as the filters must be specific to the SV caller used. Using a large cohort of high-quality normal whole genome sequencing (WGS) samples, we perform systematic false positive filtering. SVs labeled by the algorithm as somatic have evidence as actually being germline, while others are algorithmic artifacts. With such filters, bioinformaticians would have access to a set of high-quality somatic calls to manually curate, which could finally result in more robust clinical reports.

Sniphles: Phasing SVs with parallel programming. Phasing infers the correct cis or trans relationship between different heterozygous variations facilitating accurate haplotype reconstruction. Protocols and programs utilizing molecular phasing (chromosomal separation at the bench before sequencing), pedigree-based phasing (mapping parental and offspring genotypes to understand the haplotype), population-based phasing (using genotype data from large cohorts to infer haplotypes), and read-based phasing (mapping sequencing reads with the same variants to construct a haplotype) are all successful approaches to phasing next-generation sequencing data. The long-reads of third generation sequencing have bolstered our ability to phase longer and more comprehensive haplotype blocks. More comprehensive haplotype blocks increase our ability to accurately phase structural variants.

The goal of this project is to develop a wrapper script around the Sniffles SV caller to properly phase SV and augment the ability of Sniffles to accurately call SV (Figure 5). This result is obtained by using phased reads generated by SNV phasing tools such as WhatsHap or LongShot, and subsequently call SVs on the haploid phase blocks separately using temporary files before finally merging both haplotypes to obtain a single VCF file. As this algorithm processes each phase block separately this is attractive for parallelization. Our wrapper script additionally makes Sniphles compatible with alignments in the CRAM format.

Swagg: Structural Variation With Annotated Graph Genomes (Swagg). Most graphical approaches to variant calling only use genome graphs. While this information helps illustrate variation on a genomic level, it does not show variation on the individual protein level. To help leverage the power of graph approaches for SV calling, we introduce a pipeline that delivers both protein and genome graphs.

Swagg is a pipeline that enables the construction of genome graphs from read data (Figure 6). The input into the pipeline is sequence reads with or without a reference genome(s). Reads can be short reads or preprocessed (basecalled) long reads. These reads are then assembled into longer contigs which are mapped back to the reference genome to highlight discrepancies with the reference genome. These discrepancies can be caused by real mutations or sequencing artifacts and easily identified using SV tools, which output VCF files for each read set. These VCF files are taken together to make the genome graph at the end of the pipeline. The overall pipeline and intertwined modules are shown below. In addition to the
pipeline for creating graph genome and graph proteins, we also have a module for simulating reads based on an input reference genome.

PanOriginSV: detecting lab-of-origin. The advent of novel synthetic biology methods and organic bench-top synthesis toolkits like CRISPR has enabled rapid developments in genetic engineering. However, this progress has also introduced biosafety concerns surrounding the intentional or unintentional misuse of these tools. In order to increase accountability, the lab-of-origin studies attempt to map a set of plasmids to their lab-of-origin. Subsequently, the Genetic Engineering Attribution Challenge (GEAC) was announced, inviting open source tools from the community that could best predict the lab-of-origin.

Previous methods have employed machine learning or deep learning-based approaches that despite their promise, suffer from sub-optimal accuracy, long training times as well as explainability issues. Recently, a new alignment based tool PlasmidHawk reported higher accuracy than machine learning tools. PlasmidHawk relies on linear pangenome constructs to align query sequences to a pangenome in order to best determine the Top-1 and Top-10 candidate labs. Though PlasmidHawk has a higher accuracy, the runtimes to create the linear plasmid are non-scalable to larger datasets. Another drawback being the linear pangenome doesn’t incorporate SV, which could be important to predict hard-to-classify sequences. To address some of these challenges, we propose a tool PanOriginSV that combines machine learning approaches with graphical pangenome based alignment to predict lab-of-origin (Figure 7).

PanOriginSV creates multiple pangenome graphs from similar training sequences using BCALM creating a variation graph that incorporates SV and aligns the sequences back to the graph using GraphAligner. The most similar training sequences for graph construction are clustered using MMSEQ2. After this, top alignments, scores to the pangenome and sequence metadata are considered as features for a downstream machine learning model towards lab-of-origin prediction.

GeneVar: SV Browser. Next-generation sequencing provides the ability to sequence extended genomic regions or a whole-genome relatively cheaply and rapidly, making it a powerful technique to uncover the genetic architecture of diseases. However, significant challenges remain, including interpreting and prioritizing the identified variants and setting up the appropriate analysis pipeline to cover the necessary spectrum of genetic factors, which includes expansions, repeats, insertions/deletions (indels), SV and point mutations. For those outside the immediate field of genetics, a group that includes researchers, hospital staff, general practitioners, and increasingly, patients who have paid to have their genome sequenced privately, the interpretation of findings is particularly challenging. Although various tools are available to predict the pathogenicity of a protein-changing variant, they do not always agree, further compounding the problem. Furthermore, with the increasing availability of next-generation sequencing data, non-specialists, including health care professionals and patients, are obtaining their genomic information without a corresponding ability to analyse and predict lab-of-origin studies attempt to map a set of plasmids to their lab-of-origin. Subsequently, the Genetic Engineering Attribution Challenge (GEAC) was announced, inviting open source tools from the community that could best predict the lab-of-origin.

Here we present GeneVar, an open access, gene centric data browser for SV analysis (Figure 8). GeneVar takes as input a gene name or ID and produces a report that informs the user of all SVs overlapping the gene and any non-coding regulatory elements affecting expression of the gene. The tool is intended to have a clinical focus, informing the interpretation of SV pertaining to a gene name provided by the user.

SVTeaser: simulated data for SV benchmarking. SV detection tools often have a large number of wrongly detected variations requiring benchmarking to assess method quality before finalizing a workflow. Few tools are currently available to simulate data for SV benchmarking. SVTeaser is a tool for rapid assessment of SV calling fidelity with two main use-cases: 1) genotyping a set of pre-ascertained SVs and 2) benchmarking a new algorithm against pre-existing tools across a range of parameters. Users simply supply SVTeaser with a reference sequence file (.fasta) and, optionally, a set of SVs (.vcf). SVTeaser outputs simulated reads across a range of read lengths and depths and provides a downstream dataframe based analysis framework for evaluating accuracy (Figure 9). SVTeaser achieves rapid assessment by downsampling the full reference to a subset of numerous 10kb samples to which it adds SVs.

XSVLen: haplotype-resolved assemblies for benchmarking SVs. Since the development of a “gold standard” SV set, sequencing technologies and assembly algorithms have improved to enable nearly complete haplotype-resolved assemblies of human genomes. XSVLen is a framework (Figure 10) to use haplotype-resolved assemblies for benchmarking SV detection algorithms. Each variant call may be considered an operation to be applied to the reference genome. Our framework for benchmarking SV callsets is to apply SV operations to the reference genome and compare the modified
reference against the haplotype-resolved assemblies. This approach allows for SV calls that are different but produce similar sequences due to the repetitive nature of the genome to be scored as valid. In this manner, all variants in a region that is accurately assembled in both haplotypes may be benchmarked using this approach. We demonstrate the effectiveness of this approach by scoring SV calls generated from Oxford Nanopore reads on the HG002 genome and comparing against gold-standard calls by Truvari (https://github.com/spiralgenetics/truvari). This approach can be extended to use any haplotype-resolved assembly to benchmark SV callsets in additional genomes, enabling benchmarks as a distribution across call sets.

**Methods**

**Implementation**

**NibbleSV:** NibbleSV is a lightweight, scalable and easy to apply method to identify the frequency of SV events across short read data sets. As such, NibbleSV extracts kmers that are informative if an SV is present or if an SV is absent given the breakpoints of the previous predicted SV. Subsequently, NibbleSV scans the short read bam or fastq file for the presence of these k-mers and counts their number of occurrences. In the end, NibbleSV extends the VCF file with tags holding information about the number of times an SV is supported by kmers or not (Figure 1).

**CNV2SV:** CNV2SV is a tool developed to identify CNVs in the context of a whole genome sequence (Figure 2). CNV2SV requires three input files which is a BED or VCF formats (from Parliament, Control-FREEC, a putative SVs from genome-genome alignment (vcf format, from dipcall) and assembled reference files (fasta files) from both the reference and the assembled sample. In addition, we require preinstalled packages such as intavertree, mappy, pyfaidx together with python v.3.8.* or higher. The visualisation for CNV2SV requires the installment of R together with the circularize package, matplotlib, seaborn and pandas. CNV2SV currently relies on an installation of Control-FREEC which is needed for the CNV calling from short reads. The main output of CNV2SV (cnvlink.py) comprises the best matching linked SV for each CNV call, with summary statistics including alignment overlap, mismatches, respective start and end positions useful for evaluating e.g. the quality/resolution of breakpoints identified by the CNV callers. Furthermore, all additionally identified adjacent and distant SVs are reported separately for each CNV. The raw output can be further visualized to show the CNV-SV links identified across the genome in a circular plot, as well as summary statistics for the linking results (Figure 2).

**CoronaSV:** CoronaSV is a method developed for generating accurate and trustworthy SV calls across SARS-CoV-2 genomes. The tool utilises available SRA data from SARS-CoV-2 isolates that have been sequenced with both Illumina and ONT platforms. CoronaSV utilizes a combination of three different approaches: read-based SV detection with paired-end Illumina reads and ONT long-reads, as well as assembly-based SV detection using both short and long-reads (Figure 3). All the software packages used by CoronaSV can be installed via the Conda package manager (https://github.com/conda/conda).

**CleanSV:** The goal of the hackathon project was to develop filters and QC checks to remove false positive calls from common SV callers. Currently, within clinical genomics, it’s exceptionally difficult to categorize true positives from false positives, thus making accurate diagnoses virtually impossible. The situation is even more complicated within clinical oncology, as researchers need to precisely separate true somatic calls from false positives and (potential) germline calls. In order to aid with precision SV calling, the team wrote a set of scripts to be used with short-read SV callers, allowing researchers to better generate a set of high-quality SVs to further investigate manually (Figure 4).

For cancer genomics, groups normally develop in-house filters to improve the precision of SV calling. The scripts developed for this project accept as input GRIDSS, Manta, DELLY, and SvABA calls from short-read WGS data, along with a manually-curated reference set of calls designated as ground truth. The reference set was curated using a paired melanoma and normal lymphoblastoid COLO829 cell lines using four different technologies (Illumina HiSeq, Oxford Nanopore, Pacific Biosciences and 10x Genomics), along with extensive external validation. Using the reference set, we proceeded to investigate the presence and nature of false positives from the initial callsets. (Note that we focused on WGS for this hackathon, but a similar approach could be applied to other assays such as WES.) Samplot visualization of read data allowed manual curation of parameters associated with FP calls, and associations between AnnoTVS and FPR helped identify additional FP-associated SV parameters (Figure 4). Along with the manually-curated reference set, the panel of normal (PON) used for further filtering was generated from a compiled set of high-quality germline calls using 3,782 normal samples freshly-sequenced at a median depth of 38x by the Hartwig Medical Foundation.

**SnipHles:** The main idea is to phase the identified SVs. We use two approaches; the first is to extract the tagged reads from the bam file and use these reads to phase the SVs if not conflicted. The second approach is to split the haplotype bam file
based on the haplotype tag, using each split bam file to call SVs separately, this method called (Snipphes). Snipphes utilize information impeded in haplotypes, bam file and reads info support SVs. This method phases structural variants and augments the ability of Snipphes to accurately call SVs (Figure 5). Snipphes is implemented in Python 3, and it takes a haplotyped bam, and a SV VCF file as input and produces a phased vcf file as output.

**Swagg:** Structural Variation with Annotated Graph Genomes (SWAGG) is a pipeline to make genome graphs from read data. The input into the pipeline is reads either with or without reference genome(s). Reads can be short-reads or preprocessed (basecalled) long-reads. Reads are assembled into longer contigs, and contigs are mapped back to the reference genome to look for discrepancies with the reference genome. These discrepancies can be either real mutations or sequencing artifacts, and are found using structural variant tools which output VCF files for each read set. These VCF files are taken together to make the genome graph at the end of the pipeline (Figure 6).

**PanOriginSV:** This tool is a lab-of-origin prediction tool that combines machine learning approaches with graphical pangenome based alignment to predict lab-of-origin (Figure 7). PanOriginSV is implemented in Python 3 and uses the scikit-learn package for deploying machine learning models. PanOriginSV also relies on MMSEQ2 for clustering, BCALM for graph construction and minigraph for graph alignment. Given a training set of engineered plasmids and their source labs, this software can predict the lab of origin of a test set of plasmids.

**GeneVar:** The GeneVar tool was developed to help inform the clinical interpretation of structural variants pertaining to a user-provided gene. This software is an open access, gene-centric data browser for SV analysis. GeneVar is a web page application (Figure 8). After entering the gene name (HGNC, Ensembl gene (ENSG), or transcript (ENST) identifier) in the search box on the homepage, the user is directed to the summary of the gene-specific page. GeneVar is available on GitHub (https://github.com/collaborativebioinformatics/GeneVar). The repository provides detailed instructions for tool usage and installation. A bash script for automated installation of the required dependencies is also provided.

