A Sequence Distance Graph framework for genome assembly and analysis [version 1; peer review: 2 approved, 1 approved with reservations]

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Abstract
The Sequence Distance Graph (SDG) framework works with genome assembly graphs and raw data from paired, linked and long reads. It includes a simple deBruijn graph module, and can import graphs using the graphical fragment assembly (GFA) format. It also maps raw reads onto graphs, and provides a Python application programming interface (API) to navigate the graph, access the mapped and raw data and perform interactive or scripted analyses. Its complete workspace can be dumped to and loaded from disk, decoupling mapping from analysis and supporting multi-stage pipelines. We present the design and implementation of the framework, and example analyses scaffolding a short read graph with long reads, and navigating paths in a heterozygous graph for a simulated parent-offspring trio dataset.

SDG is freely available under the MIT license at https://github.com/bioinfologics/sdg

Keywords
Genome graph, genome assembly

This article is included in the Python collection.
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Introduction

Sequence graphs are the core representation of genome assemblers. Their use has increased lately thanks to the graphical fragment assembly (GFA) format for graph exchange, tools to work with genome variation graphs, and sequence to graph mappers. But a lack of inter operation between graph-based tools, and limited tools for downstream graph-based analysis, contribute to a perceived complexity which maintains linear sequences as the typical unit of exchange. This flattening of graph representations within pipelines with multiple steps, that use different types of sequencing in an iterative fashion, produces ever-longer linear genome sequences through an information loss process. As a result, genome assembly projects are prone to error propagation and difficult to reproduce and control. These problems can be addressed developing graph-based frameworks to integrate the analysis of hybrid datasets.

The Sequence Distance Graph (SDG) framework implements a SequenceDistanceGraph representation that defines sequences in nodes and their adjacency in links, and an associated Workspace containing raw data and mappings. This provides an integrated working environment to use multiple sources of information to navigate and analyse genome graphs. Datastores allow random access to short, linked, and long read sequences on disk. A mapper on each datastore contains methods to map the reads to the graph and access the mapping data. KmerCounters provide functions to compute k-mer coverage over the graph from sequencing data, enabling coverage analyses. Additional DistanceGraphs, typically representing longer-range information and different linkage levels, define alternative topologies over the SequenceDistanceGraph nodes. Finally, a NodeView abstraction provides a proxy to a node, with methods to navigate the graph and access its mapped data. This comprehensive framework can be used to explore genome graphs interactively or to create processing methods for assembly or downstream analysis.

Here we describe the SDG implementation and basic tools, providing examples of use cases that highlight its analytic flexibility. First, we show how to create a hybrid assembly by integration of long reads linkage into a short-read graph. Then we analyse a simulated parent-child trio and show how the coverage of the parent datasets can be used to navigate the graph topology. These are only two of the multiple ways integrating data and genome graphs can be used to perform simple but powerful analyses.

Methods

Implementation

The C++ core library implements SDG’s data structures and methods for WorkSpaces, graphs, datastores and mappers. Its main goal is to provide a straightforward interface to project information from raw datasets onto graphs, and enable easy access and analysis of the graph-data combination. It uses OpenMP for parallel processing, and SWIG 4.0 to export a Python API to enable interactive data analysis.

The SequenceDistanceGraph class contains a vector of nodes defining DNA sequences, and a vector of links. Every node has a positive and a negative end, and links are defined between these node ends. Links with positive distances represent gaps between linked sequences and negative distances represent overlaps. This representation, shown in Figure 1, is similar to those presented in but unifies the concept of overlap and gap. Paths can be defined as list of nodes, with the sign of the first end in the walk. Graphs can be read and written to GFA and GFA2 files.

Figure 1. A simple Sequence Distance Graph with 5 nodes, including links with d<0, representing overlaps, and a link representing a gap of 10bp. Sequences appear in only one direction and their reverse complement can be obtained by traversing the node in opposite direction, from - to +. The two largest possible paths are [1, 2, 4, 5] and [1, -3, 4, 5], and their reverse complements [-5, -4, -2, -1] and [-5, -4, 3, -1] respectively.
The `DistanceGraph` class contains a set of links over the nodes of a `SequenceDistanceGraph` object. It is used to represent alternative sources of linkage information, such as longer range linkage produced by mapped reads for scaffolding.