**SVTeaser:** SVTeaser is a tool for rapid assessment of SV call fidelity created for geneticists designing experiments to genotype a set of pre-ascertained SVs and bioinformaticians benchmarking a new algorithm against pre-existing tools across a range of parameters (Figure 9). Users are required to supply SVTeaser with a reference sequence file (.fasta) and, optionally, a set of SVs (.vcf). SVTeaser outputs assorted statistical metrics across a range of read lengths and depths. SVTeaser achieves rapid assessment by downsampling the full reference to a subset of numerous 10kb samples to which it adds SVs.

**XSVLen:** This software is a framework for haplotype-resolved assemblies for benchmarking SV detection algorithms (Figure 10). XSVLen takes a cuteSV or Sniffles VCF output file and using reference coordinates will produce modified sequences having included inserted sequences or deleted sequences within the reference sequence. By creating these modified sequences, we could check for the presence of the predicted variants in haplotype-resolved assemblies. All methods are open-source licensed and have been made available on GitHub: https://github.com/collaborativebioinformatics.

**Operation**

**NibbleSV:** NibbleSV requires a reference genome and VCF file that includes all the SV that should be genotyped (Figure 1). Next, allele kmers for the reference and alternative are extracted. The extraction process includes each site’s flanking regions. Subsequently, the occurrence of these k-mers in the reference fasta file are counted. This step is necessary to prevent k-mer miscounting between reference vs. alternative allele. To enable scaling of NibbleSV for large data sets, the results of these two steps are written into a temporary file, which is all that is needed for the actual genotyping step. During the genotyping step, NibbleSV uses this small temporary file and the bam/cram file of the sample and identifies the presence/absence of the reference and alternative k-mer across the entire sample. This is very fast and requires only minimal resources of memory as the number of k-mers is limited. On completion of NibbleSV a scanning of the bam/cram file is carried out reporting which SV have been re-identified by adding a tag in the output VCF file of this sample (Figure 1). The VCF per sample can then be merged to obtain population frequencies. The VCF per sample can then be merged to obtain population frequencies. NibbleSV requires 4Gb of memory, a single core and around 2GB hard disk space to store its index from e.g. GIAB HG002.

**CNV2SV:** An overview of the CNV2SV workflow is illustrated in Figure 2. CNV/SV are called from short read datasets using Parliament. Locations of duplications are determined using CNVnator individual calls and combined calls. Copy number estimation for regions across the genomes is implemented by Control-FREEC that is converted into BED and VCF files. The genomes to be compared are aligned using dipcall, dot plots of the alignment constructed to identify potential regions of interest and extracted to VCF output. Calls for both short read and assembly based CNV/SV calls are merged to locate regions in which the duplication events overlap. Visualization scripts will be made available in future.
versions. Here, each called CNV from the short read dataset is compared to SVs identified from the genome-genome alignment. This is achieved in two steps: First, CNVs are queried against an interval tree structure containing the SVs from the genome-genome alignment to find adjacent CNV-SV pairs (<1000bp apart, putative tandem duplication events). For CNVs for which no matching SV can be identified in this way, the search is then extended to the whole genome (putative translocation events). All potential CNV-SV links are evaluated by sequence alignment using mappy (Python binding for minimap2\(^\text{54}\), with a standard sequence identity threshold of 0.8. A quick-start example using CHM13 and GRCh38 data is available on our GitHub page. In addition, we hosted a detailed description of the output data on the GitHub page.

System requirements (see GitHub for more information): CNV2SV has been tested to work on a desktop system on the CHM13 data set with an Intel® i7-6700K Processor (4.00Ghz quad-core), 32GB RAM (less may be required), 50GB free disk space and running Unix-like operating system (e.g. Ubuntu-based distribution) or Windows subsystem for Linux running Ubuntu. The initial genome-genome alignment (CHM13 vs GRCh38) was computed on a cloud-based platform (DNANexus). CNV2SV requires Python (3.8 or newer). A full list of package dependencies is available on the GitHub page.

CoronaSV: All software packages used by CoronaSV can be installed via the Conda package manager. Additionally, the CoronaSV workflow is defined using Snakemake. Running the CoronaSV.smk snakemake pipeline handles downloading all specified data and processing of sequencing data to variant calls. Each step of the CoronaSV pipeline (Figure 3) has a defined conda environment with exact versions of software specified for easy installation. CoronaSV utilizes three approaches that includes 1) read-based SV detection with paired-end Illumina reads, 2) ONT long-reads and 3) assembly-based SV detection. Illumina paired-end short-reads are trimmed using trimmomatic\(^\text{55}\) and mapped to the SARS-CoV-2 reference using bwa mem.\(^\text{56}\) After mapping, PCR duplicates are removed with Picard MarkDuplicates

Figure 1. Overview of the NibbleSV software package workflow. Overview of NibbleSV workflow that utilises an input reference genome and file containing variant calls to generate a list of genotyped alleles using a kmer based strategy.

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\(54\) https://pypi.org/project/mappy/

\(55\) https://bcbio.readthedocs.io/en/latest/

\(56\) https://picard.readthedocs.io/en/latest/
CNV2SV: Copy Number Variant detection in the context of a full genome

Figure 2. A graphical overview of the CNV2SV pipeline. CNV2SV software pipeline that utilises both short and long read data as input to calculate the frequency of copy number variants across complete genomes.

Structural variants are identified then using Delly, Manta, Lumpy, and Tardis. Nanopore long-reads are filtered using Nanofilt and mapped to SARS-CoV-2 reference using minimap2 with default parameters. SVs are then called using Sniffles, SVIM, and CuteSV. Read quality assessment is carried out by NanoPlot. In order to integrate assembly based methods, de novo SARS-CoV-2 assemblies were generated.
using Unicycler for short-read sequencing and Flye for ONT long-reads. NucDiff and SVanalyzer tools are used for assembly-to-assembly comparisons. Followup comparative analyses across callsets is implemented by SURVIVOR\textsuperscript{59} (Figure 3).

System requirements: CoronaSV is tested on Linux-based systems with multiple illumina and nanopore sequencing data (see GitHub for full list of the testing data). The RAM usage of CoronaSV depends on the size of input data. Peak RAM usage appears during de novo assembly using Unicycler, and 16 GB of RAM is sufficient for the pipeline to run on 8 CPU cores with additional 50Gb of disk space. CoronaSV requires python (version 3.6 or newer) and snakemake. Required tools and package dependencies can be found on GitHub page.

**CleanSV:** The methods adopted to construct the CleanSVs filtration protocols are shown in Figure 4. In order to generate the data required to develop adequate filters, structural variants (SVs) were called on Illumina short reads using novoalign hs37d5 HG002 BAM (ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/technical/reference/phase2_reference_assembly_sequence/hs37d5.fa.gz) and Illumina short-read HiSeqX Ten hg19 COLOR29 BAM (https://nextcloud.hartwigmedicalfoundation.nl/s/LtKd8XxBqwaIC?path=%2FHMFTools-Resources%2FGRIDSS-Purple-Linx-Docker) using GRIDSS,\textsuperscript{42} Delly,\textsuperscript{44} and Manta.\textsuperscript{43} In HG002, the SV truthset (6) was used to determine the false positive (FP) SV calls in the short read dataset. Any SV calls that were found outside of the truthset Tier I bed regions were then filtered. In the sample COLOR29, the SV truthset (2) was used to determine the FP SV calls in the short-read dataset. Calls were inspected through manual curation by samplot (7). Generated samplots were annotated with UCSC table browser repeat tracks (8) and converted using vcfanno (9) as well as GC content by bedops file conversion from wig to bed (10).
We then compiled a set of high-quality germline calls using 3,782 normal samples freshly-sequenced at a median depth of 38x by the Hartwig Medical Foundation. We initially hypothesized that such a large cohort could be used to both perform systematic FP filtering and possibly detect calls simply incorrectly labeled as somatic. The calls were filtered if a match within 2bp of the breakpoint was found in the PON. System requirements: The scripts to filter SV calls with either VCF or BEDPE format require R version 3.6.0 or higher, which is available for Linux, Mac OS, and Windows.

Figure 4. An illustration of the approach used by CleanSV to generate and implement filtration of SV calls. CleanSV pipeline highlighting methods used to generate adequate filters that can be utilised by clinicians to filter false positive and mislabeled SV calls from short-read cancer datasets.
The analyses within were run on R version 4.0.3, with Biocondcutor version 3.12. For running SV callers, it is recommended to use a HPC environment on Linux.

**Sniphles:** The Sniphles workflow (Figure 5) requires the following dependencies: Python >= 3.6, Pysam (Version 0.16.0) ([https://github.com/pysam-developers/pysam](https://github.com/pysam-developers/pysam)), Cvcf2 (Version 0.30.2), Sniffles (Version 1.12), SURVIVOR (Version 1.0.7), Mosdepth (Version 0.2.6), Bcftools (Version 1.9), tabix (Version 1.8). The workflow partitions reads in the bam file into groups based on phase blocks and phase status, which enables parallel analysis of the data. The read coverage at each block and each phase is computed with Mosdepth and used to estimate the parameters for calling SVs by Sniffles. Next, the identified SVs per haplotype are concatenated using bcftools. SVs of two haplotypes are combined using SURVIVOR with option "1000 1 0 0 0" to merge SVs within 1 kbp between each other and to allow for different types of variants to be considered on different haplotypes. SVs are then force called with Sniffles using this combined vcf file. Force called SVs from each haplotype are combined with SVs of unphased regions as the final output (Figure 5). To facilitate workflow testing, Princess ([https://github.com/MeHelmy/princess](https://github.com/MeHelmy/princess)) was used to align, detect and phase SNVs and SVs from PacBio HiFi reads. The produced Bam from the previous step is the input for Sniphles, where pysam was used for alignment.

**Swagg:** The minimal system requirements for SWAGG are 8Gb RAM, 1 CPU and 10Gb of storage. Figure 6 demonstrates the implementation and operation of the SWAGG software package. Protein graphs are generated using a multiple sequence alignment of the proteins, then using the tool msa_to_gfa ([https://github.com/fawaz-dabbaghieh/msa_to_gfa](https://github.com/fawaz-dabbaghieh/msa_to_gfa)) after which this multiple sequence alignment is converted into a graph file in GFA format, with the original sequences embedded as paths in the graph for visualization. This tool is tested with Python 3 and does not require any extra libraries or dependencies. New sequences can be aligned to these graphs using Partial Order Alignment algorithm for example.
PanOriginSV: PanOriginSV has three additional open source dependencies which are MMSEQ2 (for clustering), BCALM (pangenome creation) and GraphAligner (for sequence-to-graph alignment). MMSEQ2 is the most memory intensive step, and MMSEQ2 requires roughly 1 byte per sequence residue. An overview of the pipeline utilised by this tool is highlighted in Figure 7. PanOriginSV performs lab-of-origin prediction in three distinct steps. Firstly, during the training phase PanOriginSV clusters similar sequences using MMSEQ2. Further, the most similar clusters having a predefined number of representative lab labels are selected for the pangenome creation. Second, in order to incorporate SV information into the pangenome a graphical pangenome is created using BCALM for each of the clusters identified in the first step. These pangenomes reflect sequence-level structural variation that reveal important differences in highly similar sequences that could belong to different labs thereby reducing the possibility of false positives. The training sequences are then mapped back to their corresponding pangenome graph to obtain important alignment information including but not limited to number of hits and percentage identity of alignment. These are then collated and embedded into a feature vector that is passed on to a machine learning model for prediction.

Figure 6. Outline of the SWAGG software package workflow. Illustration of the SWAGG pipeline that utilises both long and short read datasets for the construction of graph genomes.
Thirdly, features from the alignment steps are combined with sequence metadata and input into a random forest classifier that is trained to predict lab-of-origin in a multiclass classification task. Both the top-1 and top-10 predictions are output and compared to previous literature and Genetic Engineering Attribution Challenge (GEAC) benchmarks.

**GeneVar**: Figure 8 shows the different components of GeneVar, which is a web browser application. The webpage, including data storage, requires only one core with 1 Gb RAM and requires less than 1 Gb of storage. After entering the
gene name (HGNC, Ensembl gene (ENSG), or transcript (ENST) identifier) in the search box on the homepage, you will be directed to the gene-specific page containing: 1) Gene-level summary with number of SVs, number of clinical SVs or SVs overlapping clinical SNVs, 2) Links to the gene’s page on OMIM, GTEx, gnomAD, 3) A dynamic table with the annotated variants overlapping the gene, 4) A graph with the distribution of the allele frequency for variants matched with gnomAD-SV (50% reciprocal overlap). The profile of the SV to consider, such as type and size range, can be specified on the side bar. Each column in the dynamic table can be “searched” into or reordered dynamically. All data used by the app will be available for download in tab-delimited files. By default, allele frequency is reported based on dbVar and gnomAD genomes and exomes. Furthermore, GeneVar utilises dbVar database and links SV to genes and annotate gene impact, allele frequency, and the overlap with clinically-relevant SVs, SNVs and indels. All data, are available for download in a tab-delimited file. Each variant has been extensively annotated and aggregated in a customizable table. 

GeneVar is available on GitHub (https://github.com/collaborativebioinformatics/GeneVar). The repository provides detailed instructions for tool usage.

**SVTeaser:** SVTeaser generates regions from a user provided reference and adds in a structural variant into each region using one of two methods - 1) a call to SURVIVOR simSV which generates random, simulated SVs by introducing variation (deletions (DEL) and insertions (INS) type of SV breakpoints) in DNA sequences, or 2) automatic spike-in of a known SV from an input SV VCF file. Resultant altered reference sequences are then used for Illumina short-read simulation using ART. Parameters controlling simulated sequencing read-length, insert-size, and depth parameters can be altered. Simulated reads can then be mapped to the original, unaltered reference with any mapper of choice; here, BWA was used. Resultant BAM files can then be used to detect SVs using any mapping-based SV caller of choice; here, Parliament was used to generate calls with manta, breakseq, cnvminer, and lumpy. The resultant VCFs are then matched to the simulated SVs VCFs using Truvari and output is parsed into a pandas dataframe for report generation. SVTeaser requires installation of Python 3.7, Truvari, SURVIVOR, Vcftools and ART read simulator. The components of SVTeaser are shown in Figure 9.

**XSVLen:** The XSVLen workflow requires Python3, Minimap2, Nextflow, and R >=3.5.0. As demonstrated in Figure 10, XSVLen takes as input a haplotype-resolved de novo assembly, and a VCF file (generated by cuteSV or sniffles) of variants including only insertion and deletion calls. For each insertion or deletion call within the vcf file, a modified reference genome is generated. This modified reference will contain a 1.5kb flanking sequence that either has the sequence removed if a deletion call, or the alternate sequence added between the flanking sequences if an insertion call. The resulting ‘query’ sequences are then mapped using minimap2 to both haplotypes. Each aligned query gives rise to a map of aligned bases $P=(q_1,t_1),\ldots,(q_n,t_n)$. To score each variant call, we find two indexes i, j. These index the end of the prefix, and beginning of the suffix in the query. When the call is valid, $(P[j][i] - P[j][0]) - (P[j][1] - P[j][0])$ is equal to 0. To account for differences in alignment, we iteratively search for an $(i',j')$ combination, with $i' \leq i$ and $j' \leq j$ that gives the smallest difference. Variants are reported as valid if the difference is less than 10 bases or the intervals
defined between P $^{[iopt]}$ and P $^{[jopt]}$ are within 95% length in either haplotype. A summary report is then produced using an R script.

**Use Cases**

**NibbleSV:**

We benchmarked NibbleSV over the GIAB HG002 SV call set\(^6\). In summary, this call set was created using multiple long and short read technologies and underwent manual validation across multiple groups to ensure an overall high

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**Figure 9.** An illustration of the methodology implemented by the SVTeaser software package. SVTeaser software implementation showing how by using a reference genome a set of SVs can be simulated to benchmark SV calling tools to inform experimental design decisions prior to analyses.
quality and accuracy. Using an Illumina data set from (2x250 GIAB HG002) we benchmarked true positives (i.e. SV that should be present), false positives (i.e. parental only SV that should not be present in the proband HG002), and false negatives (i.e. SV that should be present in HG002 but were not found). Using only chr 22 from HG002, NibbleSV with a kmer size of 23 takes around 2-4 minutes on a single thread with a 80gb bam file and the provided VCF file. We assessed our recall at different k-mer sizes which increases with the kmer size. For example, k=21 (2min 3sec) achieves 0.59 recall with a precision of 0.83. Interestingly, for insertions the recall rate increases to 0.86 with a precision of 0.86.

**CNV2SV:**

Figure 11a shows the best links (based on the alignment identity score) between CNVs identified by CNVnator and duplication SVs inferred from the dipcall alignment of CHM13 and GRCh38. The genomic areas surrounding four selected adjacent duplication events are further highlighted using dot plots, revealing the architecture of the corresponding variation in the context of the genome-genome alignment. In Figure 11b, all CNV-SV links meeting a default alignment identity threshold are shown, further revealing events corresponding to putative copy number increases of greater than two. We further explored the reason why some CNV and SV could not be linked, through statistical analysis of the raw SV calls as shown in Figure 12.

**CoronaSV:** We processed more than 200 SARS-CoV-2 SRA runs with CoronaSV, and Figure 13 shows the high confidence SVs generated with SURVIVOR by taking a majority vote across multiple SV callers. We also looked for shared SVs across multiple samples. There were only a few inversions identified that were consistently and reliably called between samples. We believe those inversions are related to the transcriptional landscape of SARS-CoV-2. These inversions are small (less than 1Kb), and five of them were found in ORF1ab and one on ORF M.

**CleanSV:** We investigated filtering SV calls using both the curated reference set and a high-quality panel of normals (PON). The PON was created using GRIDSS calls from 3,782 normal samples freshly-sequenced at a median depth of 38x by the Hartwig Medical Foundation. Using this PON consisting of GRIDSS calls from 3,792 freshly-sequenced normal WGS samples, we explored how the percentage of calls from short-read SV callers which were incorrectly labeled as true somatic calls, either due to being algorithmic artifacts or germline calls. Our results show the promise of such an approach. (Figure 14). Note that this is not only a check for false positives: we know a priori that many calls from somatic SV callers are mislabeled as somatic when they’re actually germline. This is a known algorithmic error: SVs are normally first called in the normal sample and labeled as “germline”, and then the resulting SVs called using the tumor sample (separately or jointly with against the normal) are labeled as “somatic”. While such an approach is common for short-read SV callers, this frequently leads to mislabeled results, often for the simple reason that the normal sample is sequenced at a much lower coverage than the tumor sample.

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**Figure 10. Implementation pipeline of XSVLen software package.** A graphical representation of the XSVLen software pipeline showing the utilisation of haplotype-resolved de novo assembly and a VCF file to benchmarking SV detection algorithms.
We plan to continue to explore whether the false positives found exhibit distinct features which we could use for future filters to distribute to the community. With these insights, given that clinical genomics still overwhelmingly relies upon short-read sequencing, our goal would be to also apply filters to all variants (e.g. the case of Mendelian diseases).

Figure 11. Relationship between the CNV calls and identified SVs across the CHM13 genome. CNV2SV results using CHM13 in comparison to GRCh38. (A) The diagram connects the genomic location of individual CNV calls on GRCh38 (broad ends) to the location of their linked SV identified in the GRCh38-CHM13 genome-genome alignment (thin ends). The diagram reveals a number of underlying putative translocation events identified for the called CNVs. Adjacent CNV-SV links (putative tandem duplications) are shown as streaks at the respective genomic position. For four select CNV-SV links, the genome-genome alignment for the underlying SV is further shown as a dot plot on the outside of the diagram. Only the best matching SV link (thin end) identified for each CNV call (broad end), as determined by sequence alignment score between both events, is shown here. (B) Similar to A, but displaying all potential CNV-SV links that meet the default alignment identity threshold of 0.8. CNVs with multiple matching duplication SV events identified in the genome-genome alignment can be explained by copy numbers greater than two occurring in distant locations, and alternatively may suggest the involvement of complex genomic rearrangements including transposable elements.
Sniphles: We used Princess (https://github.com/MeHelmy/princess) to align, detect and phase SNVs and SVs from PacBio HiFi reads 32x coverage. The produced Bam from the previous step is the input for Sniphles, where pysam (https://github.com/pysam-developers/pysam) was used for alignment. For each phase block we used the mosdepth61 to

Figure 12. Summary the linkage statistics showing the characteristics of disparities between the CNV calls and SVs. In terms of linkage statistics, the majority of the CNVs identified have not been linked to a SV event, as indicated by A. One of the main reasons for the unsuccessful linking is the length disparity between the called CNV events and SV events as shown in B. Overall, among linked CNV and SV events, the distribution of three major categories are shown in C: adjacent events, distant events, and events spanning multiple chromosomes. Distant events are called in the case when the linked SV is at least 1Kbp away from the CNV call (either upstream or downstream). Details of the distribution of the adjacent and distant events per CNV call are given in D and E shows that alignment quality is better for adjacent matches when compared to more distant SV matches. While most links for a single CNV event are unanimously distant or adjacent, we observed an event in which a CNV was linked to both an adjacent and a distant SV which occurs on chromosome 7 (F: adjacent: chr7:1009997804 length 8325 and distant: chr7:100994092 length 3249).
detect coverage. Later, we called SVs using Sniffles\textsuperscript{10,39} with the adequate numbers of reads to support SV. The identified SVs per phase block were sorted and concatenated using bcftools version 1.967, and both the haplotypes were merged using SURVIVOR.

\textbf{Swagg:} The main results from the Swagg package includes the development of the graph module and the protein graphical application (Figure 15). The graph module is able to retrieve basic statistics from a pangenome graph in GFA format from either reference-based (fasta + VCF) or a set of assemblies (fasta), followed by conducting a pairwise comparison of all paths in the graph, and outputting a matrix in TSV format with the path names and corresponding samples in the first position. After creating the pairwise matrix, the module can plot an SV pileup over any path in the graph, counting the number of other paths that contain an SV overlapping each position. In addition to utilizing the pairwise comparisons where hotspots are references to a given sample, this approach also allows for graphs derived from

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure13.png}
\caption{An analysis of size and number of identified SARS-CoV-2 SVs. Histogram showing size of the SVs and the total number of SVs across multiple Nanopore and Illumina datasets. The y-axis of the histogram is log compressed.}
\end{figure}

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure14.png}
\caption{Filtering somatic SV calls using a Panel of Normals. Using a cohort of 3,792 freshly-sequenced WGS normal samples to create a Panel of Normals (PON) from GRIDSS calls and a set of curated calls from sample COLO829 to classify false positives, we discover that a sizeable number of false positives were found within the panel of normals, suggesting that these were miscategorized due to algorithmic errors. PON filtering based on GRIDSS calls is effective for the modern callers such as GRIDSS, Manta and SvABA which (partially) rely upon localized assembly.}
\end{figure}
vcf files. The objective from the protein graph mapping application was to show if the variants introduce new amino acids or stop a stop codon. Similarly, a pangenome graph can be constructed from DNA sequences as panproteome graphs can be built from different amino acid sequences of a protein. These graphs can then help visualize the variants between the sequences and show the paths each sequence take through the graph. Another layer of information can be added to the nodes, e.g. does a node represent a conserved or a non-conserved side, does a path divergence in the graph have any significant phenotypic characteristics, relating genome-wide association studies to these proteins graph68. Therefore, when aligning a new sequence to the graph, one can check the path the new sequence took in the graph and what information are related to this path.69 Figure 15 shows an example of an annotated graph of the Nucleocapsid Phosphoprotein in SARS-COV-2.

PanOriginSV: It became apparent that the quality and representation of the clusters was a main factor in prediction accuracy. To this end, we tested PanOriginSV on a range of clusters of at least 500 sequences and only considered cases where the test sequence had a training representative in the assigned cluster. The CPU time used by PanOriginSV was 10-50x less than the linear model (depending on the cluster). We observed a range of results, with the linear prediction model outperforming PanOriginSV by up to 5% in some clusters and PanOriginSV outperforming the linear model by up to 6% in others (Table 1). With deeper analysis of the input data, we hope to achieve better clusters and thus better prediction results with the graph model. It is also worth noting that our graph construction method can be improved using more recent pangenome graph construction tools.

GeneVar Databrowser. Upon querying a gene or transcript, the data browser will visualize a rare-variant burden test, allele frequency distribution, and variant level information for known SVs within dbVar. The data browser does not have a login requirement and integrates multiple public resources (Figure 16). To illustrate whether a particular gene/transcript or exon has been adequately covered to detect variation, the average depth of coverage is graphically represented. An additional panel shows gene expression levels across all general tissues included in GTEx71. Report summary. Analysis
results are enriched with information from several widely used databases such as ClinVar^72 and gnomAD,^73 as well as graphical visualization utilities integrated in the pipeline as part of GeneVar. Resulting variants are reported in a tab delimited format to favor practical use of worksheet software such as iWork Number, Microsoft Excel or Google Spreadsheets. All information can be downloaded in tabular form.

SVTeaser: SVTeaser was able to simulate SV data on average 14 min per sample when tested on chromosome two. SVTeaser output includes organized VCF results of true positive, false positive, and false negative from evaluated SV

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Figure 16. User interface of GeneVar data browser. A description of the elementary transcript details for the gene of interest. This includes the Ensembl transcript ID, Ensembl Gene ID, number of exons and genomic coordinates as described in the GRCh38 build.
callers. Furthermore, performance scores are reported alongside automated plots for quick visual evaluation of SV callers (Figure 17). SVTeaser is able to simulate sequencing for known deletions and insertions with various coverage, read length, and insert length. This enabled thorough evaluation for understanding the strengths and boundaries of the SV callers in question (Figure 18).

XSVLen: A diploid assembly of the MHC locus of HG002 from Nurk et al.\textsuperscript{74} was used to benchmark variants. The assembly has an N50 of 16.1 and 18.0 Mb per haplotype. Variants were called using 50-fold coverage of ONT reads using CuteSV version v1.0.8. The number of INS and DEL per SV size can be observed in Table 2. The number of INS/DEL overlapping with the haplotype-resolved assemblies are shown in Table 2 as well as a comparison of assembly-based calls and gold-standard Truvari calls.

**Figure 17. Summary report output of SVTeaser.** A report generated from benchmarking a real HG002 Manta 30x Illumina sample against GIAB Tier1 SVs. A) Counts of SVs by SVType and their intersection state with GIAB Tier1 benchmark SVs. B) Summary table of benchmarking performance. C) Proportions of SV intersection states with the benchmark by SV size bin.
Conclusion

The results of the 2020 Baylor College of Medicine/DNANexus hackathon described here represent novel work that pushes the field forward for human genome SV detection but also for Covid related research. Both are needed to further current findings about diversity and the complexity of organisms and their genotypes. To further facilitate this progress in a FAIR-compliant manner, 80 people came together from across the world in October 2020 completed 10 groundbreaking prototypes. Hackathons like these not only represent short bursts of prototype development, but are essential to form groups and communities, inspire communication across countries and research institutions, and form novel collaboration networks. As such, this year’s hackathon not only sparked the projects described here, but also highlighted the need for unified databases for SVs and other genomic features hosted on DNAnexus, Anvil, and other platforms as well as

Table 1. Summary: Insertion and deletion events are compared according to structural variant size and number and overlaps with 1) haplotyped assemblies and 2) Truvari callsets are counted.

| SVType | Insertions | | Deletions | |
|---|---|---|---|---|
| SVLen | <50 | >=50 | <50 | >=50 |
| Number of insertion/deletion calls by cuteSV | 20007 | 14942 | 38331 | 10926 |
| Number of these calls that overlap assembly contigs | 16054 | 17481 | 30494 | 6475 |
| Number of truvari TP/FP/no-call (NP) | 3351 FP 7200 TP 15974 NP | 2835 FP 4095 TP 80 NP | 880 FP 1925 TP 30196 NP | 1571 FP 2099 TP 298 NP |
| Number of assembly TP/FP/no-call (NP) | 5220 FP 10834 TP | 8979 FP 8502 TP | 23202 FP 7292 TP | 3444 FP 3031 TP |
larger standards and references (e.g. GIAB, UKBB). This is essential to ensure quality standards for benchmarking and comparability, which will further advance the science and medical research.

Rapidly switching to a completely remote hackathon that enabled increased international participation was made necessary by the COVID-19 pandemic. This allowed for an open science effort across an even more diverse population of individuals and professional backgrounds. That diversity made it possible to complete 10 projects, which spearheaded novel insights in the understanding of structural variants in humans, as well as COVID19 genome structure. More importantly, it led to new synergies among participants, an active online community, and new friendships across borders.

Data availability
Associated code is available at: DOI 10.17605/OSF.IO/ME62X

Data sources utilized:

NibbleSV: Genome in a Bottle, HG002, SV callset ([ftp://ftp-trace.ncbi.nlm.nih.gov/ReferenceSamples/giab/data/AshkenazimTrio/HG002_NA24385_son/PacBio_CCS_15kb/])

CNV2SV: CHM13 ([https://github.com/nanopore-wgs-consortium/CHM13#telomere-to-telomere-consortium]) and GRCh38

CoronaSV: Data sources available on [https://github.com/collaborativebioinformatics/coronasv]

CleanSV: hs37d5 HG002 BAM ([ftp://ftp.1000genomes.ebi.ac.uk/vol1/ftp/technical/reference/phase2_reference_assembly_sequence/hs37d5.fa.gz]), hg19 COLO829 BAM ([https://nextcloud.hartwigmedicalfoundation.nl/s/LTtKt8XxBqwaiC?path=%2FHMFTools-Resources%2FGRIDSS-Purple-Linx-Doctet])

SWAG: Data sources available on [https://github.com/collaborativebioinformatics/swag/blob/main/sample_manifest.tsv]

XSVLen: MHC locus of HG002 (74)

GeneVar: Data sources available on [https://github.com/collaborativebioinformatics/GeneVar]

PanOriginSV: genetic engineering attribution challenge (GEAC)
[https://www.drivendata.org/competitions/63/genetic-engineering-attribution/]

SVTeaser: All data utilized was based on simulations

Sniphles: Genome in a Bottle, HG002, SV callset ([ftp://ftp-trace.ncbi.nlm.nih.gov/ReferenceSamples/giab/data/AshkenazimTrio/HG002_NA24385_son/PacBio_CCS_15kb/])

Software availability

NibbleSV

Source code available from: [https://github.com/collaborativebioinformatics/nibSV]

Archived source code at time of publication:

License: MIT license

CNVSV

Source code available from: [https://github.com/collaborativebioinformatics/CNV2SV]

Archived source code at time of publication:

License: MIT license
CoronaSV
Source code available from: https://github.com/collaborativebioinformatics/coronasv
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CleanSV
Source code available from: https://github.com/collaborativebioinformatics/CleanSV
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XVSLen

Source code available from: https://github.com/collaborativebioinformatics/The_X_team

Archived source code at time of publication:

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Acknowledgements

The authors like to thank Richard Gibbs, Aaron Wenger and Stephen Rudd for their helpful discussions during the hackathon. In addition, Gerald Wright and Brenton Pyle for helping with advertisement and organising. DNA nexus provided computational resources for the hackathon. The authors would also like to thank Oxford Nanopore Technologies and Pacific Biosciences for sponsored prizes.

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Loren L. Flynn
McCartney and colleagues describe a compelling and comprehensive account of the outcomes of an international hackathon to develop new tools for the discovery and analysis of structural variants (SVs). The authors describe 10 novel software tools to overcome gaps in the available tools to more accurately identify, genotype and map SVs, as well as provide predictive functional effects of the variants to gene expression. The identification of new and accurate tools to identify SVs is of critical importance for understanding the genetic underpinnings of complex disease with no monogenic pattern of disease inheritance. This manuscript is the product of a virtual international hackathon, describing newly developed tools as a foundation for future discussion and development, and the links to the already available online tools provide additional and necessary information for understanding the software applications and outputs. I recommend this manuscript for indexing following amendments. From a functional genetics perspective, I provide the following comments and suggestions:

1. The manuscript does not provide full detail about the code, methods and analysis of the individual tools, however, the purpose of the manuscript is to describe the hackathon and the resultant projects, with the intention for future development of these tools. This could be made clearer in the text with the development status and future requirements for each tool discussed.

2. A summary table of the main applications, benefits, and stage of development for each of the tools and the future direction would be helpful for readers. In particular, more information on the specific applications of each of the tools would be helpful for those reading the manuscript without a bioinformatics background.

3. The manuscript is written with an introduction to each tool, the operation for each and finishing with use cases or results for each. As a suggestion, describing each tool separately under its own heading and incorporating each of these sections would improve the flow of the manuscript.

4. The current tools for accurate SV calling and genotyping from GWAS data are limited for polynucleotide variations. Do any of the described tools address this limitation?
5. The Swagg software describes a tool for SV calling and mapping at both the gene and protein level. However, the advantage of SV calling and mapping at the protein level is not clear given that many SVs are found in non-coding regions. The protein function is not further described in the methods and the interpretation of the figure in the results section was not clear. It would be useful to provide more information for the protein mapping. Can this software show predictive changes to mRNA or protein structures induced by SVs in non-coding regions?

6. The GenVar tool offers a clinically relevant analysis with predictions to SV effects on regulation of gene expression. Does the tool identify SV regions based only on the reference sequence or is it able to further compare a patient sequence to the RefSeq and provide information as to the carriage of potential risk variants/variants known to effect gene expression?

7. There is no figure displaying the result output for the phase sequencing analysis Sniphles tool, this would help in explaining the data analysis and output.

Is the rationale for developing the new software tool clearly explained?
Yes

Is the description of the software tool technically sound?
Partly

Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others?
Partly

Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool?
Partly

Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?
Partly

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: Genetic therapies, functional genetics

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.
This manuscript describes ten Structural Variation related software tools developed at the Second Baylor College of Medicine & DNAnexus hackathon. These tools successfully cover relevant topics in the SV field and certainly will be appreciated by the scientific community. As other reviewers pointed out, my primary concerns are that the text still needs some work to improve readability and fix inconsistencies (e.g., American vs. British spelling). Also, a table summarizing all tools described will be very welcome. There are also some issues specific to some tools that should be addressed. Comments for each tool are described below.

**NibbleSV**: The tool seems great, performs efficiently, and the results are impressive. I just have few comments that would be interesting to see. First, the authors could add more details about the tests using the HG002. If I understood right, the GiaB tier 1 v0.6 was used as candidates to genotyping using the Illumina reads (250bp), is that correct? Also, how was the evaluation done? Second, if possible, it would be interesting to see how NibbleSV compares with other tools (e.g., paragraph). Finally, it would be nice to see how it performs in more complicated regions (e.g., segmental duplications). I wonder if NibbleSV could help filter artifact SVs found by misalignment in those regions.

**CNV2SV**: The motivation and the tool description must be improved in the manuscript. Also, the results are not well described and discussed, being restricted to the legends of Figures 11 and 12.

**Minor:**
In the methods, CNV2SV circularize should be circlize to refer to the R package.
Missing description of letter C in figure 11 legends.

**CoronaSV**: The method proposes a Snakemake pipeline for SV discovery in SARS-CoV-2 genomes. It integrates short and long reads SV discovery tools, as well as assembly-based methods. CoronaSV also handles downloading sequencing data from SRA for variant calling. As a use case, the pipeline was applied to over 200 genomes, and a basic report of the results was shown. I'm not familiar with viral genomes, but I'm wondering what is the actual gain of applying such complex pipeline instead of performing de novo assembly and getting the SVs from it instead.

**CleanSV**: The tool proposes a systematic way for filtering false-positive somatic SVs in cancer genomics. The approach uses germline SVs identified in normal tissue samples to find the best filtering thresholds for specific SV callers. There is no description of how to run the tool, and an example dataset could be added to the GitHub page.

**Sniphles**: More details could be added to the diagram of the pipeline, the calling and merging steps are oversimplified. Is there a reason why Sniffles need Mosdepth to estimate its parameters? It would be great to see how it compares with the “first” approach of phasing mentioned by the
authors, using tagged reads to phase SVs.

**Swagg:** The motivation and methods are not very clear in the manuscript. It seems to me that the protein-graph is basically capturing small-variants. Where SVs fit in this context? The example showed in Figure 15 needs a more detailed explanation (e.g., what the color mean? Is there any SV affecting this protein?). Also, some phrases are repeated in the introduction and methods sections.

**PanOriginSV:** I'm unfamiliar with the topic addressed by this tool. For the little information I could get, the proposal seems very relevant, and the results look promising. In the introduction, the authors discuss the drawbacks of using machine learning approaches and highlight PlasmidHawk advantages, but in the end, propose another machine learning method? What is different in this method compared to existing ones? I'm wondering if extending that tool to work non-linear pangenomes would be possible. Also, it would be nice to see the same benchmarking against PlasmidHawk.

**GeneVar:** The proposal of GeneVar seems interesting and novel. However, not all features described in the manuscript seemed to be implemented. For instance, the authors state that a rare-variant burden test is performed but, I could not find that on the demo website. Also, the annotations are still very basic, the presentation of the overlap is not user-friendly. I would suggest improving how information is shown, with more summarized information about the overlap.

**SVTeaser:** SVTeaser is a framework for benchmarking SVs using simulations. Simulations are performed using SURVIVOR. Overall, the method seems well developed, and the contribution is relevant for the field.

**XSVLen:** The methods sound truly relevant and novel. I'm very interested in applying this method in the future. I'm wondering if a more refined report could be implemented. It would be interesting to see results summarized for each haplotype separately. In the manuscript, the results table is hard to understand. For example, the first row shows 14942 INS >=50, while the second row shows a higher number when I would expect just a subset based on the description.

**Is the rationale for developing the new software tool clearly explained?**
Yes

**Is the description of the software tool technically sound?**
Partly

**Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others?**
Partly

**Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool?**
Partly
Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?
Partly

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Bioinformatics, genomics, genetics

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

Author Response 14 Jun 2021

**Ann Mc Cartney**, NIH, Washington, USA

*We would like to thank Ricardo for their time on giving the authors of this manuscript feedback and comments. Upon receiving these comments the corresponding authors reached out to the developers of each of the tools developed as part of this hackathon, each of which have done their utmost to respond to each comment or question.*

**Comment 1:** General: This manuscript describes ten Structural Variation related software tools developed at the Second Baylor College of Medicine & DNAnexus hackathon. These tools successfully cover relevant topics in the SV field and certainly will be appreciated by the scientific community. As other reviewers pointed out, my primary concerns are that the text still needs some work to improve readability and fix inconsistencies (e.g., American vs. British spelling). Also, a table summarizing all tools described will be very welcome. There are also some issues specific to some tools that should be addressed.

*Response: We agree that the format of the manuscript is somewhat cumbersome to read, however this is a pre-requisite determined by the journal for manuscripts associated with hackathons.*

**Comment 2:** For the Developer of NibbleSV: The tool seems great, performs efficiently, and the results are impressive. I just have a few comments that would be interesting to see. First, the authors could add more details about the tests using the HG002. If I understood correctly, the GiaB tier 1 v0.6 was used as a candidate for genotyping using the Illumina reads (250bp), is that correct? Also, how was the evaluation done? Second, if possible, it would be interesting to see how NibbleSV compares with other tools (e.g., paragraph). Finally, it would be nice to see how it performs in more complicated regions (e.g., segmental duplications). I wonder if NibbleSV could help filter artifact SVs found by misalignment in those regions.

*Response: Yes, the tests were done using GIAB Tier 1 v0.6 truth SV set and alignment file of illumina 2x250 bp dataset for HG002 sample. The true positive count is the number of genotyped SVs that are not parental SVs i.e. HG2count > 0 and the FP count is the parental SVs i.e. HG2count=0. The parental exclusive SVs that are not genotyped are counted as TNS.*
and the HG002 SVs that are not genotyped by nibSV are counted as FN. The following
metrics are used to evaluate performance of nibSV:

Recall = TP/(TP+FN); Precision = TP/(TP+FP); Accuracy = (TP+TN)/(TP+FP+FN+TN); False
discovery rate = FP/(TP+FP). The performance of nibSV was compared with SVTyper.
SVTyper can genotype only non-insertion (i.e. here deletions) type SVs, so we compared
nibSV with SVtyper only for deletion type of SVs. nibSv was run with parameters k=23 and
m=0.

| Tool     | TP   | FP   | FN   | TN   | Recall | Precision | Accuracy | FDR |
|----------|------|------|------|------|--------|-----------|----------|-----|
| nibSV    | 12152 | 2055 | 17957 | 5248 | 0.4    | 0.86      | 0.47     | 0.14 |
| SVTyper  | 5873  | 212  | 24236 | 7091 | 0.20   | 0.97      | 0.35     | 0.03 |

The high recall score of nibSV shows that it outperformed SVTyper for genotyping the non-
insertion type SVs. The high precision rate and low false discovery rate of SVTyper is due to
the very low number of total genotyped SVs. The Paragraph tools required to add the
padding bases to the SVs in the truth set. We could not successfully run Paragraph without
making any changes to the truth set. Therefore, we have not included that tool in this
result.

Comment 3: For the CNV2SV Developers: The motivation and the tool description must be
improved in the manuscript. Also, the results are not well described and discussed, being
restricted to the legends of Figures 11 and 12. Minor: In the methods, CNV2SV circularize should
be circlize to refer to the R package. Missing description of letter C in figure 11 legends.

Response: Thank you for your assessment. We have restructured the respective sections in
the manuscript in order to clarify the tool motivation and description of operation. We
have also corrected the figure accordingly.

Comment 4: For the CoronaSV Developers: The method proposes a Snakemake pipeline for
SV discovery in SARS-CoV-2 genomes. It integrates short and long reads SV discovery tools,
as well as assembly-based methods. CoronaSV also handles downloading sequencing data
from SRA for variant calling. As a use case, the pipeline was applied to over 200 genomes,
and a basic report of the results was shown. I'm not familiar with viral genomes, but I'm
wondering what is the actual gain of applying such a complex pipeline instead of
performing de novo assembly and getting the SVs from it instead.

Response: The reviewer makes a good point with respect to de novo assembly from short
reads, which would accurately capture small structural variants. However, given SARS-CoV-2
is sequenced from an intrahost population, which often is from metatranscriptomes
(RNA-sequencing of a microbiome sample) containing subgenomic RNAs, SVs reflect the
transcriptional landscape of SARS-CoV-2. Thus we feel it is of use to capture the
concordance across short reads, long reads, and assembly for structural variants calling.

Comment 5: For the CleanSV Developers: The tool proposes a systematic way for filtering
false-positive somatic SVs in cancer genomics. The approach uses germline SVs identified in
normal tissue samples to find the best filtering thresholds for specific SV callers. There is no
description of how to run the tool, and an example dataset could be added to the GitHub
Response: Thank you to the reviewer for the response. As mentioned in the previous reviews, our contribution to the hackathon was a proof of concept. Releasing the filters for general use would require more work specifically investigating tumors of a given sequencing depth and tumor content. The panel of normals used for the paper are accessible, as cited in the paper.

Comment 6: For the Sniphles Developers: More details could be added to the diagram of the pipeline, the calling and merging steps are oversimplified. Is there a reason why Sniffles need Mosdepth to estimate its parameters? It would be great to see how it compares with the “first” approach of phasing mentioned by the authors, using tagged reads to phase SVs.

Response: We would like to thank the reviewer for the comments and questions. We agree with the reviewer regarding the merging step, while this is a prototype of the idea, we will be working on enhancing the merging step. Additionally, we stated that in the manuscript.

Comment 7: For the Sniphles Developers: “Sniffles need Mosdepth to estimate its parameters”. Response: Sniffles use a minimum of 10 reads to call an SV, we suggest to use mosdepth to calculate the coverage for the region we call SVs in, so it could adapt to low and high coverage as well as technology used for sequencing (PacBio or Oxford Nanopore Technology).

Comment 8: For the Sniphles Developers: “It would be great to see how it compares with the “first” approach of phasing mentioned by the authors, using tagged reads to phase SVs.”.

Response: We will be working on that, as the second approach still needs more tuning.

Comment 9: For the Swagg Developers: The motivation and methods are not very clear in the manuscript. It seems to me that the protein-graph is basically capturing small-variants. Where do SVs fit in this context? The example shown in Figure 15 needs a more detailed explanation (e.g. What does the color mean? Is there any SV affecting this protein?). Also, some phrases are repeated in the introduction and methods sections.

Response: We would like to thank the reviewer for their comments and questions regarding the manuscript. In response to their questions, the different colors are associated with different sequences used to build the graph, with each sequence taking a certain path in the graph, which yields the colors.

Comment 10: For the PanOriginSV Developers: I’m unfamiliar with the topic addressed by this tool. For the little information I could get, the proposal seems very relevant, and the results look promising. In the introduction, the authors discuss the drawbacks of using machine learning approaches and highlight PlasmidHawk advantages, but in the end, propose another machine learning method? What is different in this method compared to existing ones? I’m wondering if extending that tool to work non-linear pangenomes would be possible. Also, it would be nice to see the same benchmarking against PlasmidHawk.

Response: We’d first like to thank the reviewer for their feedback. While PanOriginSV does
use machine learning, it has the advantage that random forests are much easier to inspect, train, and interpret than other ML models. In addition, the step of constructing the pangenome already constructs features of the sequence. The results of PlasmidHawk are shown as the “linear” model in Table 1 as opposed to PanOriginSV, the graph model.

Comment 11: For the GeneVar Developers: The proposal of GeneVar seems interesting and novel. However, not all features described in the manuscript seemed to be implemented. For instance, the authors state that a rare-variant burden test is performed but, I could not find that on the demo website. Also, the annotations are still very basic, the presentation of the overlap is not user-friendly. I would suggest improving how information is shown, with more summarized information about the overlap.

Response: Thank you reviewer, for your thoughts and assessment. Indeed, GeneVar is a work in progress and what you see now is the result of only 3 days of work. The developers agree that to make this a public tool, additional time and effort is needed.

Comment 12: For the SVTeaser Developers: SVTeaser is a framework for benchmarking SVs using simulations. Simulations are performed using SURVIVOR. Overall, the method seems well developed, and the contribution is relevant for the field.

Response: We appreciate the reviewer’s careful and detailed assessment of our work, as well as his complimentary remarks. We believe that our tool will be useful in genetic research on SV predictions.

Comment 13: For the XSVLen Developers: The methods sound truly relevant and novel. I’m very interested in applying this method in the future. I’m wondering if a more refined report could be implemented. It would be interesting to see results summarized for each haplotype separately. In the manuscript, the results table is hard to understand. For example, the first row shows 14942 INS >=50, while the second row shows a higher number when I would expect just a subset based on the description.

Response: Many thanks to the reviewer for pointing this out and for the interest in the method, we have updated the main table for including the right numbers and also to help comprehension of categories. This was due to a bug in the annotation script that has been fixed. We did not perform a detailed exploration of haplotype-overlapping calls or the generation of a more detailed report as this was a proof-of-concept of the method but we agree it would be interesting if future work is performed.

We would like to thank the reviewer for their kind comments and feedback and hope that they have addressed all questions or concerns posed.

Competing Interests: No competing interests were disclosed.
General comments

This manuscript is a write-up report of an apparently very successful hackathon. Authors present an overview of 10 different approaches to solve various issues in the genomics world of structural variants, some of then sound really promising. I appreciated the modest tone authors haven chosen when describing the current state of each tools. All the tools have at least some exploratory value, and all of them are publicly available, therefore I have no major objections for the manuscript to be indexed.

This being said, I do agree with the first reviewer, I would greatly appreciate if before diving into details of individual methods, a big picture overview would be provided, for example a table with short descriptions, urls and development stages would be really handy. Disclosing the development stages would be particularly useful for a reader to know what to expect if proceeding further in reading.

I do think this manuscript will spark a lot of ideas in the SV community, so I also wondered, what are the contribution policies to individual tools? Many of the tools have some future plans and are not yet production ready. Who would be the people to contact if someone would be interested in contributing and helping out with finishing a tool that would be particularly helpful for their research project?

The manuscript could benefit immensely from a big reorganisation. I do believe that classical Intro/Meth/Results organisation is not doing this manuscript a favour. It was really hard for me to keep track of ten threads in parallel and at some point I just gave up on top-bottom reading. I ended up scrolling back and forth between different sections to get information regarding one tool I tried to understand at the time. I also think most of the people will be mostly interested in one of the tools only when opening the manuscript for reading, therefore I would propose to keep the common Introduction and Conclusion sections but restructure the rest of the paper on per method basis.

I also appreciated authors' effort to comply with FAIR principles. Although I have not check every tool, those I did were neatly documented. Thoughts regarding individual tools are listed bellow.

NibblerSV

If I understand right. The genotyping using NibblerSV is done in two phases - 1. calculating a dictionary of specific kmers for each allele, I would call those diagnostic kmers (e.g. allele specific kmers absent in the reference genome) and 2. matching kmers of resequencing data to the diagnostic kmers. I did find the idea very clearly explained.
The part I did not understand is how it's decided if a sample does or does not carry a variant. Does it need to have all the diagnostic kmers at a certain coverage? Or at least some proportion? What if the sample contains kmers of multiple diagnostic markers? Is it possible to genotype a heterozygous SV? Is there any measure of confidence in genotyped alleles? I personally would prefer to read more about these details over technical aspect such as different types of temporary files that the software is producing that actually are mentioned.

I also wondered, given the approach of generating catalogues of SV-diagnostic kmers, I would anticipate very variable power to detect SVs of different sizes and types. For example, recent inversions will be identified only through very few kmers found at the edge of the inversion, compared to large scale insertions that will probably harbour many diagnostic kmers and clearer signal. Did you look at the sensitivity/precision in respect to size/type? Would that be a consideration one should have when deciding if to use nibSV or paragraph or VG?

**CNV2SV**

This tool has the least intuitive motivation of all the tools presented. I would naturally assume all loci with CNV should also be called as SVs, hence I was not sure why it is interesting at all getting the same information twice. However, the results of this analysis proved my intuition wrong. Only very few CNVs are linked to SVs. However, the suggested explanation with length disparity was a bit dissatisfying to me. How comes that the SV and CNV callers call variants of so different size? Which are shorter/longer? Do you think the CNV/SV reconciliation could be more overlapping with a different choice of SV/CNV caller?

This software has absolutely outstanding description on the GitHub repository.

**Minor comments:**

Figure 12, panel B has a cropped y-axis label.

In the Methods the text "formats (from Parliament" has an opening bracket that is never closed.

**CoronaSV**

I did appreciate the idea of scalable pipeline for conservative analyses of SARS-CoV-2 genomes. Authors made many unjustified choices. For example, why did you chose in particular Manta, Delly, Lumpy and Tardis to detect SV using short reads? But I suppose it is understandable why it would be challenging justify every choice in limited time of a hackathon.

The step of the analysis I am worried about is the merging by SURVIVOR, it also happens to be the only one that does not seem to be included in the GitHub repository. What parameters were used for individual merging steps using SURVIVOR? Given the genome of SARS-CoV-2 is really small, I believe that the outcome of this pipeline will be very sensitive to the chosen thresholds for merging variants.

This project aims to get "trustworthy SV calls across SARS-Cov-2", but there was only very little effort in validation of the SV set detected as far as I can tell. I was honestly quite surprised to see how many deletions that are >20k long you reported as confident. Is it really possible for SARS-
CoV-2 to lose more than 2/3 of its genome? Could they represent some SV calling artefacts? One way how to check the quality of called SVs would be compare phylogenetic relationships of the analysed viruses that are probably available for all the sequenced data anyway and verify that the common SVs are shared within monophyletic clusters.

Finally, the provided data frame with the input data contains various types of libraries. How does SV calling works on RNA-seq data? To be honest, I am not even sure what "RNA-seq" means in the context of RNA viruses, but I do find strange that all the libraries seem to be treated the same. Could some of those huge deletion be simply the selection process of the non-WGS libraries?

I understand this is a hackathon paper, and therefore it would be unreasonable to ask for more analyses, but I do believe that the wording of this tool should be toned down a lot. I think this is a good first draft for this ambitious pipeline, but loads of work and thought should be given to it before the called SV sets are called trustworthy.

**CleanSV**

The idea here is to figure out empirical filtering thresholds to minimise false positives on called SVs. This is a great idea and it's desperately needed to have a better community resources to reduce the need for manual curation.

I am also very sure that these thresholds would have great applicability in other species and germline SV calling. Is there any chance of more general applicability?

Given authors have decided not to share the filtering thresholds, I think it should be clearly stated early on that this is only a conceptual description of the workflow and results are not provided yet.

**Minor comments:**

I resume hs37d5 in "Illumina short reads using novoalign hs37d5 HG002 BAM" means it a human data, but I do think it should be explicitly mentioned.

A bunch of citations are in a different format, something like "(7)" and without actual reference.

**Sniphles**

Again, a very timely contribution. Phasing of SVs will me more and more important as long read technologies are more and more available even for population genomics studies.

I found the idea very clear, and sound. I did wonder, however, about the merging step. Technically, SVs from different halplotype need to be merged as if only the SV is homozygous in the alternative allele, therefore I wonder if it is appropriate to lump together everything that is closer than 1000 bases from each other. How many such variants were detected?

That's kind of related to the only thing I was missing - results. The test case of this method is basically a more confusing repetition of what is already written in methods without any actual numbers of phased SVs. How successful was phasing? How many and how long blocks were phased? If unavailable yet, you should be explicit that it's untested approach.
Although I have not tried to install the software, I appreciated that the GitHub repository was very neat and the code very well organised.

**Swagg**

This was another tool with rather difficult text and figure. It is still not very clear to me how using protein graph should help SV calling.

I was also rather confused about the construction of protein graph. In methods the you write, "Protein graphs are generated using a multiple sequence alignment of the protein", but what is "the protein"? I originally imagined SWAGG would use genome annotation, where the multiple sequence alignment comes into play? I got it lot better from the presentation I found on the GitHub, but I am still not 100% sure how it's done. The presented figure is very pretty, but I did not understand at all what I was supposed to look at.

I just think it's pity, it does look like a loads of work with a really interesting idea. I would recommend to spend more effort on explanation of the reasoning behind and general clarity of the text.

I also wondered, is the protein information used for SV calling in the end? If no, why is not the input to your software a vcf file with SVs? Why it's important they are called by GATK and deepVariant? That was another confusing bit for me.

**PanOriginSV**

A machine learning approach that takes genome variation graph that includes both SNVs and SVs and determines lab of origin. First, I would like to disclaim I am a bit out of my depth reviewing this method, as I never worked with lab strains and genetically modified organisms or thought about the problem of lab attribution.

I was a bit confused about the benchmarking. The Genetic Engineering Attribution Challenge seem to have plenty of approaches with really successful prediction algorithm, why is the approach compared to a single method (PlasmidHawk) that does not seem to be compared to the other approaches in the challenge? I also wondered why the results were shown in per-cluster basis, not overall accuracy, so they could be comparable to the approaches from the Genetic Engineering Attribution Challenge.

Perhaps a silly thought, but do you think that machine learning approach is scalable? Having comparable training dataset to the unclassified dataset is essential for meaningful prediction, would the approach work for novel strains?

**Minor comments:**

This sentence "PanOriginSV has three additional open source dependencies which are MMSEQ2 (for clustering), BCALM (pangenome creation) and GraphAligner (for sequence-to-graph alignment)" should probably contain references to respective software.
"10-50x less than the linear model (depending on the cluster)." Less than PlasmidHawk?

**GeneVar**

This tool has a functional prototype and it works! It allows a user to browse variants related to any human gene, which is fun. However, I do think it should be called "human SV browser" (I originally imagined the intend is to make a generic SV browser).

I appreciated attempt of making information about human SVs accessible to more general public, but I do wonder how close to the goal authors got. I don't think allele frequency spectrum is an easy plot for a user without background in genetics. Also, there are loads of variants shown for every gene, would it be possible in future to somehow prioritise to the "most likely relevant" variants? What you think could be done to improve accessibility of the tool by for example GPs?

*Minor comment:*

In the app, it's a bit confusing that SV length is filtered on the panel on left, while all the remaining variables are filtered by clicking on the empty "all" fields bellow column names.

**SVTeaser**

I really like this idea for a framework for simulating reads and benchmarking SV callers. Beside previously utilised approach of generating insilco generated SVs, SVTeaser also allows a lexicon based generation of SVs. I also found the writing and description of this method very nice and understandable.

One of the main general problems of simulating sequencing reads is that it is very hard to take into consideration all biases of real life sequencing. For example, cross sample or bacterial contamination are really hard to consider. As a consequence, benchmarking using simulations is (probably) always overly optimistic. Would it be useful to attempt to quantify the optimism? For example by simulating reads from a sequenced genome with known SVs? Then the precision and recall of SVs could be compared between the simulated and real sequencing data.

**XSVLen**

Sound approach, but took me while to see any added value compared to the original study by Chin *et al.* (2020)¹. Do I understand right that what was done on MHC locus you generalised in a framework for any region with haplotype-resolved assemblies? I think it should be more clearly stated what was done by by Chin *et al.* before and how your approach differs. Also, in the text you refer to Nurk *et al.*, although the reference is Chin *et al.* (I think it was confused with this other paper presenting HiCanu²).

In the table, the second row says "Number of these calls that overlap assembly contigs". First, how can be any variant called outside of an assembly contig? Do you mean overlapping with haplotype-resolved contigs? You also write "these calls", which made me think it's a subset of the first row, but insertions >50bp seems to have a higher number in the first row (14942 vs 17481). Actually, I did not really understand the third row either, where the "Number of truvari" come from? Are these cuteSV compared to the gold standard? If so, how comes the FP and TP do not sum up to
the same number as the first row?

Similarly to SWAGG, I think this could be a useful tool and you seem to do loads of work on the coding part. However, the text was very confusing to me and I do think with more effort in it you could reach much wider readership.

**Minor comments:**

This sentence is hard to read "This software is a framework for haplotype-resolved assemblies for benchmarking SV detection algorithms (Figure 10).", what about "This software is a framework for benchmarking SV detection algorithms against haplotype-resolved assemblies (Figure 10)."?

"All methods are open-source licensed and have been made available on GitHub: https://github.com/collaborativebioinformatics." is an unnecessary statement as FAIR principles are disclosed for all 10 tools and all the links are provided at the bottom.

**References**

1. Chin C, Wagner J, Zeng Q, Garrison E, et al.: A diploid assembly-based benchmark for variants in the major histocompatibility complex. *Nature Communications*. 2020; 11 (1). [Publisher Full Text](https://www.nature.com/articles/s41467-020-15289-8)
2. Nurk S, Walenz BP, Rhie A, Vollger MR, et al.: HiCanu: accurate assembly of segmental duplications, satellites, and allelic variants from high-fidelity long reads. *Genome Res*. 30 (9): 1291-1305 [PubMed Abstract](https://www.ncbi.nlm.nih.gov/pubmed/32627751) | [Publisher Full Text](https://www.genome.org/article-pdf/30/9/1291/34861849.pdf)

**Is the rationale for developing the new software tool clearly explained?**

Partly

**Is the description of the software tool technically sound?**

Partly

**Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others?**

Yes

**Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool?**

Yes

**Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?**

Yes

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Genomics, evolutionary biology, bioinformatics

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have...
significant reservations, as outlined above.

Author Response 06 Jun 2021

Ann Mc Cartney, NIH, Washington, USA

We would like to thank Kamil Jaron for his time on giving the authors of this manuscript feedback and comments. Upon receiving these comments the corresponding authors reached out to the developers of each of the tools developed as part of this hackathon, each of which have done their utmost to respond to each comment or question. The responses to each of the reviewer's comments are outlined below in bold and italic.

Comment 1: General: This manuscript is a write-up report of an apparently very successful hackathon. Authors present an overview of 10 different approaches to solve various issues in the genomics world of structural variants, some of them sound really promising. I appreciated the modest tone authors have chosen when describing the current state of each tool. All the tools have at least some exploratory value, and all of them are publicly available, therefore I have no major objections for the manuscript to be indexed. This being said, I do agree with the first reviewer, I would greatly appreciate it if before diving into details of individual methods, a big picture overview would be provided, for example a table with short descriptions, urls and development stages would be really handy. Disclosing the development stages would be particularly useful for a reader to know what to expect if proceeding further in reading.I do think this manuscript will spark a lot of ideas in the SV community, so I also wondered, what are the contribution policies to individual tools? Many of the tools have some future plans and are not yet production ready. Who would be the people to contact if someone would be interested in contributing and helping out with finishing a tool that would be particularly helpful for their research project?

Response: This is a great question. In general the tools are provided as is. The individual authors are part of their github pages. Furthermore these tools might be extended in future hackathons. Some tools are still being worked on (e.g. NibSV). In terms of contacting authors, please contact the authors associated with each tool; this should be listed in each README of the respective tool. Naturally, feel free to contact the lead authors of this paper as well.

Comment 2: General: The manuscript could benefit immensely from a big reorganization. I do believe that the classical Intro/Meth/Results organization is not doing this manuscript a favor. It was really hard for me to keep track of ten threads in parallel and at some point I just gave up on top-bottom reading. I ended up scrolling back and forth between different sections to get information regarding one tool I tried to understand at the time. I also think most of the people will be mostly interested in one of the tools only when opening the manuscript for reading, therefore I would propose to keep the common Introduction and Conclusion sections but restructure the rest of the paper on a per method basis.

Response: While we agree, this is a given from the journal for hackathons. We are following this format.

Comment 3: General: I also appreciated the authors' effort to comply with FAIR principles.
Although I have not checked every tool, those I did were neatly documented. Thoughts regarding individual tools are listed below.

Response: We thank the reviewer for going over these and highly appreciate the feedback.

Comment 4: For the NibblerSV Developers: If I understand right. The genotyping using NibblerSV is done in two phases - 1. calculating a dictionary of specific kmers for each allele, I would call those diagnostic kmers (e.g. allele specific kmers absent in the reference genome) and 2. matching kmers of resequencing data to the diagnostic kmers. I did find the idea very clearly explained. The part I did not understand is how it's decided if a sample does or does not carry a variant. Does it need to have all the diagnostic kmers at a certain coverage? Or at least some proportion? What if the sample contains kmers of multiple diagnostic markers? Is it possible to genotype a heterozygous SV? Is there any measure of confidence in genotyped alleles? I personally would prefer to read more about these details over technical aspects such as different types of temporary files that the software is producing that actually are mentioned.

Response: NibSV currently screens the diagnostic kmers against the reference genome. Right now if one of these kmers are found in the reads it is reported that the SV is found in general. This can be improved upon easily (e.g. weight with reference allele kmers) but that is the current state. NibSV does not include a genotyping model nor ref alleles at the moment. Still the early results are very promising. The tests were done using GIAB Tier 1 v0.6 truth SV set and alignment file of illumina 2x250 bp dataset for HG002 sample. The true positive count is the number of genotyped SVs that are not parental SVs i.e. HG2count > 0 and the FP count is the parental SVs i.e. HG2count=0. The parental SVs that are not genotyped are counted as TNs and the non-parental SVs that are not genotyped by nibSV are counted as FN. The following metrics are used to evaluate performance of NibSV: Recall = TP / (TP+FN); Precision = TP/(TP+FP); Accuracy = (TP+TN)/(TP+FP+FN+TN); False discovery rate = FP/(TP+FP). We have compared the performance of nibSV with SVTyper (will update the results in the manuscript). The Paragraph tools required to add the padding bases to the SVs in the truth set. We could not successfully run paragraph without making any changes to the truth set. Therefore, we have not included the paragraph tool in this result.

Comment 5: For NibSV Developers: I also wondered, given the approach of generating catalogues of SV-diagnostic kmers, I would anticipate very variable power to detect SVs of different sizes and types. For example, recent inversions will be identified only through very few kmers found at the edge of the inversion, compared to large scale insertions that will probably harbour many diagnostic kmers and clearer signals. Did you look at the sensitivity/precision in respect to size/type? Would that be a consideration one should have when deciding if to use nibSV or paragraph or VG?

Response: This is again an excellent point. We only focused on insertion and deletions at this moment. Indeed we see a higher recall on insertion compared to deletions. This is also driven by the fact that we see a deletion more often within repeats. Nevertheless, nibSV does not consider the entire length of an insertion but rather only the breakpoints. We think that nibSV has some advantages over runtime and its not reliant over the genome version of the reads. However, Paragraph etc. will likely have more control of different
Comment 6: For the CNV2SV Developers: This tool has the least intuitive motivation of all the tools presented. I would naturally assume all loci with CNV should also be called as SVs, hence I was not sure why it is interesting at all getting the same information twice. However, the results of this analysis proved my intuition wrong. Only very few CNVs are linked to SVs. However, the suggested explanation with length disparity was a bit dissatisfying to me. How come that the SV and CNV callers call variants of so different sizes? Which are shorter/longer? Do you think the CNV/SV reconciliation could be more overlapping with a different choice of SV/CNV caller? This software has an absolutely outstanding description on the GitHub repository. Minor comments: Figure 12, panel B has a cropped y-axis label. In the Methods the text “formats (from Parliament” has an opening bracket that is never closed.

Response: Typically CNV is called against a locus on a reference genome. We take the view that a CNV is caused by one or more SVs but most CNV callers do not identify a specific SV (insertions or deletions with the coordinate associated.) For example, if we know there are additional copies for region A. A CNV caller will call an additional copy of A in the reference coordinate, but one would not know where the additional copy really is and what the exact sequences and the inserted locus. CNV2SV is partially developed to understand the spectrum of CNV. By connecting the CNV calls for SV calls from de novo assembly results, we get a chance to understand the nature of CNVs is the light full de novo assembly results. The differences of the sizes from the different CNV caller and SV caller actually reflects that we still need more work to understand the relationship of CNVs and SVs. (Most CNV callers use short read data with coverage statistics. This is fundamentally different from calling SVs from assembly-assembly alignment in terms of detection methodology.) We anticipate more de novo assemblies of human genomes will help us eventually get better pictures.

Comment 7: For the CoronaSV Developers: I did appreciate the idea of a scalable pipeline for conservative analyses of SARS-CoV-2 genomes. Authors made many unjustified choices. For example, why did you choose in particular Manta, Delly, Lumpy and Tardis to detect SV using short reads? But I suppose it is understandable why it would be challenging to justify every choice in the limited time of a hackathon. The step of the analysis I am worried about is the merging by SURVIVOR, it also happens to be the only one that does not seem to be included in the GitHub repository. What parameters were used for individual merging steps using SURVIVOR? Given the genome of SARS-CoV-2 is really small, I believe that the outcome of this pipeline will be very sensitive to the chosen thresholds for merging variants. This project aims to get “trustworthy SV calls across SARS-Cov-2”, but there was only very little effort in validation of the SV set detected as far as I can tell. I was honestly quite surprised to see how many deletions that are >20k long you reported as confident. Is it really possible for SARS-CoV-2 to lose more than 2/3 of its genome? Could they represent some SV calling artefacts? One way to check the quality of called SVs would be to compare phylogenetic relationships of the analysed viruses that are probably available for all the sequenced data anyway and verify that the common SVs are shared within monophyletic clusters. Finally, the provided data frame with the input data contains various types of libraries. How does SV calling work on RNA-seq data? To be honest, I am not even sure what
"RNA-seq" means in the context of RNA viruses, but I do find it strange that all the libraries seem to be treated the same. Could some of those huge deletions be simply the selection process of the non-WGS libraries? I understand this is a hackathon paper, and therefore it would be unreasonable to ask for more analyses, but I do believe that the wording of this tool should be toned down a lot. I think this is a good first draft for this ambitious pipeline, but loads of work and thought should be given to it before the SV sets are called trustworthy.

Response: The reviewer makes several helpful points. To the first point, we selected a handful popular SV callers from short reads suitable for the task, and given the short duration of the hackathon, we unfortunately were unable to evaluate a more comprehensive set. To the general point of do we expect to see such large SVs, given SARS-CoV-2 is sequenced from an intrahost population, which often is from metatranscriptomes (RNA-seq applied to a microbiome sample) containing subgenomic RNAs, large SVs can be observed as it captures the transcriptional landscape of SARS-CoV-2. And we have added details of SURVIVOR to the Github repository, thank you for pointing this out.

Comment 8: For the CleanSV Developers: The idea here is to figure out empirical filtering thresholds to minimise false positives on called SVs. This is a great idea and it's desperately needed to have better community resources to reduce the need for manual curation. I am also very sure that these thresholds would have great applicability in other species and germline SV calling. Is there any chance of more general applicability?

Response: We thank the reviewer for the assessment and insightful question. In terms of more general applicability, we agree with the reviewer entirely. However, in order to release these filters/thresholds to the larger community for general use, the optimal approach would be to focus on a particular use of SV calling (e.g. translational genomics in oncology) with a particular assay of a certain average coverage and tumor purity. These filters need to be tailored specifically for these parameters. We suspect generating filters per SV caller for general use would result in minimal quality control; this is especially true within cancer genomics in terms of calling somatic and germline SVs accurately. As for focusing on other species (even within a germline context), the main impediment is that we would need a sufficient number of samples from species X at a specific time in order to create these filters.

Comment 9: For the CleanSV Developers: Given authors have decided not to share the filtering thresholds, I think it should be clearly stated early on that this is only a conceptual description of the workflow and results are not provided yet.

Response: We agree with this assessment, and have tried to clarify this in the manuscript. This paper is the result of a hackathon, and the ideas within are largely proof of concepts which have value outside of a more comprehensive paper.

Comment 10: For the CleanSV Developers: Minor comments: I resume hs37d5 in "Illumina short reads using novoalign hs37d5 HG002 BAM" means it is human data, but I do think it should be explicitly mentioned. A bunch of citations are in a different format, something like "(7)" and without actual reference.
Response: We have clarified and fixed these issues in the manuscript.

Comment 11: For the Sniphles Developers: Again, a very timely contribution. Phasing of SVs will be more and more important as long read technologies are more and more available even for population genomics studies. I found the idea very clear, and sound. I did wonder, however, about the merging step. Technically, SVs from different haplotype need to be merged as if only the SV is homozygous in the alternative allele, therefore I wonder if it is appropriate to lump together everything that is closer than 1000 bases from each other. How many such variants were detected? That's kind of related to the only thing I was missing - results. The test case of this method is basically a more confusing repetion of what is already written in methods without any actual numbers of phased SVs. How successful was phasing? How many and how long blocks were phased? If unavailable yet, you should be explicit that it's an untested approach. Although I have not tried to install the software, I appreciated that the GitHub repository was very neat and the code very well organised.

Response: We would like to thank the reviewer for the ideas and suggestions. Also, we would like to mention that, this is a prototype of the ideas, and we opened an issue regarding the suggestions. We shall implement a parameter to set the minimum accepted length to identify 2 heterozygote SVs as one.

“How many such variants were detected? That's kind of related to the only thing I was missing - results.”

We used chromosome 20, it contained 652 SVs with minimum 13 to support SVs. 239 SVs are heterozygote we managed to phase 182 of those (76.15%) in 76 phase blocks using the first method. For the second method, while we completed the tool, and it worked as expected, but sadly it did not give the expected results, we still need to work on debugging and enhancing the second method.

Comment 12: For the Sniphles Developers: “I did wonder, however, about the merging step. Technically, SVs from different haplotype need to be merged as if only the SV is homozygous in the alternative allele, therefore I wonder if it is appropriate to lump together everything that is closer than 1000 bases from each other.”

Response: We agree with the suggestion, we shall implement a parameter that takes as an input the maximum distance to merge heterozygote SVs.

Comment 13: For the Sniphles Developers: “The test case of this method is basically a more confusing repetion of what is already written in methods without any actual numbers of phased SVs.”

Response: We changed it accordingly.

Comment 14: For the Sniphles Developers: “How successful was phasing? How many and how long blocks were phased? If unavailable yet, you should be explicit that it's an untested
approach."

Response: This is a prototype we worked only on chromosome 20 using HiFi data, we still need to develop our method, we updated the manuscript to state that.

Comment 15: For the Swagg Developers: This was another tool with rather difficult text and figure. It is still not very clear to me how using a protein graph should help SV calling. I was also rather confused about the construction of the protein graph. In the methods you write, "Protein graphs are generated using a multiple sequence alignment of the protein", but what is "the protein"? I originally imagined SWAGG would use genome annotation, where the multiple sequence alignment comes into play? I got it a lot better from the presentation I found on GitHub, but I am still not 100% sure how it's done. The presented figure is very pretty, but I did not understand at all what I was supposed to look at. I just think it's a pity, it does look like a load of work with a really interesting idea. I would recommend spending more effort on explanation of the reasoning behind and general clarity of the text. I also wondered, is the protein information used for SV calling in the end? If not, why is not the input to your software a vcf file with SVs? Why is it important that they are called by GATK and deepVariant? That was another confusing bit for me.

Response: We would like to thank the reviewer for their comments and questions, specifically with respect to why we should build a variation graph and not just keep variations in a VCF file. Building graphs from amino acid sequences can be used to align new amino acid sequences to these graphs, thereby finding similarities to other sequences. A modified Smith-Waterman algorithm that works for DAGs (Directed Acyclic Graphs) can be used for this job. Due to how proteins work and fold, it is possible to have a gene between two samples from two different genres in bacteria that are very distant (20-30% sequence similarity), however, due to nonsense mutations, the amino acid sequence similarity is higher (50-60%). For example, in Myxobacteria, looking at two different samples from two different genres or families, building a pangenome graph out of the DNA is not really feasible due to the amount of variations between them. However, looking at amino acid sequences, more similarities can be found. Therefore, with building a graph out of the amino acid, paths can be labeled with metadata (certain conserved nodes for example are associated with antibiotic resistance, aligning a new sequence to this graph, can also tell us about this new sequence). This can additionally be accomplished via HMM modeling, but, building a graph adds a visualization element, and with the recent optimizations in graphical mapping algorithms, it may be faster at aligning a sequence to a graph than using some HMM model. Building on the idea that we can add metadata to paths in the graph, looking at this paper for example, “Jaillard et al. (2018). “A Fast and Agnostic Method for Bacterial Genome-Wide Association Studies: Bridging the Gap between K-Mers and Genetic Events.” We see how from looking at the graphs, certain nodes can be associated with resistance. However, for example, this method breaks down when the similarity between the sequence we’re looking at drops, as it’s hard to find shared k-mers then. However, going to amino acid space, the similarity increase and similar methods and algorithms can then be used but with the amino acid alphabet.

Comment 16: For the PanOriginSV Developers: A machine learning approach that takes a genome variation graph that includes both SNVs and SVs and determines lab of origin. First,
I would like to disclaim I am a bit out of my depth reviewing this method, as I never worked with lab strains and genetically modified organisms or thought about the problem of lab attribution. I was a bit confused about the benchmarking. The Genetic Engineering Attribution Challenge seems to have plenty of approaches with really successful prediction algorithms, why is the approach compared to a single method (PlasmidHawk) that does not seem to be compared to the other approaches in the challenge? I also wondered why the results were shown on a per-cluster basis, not overall accuracy, so they could be comparable to the approaches from the Genetic Engineering Attribution Challenge. Perhaps a silly thought, but do you think that machine learning approach is scalable? Having a comparable training dataset to the unclassified dataset is essential for meaningful prediction, would the approach work for novel strains? Minor comments: This sentence "PanOriginSV has three additional open source dependencies which are MMSEQ2 (for clustering), BCALM (pangenome creation) and GraphAligner (for sequence-to-graph alignment)" should probably contain references to respective software. "10-50x less than the linear model (depending on the cluster)." Less than PlasmidHawk?

Response: We’d first like to thank the reviewer for their feedback and questions. While other tools have been developed for the same problem, Plasmidhawk was recently shown to outperform the other state-of-the-art methods while also being more straightforward to run. Additionally, Plasmidhawk does not require a GPU in order to train efficiently. The main reason for showing per-cluster accuracy was to show the performance of our method on large groups of highly similar sequences. In addition, machine learning methods are not well suited for small cluster scenarios due to the lack of training data. Unfortunately, the code for all other methods in the GEAC was unavailable, therefore we were unable to compare results without submitting to the challenge. However, we do think that machine learning is scalable, although the preprocessing of the data will become more important as the dataset grows. For novel strains that have no similarity to the database, our method would not work but we are confident that no other method would. However, for novel strains that have “signatures” in the database, these would be represented by nodes in the pangenome that are relatively unique to certain labs.

Comment 17: For the GeneVar Developers: This tool has a functional prototype and it works! It allows a user to browse variants related to any human gene, which is fun. However, I do think it should be called a “human SV browser” (I originally imagined the intent is to make a generic SV browser). I appreciated the attempt to make information about human SVs accessible to the general public, but I do wonder how close to the goal the authors got. I don't think the allele frequency spectrum is an easy plot for a user without a background in genetics. Also, there are loads of variants shown for every gene, would it be possible in future to somehow prioritise to the "most likely relevant" variants? What you think could be done to improve accessibility of the tool by for example GPs? Minor comment: In the app, it's a bit confusing that SV length is filtered on the panel on the left, while all the remaining variables are filtered by clicking on the empty "all" fields below column names.

Response: Thank you reviewer for your thoughts and questions. We agree that ranking the variants shown could be helpful for genes with a lot of variants. While right now a long list of variants suggest more filtering needed by the user, it would be better to have the most
impactful SVs shown first by default. In terms of improving accessibility of the tool to GPs, etc. We think adding documentation and tutorials to the tool would be helpful, for example to interpret the frequency histogram, and will be added as the tool matures. We are also keen on receiving feedback directly from clinicians and GPs. We are exploring ways to reach out to them to learn what should be changed, improved, or added.

Comment 18: From the SVTeaser Developers: I really like this idea for a framework for simulating reads and benchmarking SV callers. Beside the previously utilised approach of generating in silico generated SVs, SVTeaser also allows a lexicon based generation of SVs. I also found the writing and description of this method very nice and understandable. One of the main general problems of simulating sequencing reads is that it is very hard to take into consideration all biases of real life sequencing. For example, cross sample or bacterial contamination are really hard to consider. As a consequence, benchmarking using simulations is (probably) always overly optimistic. Would it be useful to attempt to quantify optimism? For example by simulating reads from a sequenced genome with known SVs? Then the precision and recall of SVs could be compared between the simulated and real sequencing data.

Response: We agree that benchmarking against simulated data is overly optimistic and that it's hard to accurately replicate library preparation errors such as what you have described. A primary goal of SVTeaser is to assist developers of SV calling algorithms with assessing how their tool would perform without needing to quantify their algorithm's problems versus their sequencing's problems. A well benchmarked, production-ready tool shouldn't rely on only SVTeaser and would always have real data to test against. However, should someone want to simulate more realistic sequencing experiments by adding batch effects such as cross-sample contamination, it would be possible for them to run SVTeaser twice - once over their target sample's SVs at e.g. 28x coverage and again with a second sample's SVs at e.g. 2x coverage. Then, simply concatenating the sets of generated reads prior to read-mapping would create a simplistic simulation of ~5% cross-sample contamination. While SVTeaser doesn't currently automate this kind of operation, we will strongly consider how best to incorporate the functionality.

Comment 19: For the developers of XSVLen: Sound approach, but took me a while to see any added value compared to the original study by Chin et al. (2020)1. Do I understand right that what was done on MHC locus you generalised in a framework for any region with haplotype-resolved assemblies? I think it should be more clearly stated what was done by Chin et al. before and how your approach differs. Also, in the text you refer to Nurk et al., although the reference is Chin et al. (I think it was confused with this other paper presenting HiCanu2). In the table, the second row says "Number of these calls that overlap assembly contigs". First, how can there be any variant called outside of an assembly contig? Do you mean overlapping with haplotype-resolved contigs? You also write "these calls", which makes me think it's a subset of the first row, but insertions >50bp seems to have a higher number in the first row (14942 vs 17481). Actually, I did not really understand the third row either, where does the "Number of truvari" come from? Are these cuteSV compared to the gold standard? If so, how come the FP and TP do not sum up to the same number as the first row? Similarly to SWAGG, I think this could be a useful tool and you seem to do loads of work on the coding part. However, the text was very confusing to me.
and I do think with more effort in it you could reach much wider readership. Minor comments: This sentence is hard to read "This software is a framework for haplotype-resolved assemblies for benchmarking SV detection algorithms (Figure 10).", what about "This software is a framework for benchmarking SV detection algorithms against haplotype-resolved assemblies (Figure 10)."

Response: Many thanks for the comments and all the interest about the work we performed. The assembly we used for benchmarking is the one generated by Wenger et al. 2020 (NCBI Assembly with accessions GCA_004796485.2 (maternal) and GCA_004796285.1 (paternal) - so we worked with the whole-genome and not only CHM. The reference of CHM locus has been removed from the text and we included the right reference. We have updated the methods section for improving comprehension on the approach we implement. Here, we benchmarked sequence-resolved variants by checking their presence in a haplotype-resolved assembly available and so it works on top of an haplotype-resolved assembly. We have also updated the main table to include the correct numbers and also to help comprehension of categories. The "Number of these calls that overlap assembly contigs" row makes reference to SV called by cuteSV (sequence-resolved SV calling) that would be found also in the assembly by the approach we tested (construct SV query and map queries back to assembly). In the "Number of truvari" (updated to Number of overlapping calls classified with truvari) row we include the classification obtained using truvari – we compare the cuteSV calls and GIAB truth set). This was due to a bug in the annotation script that has been fixed. The minor suggested change has been applied to the text.

Again, we thank the reviewer for his thoughtful comments, suggestions and feedback and hope that the following responses have clarified any queries or concerns.

Competing Interests: No competing interests were disclosed.
The paper has been written collectively, each software participants writing their part, and while of high interest in terms of opening, it is the main limit in the manuscript, in my opinion.

Indeed there is no homogeneity in the way tools are described (global aim, method, implementation, figure, results/tests), and thus it is quite difficult to have a global idea of the different level of development. An initial table with level of advancement (alpha, beta, ready for use, in testing, etc or something like it) would be useful. In addition, a final reformatting by a single writer would smooth the reading.

In details, going for each tools:

**NibbleSV**: what is the impact of SNP on the detection through k-mers? I mean if the individual has more than a very low level of variation, it may increase the FP rate? what is the RAM usage per sample? why providing info of time for 23mers but results of recall/precision with 21.

**CNV2SV**: page 8, output of CNV2SV is said to be a python script (envlink.py)? Why computing the pre-alignment on DNAnexus? Which stats methods are used?

**CORONASV**: are the three approaches (SR, LR, assembly) required in all analyses or you can choose? Why does Flye not represented in Figure 3?

**CleanSV**: on page 6, I cannot see the link with aneuploidy in this specific case. Can you estimate the false negative rate? Do you have an idea of the RAM usage?

**Sniphles**: "the produced BAM from the previous step" is the one produced by Princess? The first sentence of results is the same as in the method. It missed also here a short summary of the results.

**Swagg**: the implementation text is almost the same as in the introduction of the tool. The implementation part is quite short and the fig 6 did not shown any info about the protein graph possibility. Finally, what is the advantage of having the protein graph compared to an alignment showing the variation?

**PanOriginSV**: What are the alignment information included in the training part? Which 'more recent pangenome graph construction tool' are you thinking about?

**GeneVar**: a really interesting tool for praticians. My only question is will it be possible to add new informations about variations outside of strandard DB?

**SVteaser**: do you have any information of time per SV? What are the size of variations you can include? Can we add also translocations or SNP?

**XSVLen**: the figure 10 is quite drafty and would need to be improved for more information. Do you have any information about the recall/precision?

In conclusion, I was very impressed by the high number of tools and of their quality and think this paper is really worthy.
Is the rationale for developing the new software tool clearly explained?
Yes

Is the description of the software tool technically sound?
Partly

Are sufficient details of the code, methods and analysis (if applicable) provided to allow replication of the software development and its use by others?
Partly

Is sufficient information provided to allow interpretation of the expected output datasets and any results generated using the tool?
Partly

Are the conclusions about the tool and its performance adequately supported by the findings presented in the article?
Partly

**Competing Interests:** No competing interests were disclosed.

**Reviewer Expertise:** Genomics, bioinformatics, evolution

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard, however I have significant reservations, as outlined above.

**Author Response 14 Jun 2021**

**Ann Mc Cartney,** NIH, Washington, USA

We would like to thank Francois Sabot for their time on giving the authors of this manuscript feedback and comments. Upon receiving these comments the corresponding authors reached out to the developers of each of the tools developed as part of this hackathon, each of which have highly appreciated your work and thoughtful comments. The responses to each of the reviewers comments are outlined below in bold and italic.

**General Comment 1:** The Pangenome Hackathon paper led by A. McCartney and FJ Sedlazeck was a great moment of intellectual development and findings for pangenomics research on human and coronavirus, pandemics time “oblige”. Ten softwares/methods were developed, some being even released, some being more in alpha state. All these tools are of interest, in their targeted human-related field but also in non-human fields (working on plants, I am interested in more than one of them). The paper has been written collectively, each software participant writing their part, and while of high interest in terms of opening, it is the main limit in the manuscript, in my opinion. Indeed there is no homogeneity in the way tools are described (global aim, method, implementation, figure, results/tests), and thus it is quite difficult to have a global idea of the different levels of
development. An initial table with level of advancement (alpha, beta, ready for use, in testing, etc or something like it) would be useful. In addition, a final reformatting by a single writer would smooth the reading.

Response: We thank the reviewer for this suggestion. Indeed we also struggled as this manuscript has over 50 authors. The tools have so far only been developed within a few days in the context of a remote hackathon and thus represent more prototypes / proof of concepts than production ready methods.

Comment 2: For NibbleSV Developers: “what is the impact of SNP on the detection through k-mers? I mean if the individual has more than a very low level of variation, it may increase the FP rate? What is the RAM usage per sample? why providing info of time for 23mers but results of recall/precision with 21.

Response: That’s a great question. We predict that the impact of SNP is minimal because the SNV must be directly within +/- e.g. 10bp of the SV breakpoint on either side. If there is a SNV it will lower our recall rate. We have further experimented with spaced kmers, that would help in this regard. RAM usage is ~4Gbp per sample.

Comment 3: For CNV2SV Developers: page 8, output of CNV2SV is said to be a python script (envlink.py)? Why computing the pre-alignment on DNAnexus? Which stats methods are used?

Response: Thank you for your question. The CNV2SV script performs the CNV-SV linking steps independently of the tools used to generate the three input files (2.vcf files containing SV calls and CNV calls, respectively, as well as the reference genome as .fasta). In the scope of the Hackathon, we generated the test input VCF files using dipcall (genome-genome-alignment for SVs) and CNVnator (a CNV caller), for a human data set. These steps were executed on DNANexus, since unlike the actual CNV2SV script itself, high processing power and working memory were required, especially for the genome-genome alignment for the two human genome versions. Users can use VCF files as generated by tools and parameters of their choice as input for CNV2SV. While we expect output from most relevant tools that produce calls in VCF format to be compatible, we specifically tested the before mentioned tools in the scope of the Hackathon. While the statistics applied to generate the input SV/CNV calls vary based on the tools used, CNV2SV itself performs the linking using default cutoff values for parameters such as minimum CNV length, minimum alignment identity and maximum distance for adjacent duplication events. Furthermore, basic statistics are recorded for matched CNV-SV pairs as well as the failure reasons for CNVs that could not be linked to a corresponding SV. In response to the final question "Which stats methods are used?" the developers were not sure about which stat this is in reference to.

Comment 4: For the CORONASV Developers: are the three approaches (SR, LR, assembly) required in all analyses or you can choose? Why is Flye not represented in Figure 3?

Response: We thank the reviewer for catching this; we have updated the text to indicate that long read assembly with Flye is not currently supported for such a small genome with
default parameters. Specifically, we have modified “In order to integrate assembly based methods, de novo SARS-CoV-2 assemblies were generated using Unicycler for short-read sequencing and Flye for ONT long-reads.” to instead read “In order to integrate assembly based methods, de novo SARS-CoV-2 assemblies were generated using Unicycler for short-read sequencing.”

**Comment 5:** For the CleanSV Developers: Can you estimate the false negative rate? Do you have an idea of the RAM usage?

*Response:* Here we refer to the presence of somatic structural variations to be synonymous with “aneuploidy”. Aneuploidy itself is an umbrella term used within oncology to refer to somatic abnormalities of a cell’s chromosomes. We try to clarify this with a sentence in the manuscript. In terms of the false negative rate, so far this has been estimated with simulated benchmarks, which surely don’t reflect the complexities of actual clinical data---any real data will be of variable tumor content (purity) and coverage. The best source of collective knowledge within the field is probably the results of DREAM challenges. Indeed, the question of false negatives is one of the deeper questions for all variant calling methods used within cancer genomics. Systematically investigating this problem would require benchmarks from real data, which don’t exist---even robustly characterized cell lines would not be realistic enough. With that in mind, we also wish to emphasize that clinicians are primarily concerned with the presence of false positives. These are potential therapeutic biomarkers which, if missed/ignored by bioinformatic methods, could result in overlooked therapeutic interventions. In terms of RAM usage, the tool provided uses standard functions in R, and will not be memory intensive.

**Comment 6:** For the Sniphles Developers: “the produced BAM from the previous step” is the one produced by Princess? The first sentence of results is the same as in the method. I also missed a short summary of the results.

*Response:* We would like to thank the reviewer, a change was made accordingly in the manuscript.

**Comment 7:** For the Swagg Developers: the implementation text is almost the same as in the introduction of the tool. The implementation part is quite short and the fig 6 did not show any info about the protein graph possibility. Finally, what is the advantage of having the protein graph compared to an alignment showing the variation?

*Response:* Generating the graphs out of the amino acid has a visualization factor, as different sequences can be represented as different paths in the graph, conserved sequences in the MSA will show up as a single node and bubbles are generated from the variations. One can extend this by aligning new sequences to the graph, e.g. using a modified Smith-Waterman algorithm after topologically sorting the graph, this way, the alignment can tell us quickly which path this new sequence takes through this graph, moreover, one can add another layer of data to the paths, for example in bacteria, certain mutations or conserved sequences in the protein are related to antibiotic resistance.

**Comment 8:** For the PanOriginSV Developers: What is the alignment information included
in the training part? Which 'more recent pangenome graph construction tool' are you thinking about?

**Response:** *We’d first like to thank the reviewer for their questions. We thread/align the query sequences to the pangenome graph created using BCALM. The nodes that are hit are used as the features for machine learning. We have experimented with other information as well such as alignment length, %id, etc, but overfitting is an issue that requires further work. Minigraph would be a more recent pangenome graph construction tool that we could use.*

**Comment 9:** For the GeneVar Developers: a really interesting tool for practitioners. My only question is will it be possible to add new information about variations outside of standard DB?

**Response:** *Thank you reviewer, for the question. At this time, we don’t plan to support non-standard data types. However, if you have a specific use case in mind, the developers would be open to a collaboration.*

**Comment 10:** For the SVteaser Developers: do you have any information of time per SV? What are the sizes of variations you can include? Can we also add translocations or SNP?

**Response:** *Thank you to the reviewer for the feedback. In terms of running time per SV, this correlates with the size of SV simulated and the nature of the SV (e.g. translocations being more difficult). However, these operations are relatively straightforward operations manipulating the strings from the FASTA file. Users are able to set the size of SVs within the simulation framework, and in principle could handle any organism. Yes, translocations and SNPs are possible.*

**Comment 11:** For the XSVLen Developers: the figure 10 is quite drafty and would need to be improved for more information. Do you have any information about the recall/precision?

**Response:** *Many thanks for the suggestion and the interest, the figure is intended to give an overview of the method and not a detailed explanation of the pipeline. Regarding precision/recall numbers, we did not check this specifically but in Table 2 the number of TP and FP are summarized for the assembly calls.*

**Competing Interests:** No competing interests were disclosed.
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