The `WorkSpace` contains a single `SequenceDistanceGraph`, multiple `DistanceGraphs`, datastores and mappers, and its structure in memory represents the status of the SDG framework. It can be dumped and loaded from disk, providing persistence and checkpoints between different steps on SDG-based pipelines. Raw reads and `k-mer` counts are kept in separate files, pointed from the `WorkSpace`, to avoid duplication when using multiple `WorkSpaces` around the same dataset.

The `DataStores` and `Mappers` provide access and management to raw data and its mapping on the graph. `DataStores` do not load read data into memory, but rather provide random access to the on-disk data. The `PairedRedMapper` and `LinkedReadMapper` classes use a unique `k-mer` index to map reads to single nodes, with single reads mapping to multiple nodes not being mapped\(^1\)\(^2\)\(^3\). The `LongReadMapper` class generates multiple mappings from each read to nodes, using a short non-unique `k-mer` index (k=15 by default)\(^1\)\(^3\)\(^4\)\(^5\)\(^6\) Long read mapping filtering is left to later stages of the processing.

The `KmerCounters` creates an index with all the `k-mers` at a given k up to k=31 and counts occurrences of these `k-mers` on the graph, allowing them to count occurrences in datastores or fastq files. These counts, persisted in the `KmerCounter` with a name, can be then accessed to perform `k-mer` coverage analyses. Projections of raw `k-mer` coverage in the reads and the assembly over a particular sequence for a node or path, similar to those produced by the “sect” tool of K-mer Analysis Toolkit (KAT)\(^7\) are valuable for content analysis. Spectra analysis of these frequencies can provide further insight into genome composition and representation on the assembly.

Two processing classes, `LinkageUntangler` and `LinkageMaker`, work with alternative linkage configurations. The `LinkageMaker` is used to condense information via one of its `make_linkage*` methods, from evidence in the `WorkSpace` into links in a `DistanceGraph`. The `LinkageUntangler` class works on a `DistanceGraph` to simplify, condense and/or linearise its linkage. In the second use case below it can be seen how a combination of `LinkageMaker` and `LinkageUntangler` can be used for scaffolding with long reads.

Finally, the `NodeView` class, and its associated `LinkViews`, provide a single-entry point for node-centric analyses. A `NodeView` from either a `DistanceGraph` or `SequenceDistanceGraph` is a wrapper containing a pointer to the graph and a node id, and will provide access to its nodes’ previous and next linked nodes, mapped reads, or `k-mer` coverage. A user with good understanding of the `NodeView` class should be able to access most information in the `WorkSpace` through it, making it the default choice for analysing the graph.

**Operation**

**Requirements and installation.** SDG can be run on Linux and MacOS, and requires enough RAM to hold the `WorkSpace` completely in memory, which will depend on the dataset. Space to hold the uncompressed sequences on the datastores on disk will also be required.

SDG can be installed via pre-compiled binaries from https://github.com/bioinfologics/sdg/releases. The binaries have been built using Python3 and GCC version 6 from the Ubuntu package manager for the Linux version. The MacOS version dependencies were obtained using Homebrew (Python3, GCC-6 and SWIG). SDG can be compiled using CMake, Python3, SWIG version 4 and GCC version 6 onwards. Detailed instructions can be found at https://bioinfologics.github.io/sdg/sdg/README.html#installation.

**Typical workflow.** Working with SDG typically involves two different stages: creating a `WorkSpace` with the data and mappings, and analysing this `WorkSpace`. SDG includes command line tools to create `DataStores`, `KmerCounts`, and `WorkSpaces`, and map reads within a `WorkSpace`.

- `sdg-datastore`: creates a `Datastore` from raw reads and can process paired, 10x or long reads. An output prefix is specified as a parameter and a `<prefix>.prseq`, `<prefix>.lrseq` or `<prefix>.loseq` file is generated.
- `sdg-kmercounter`: creates a `KmerCounter` indexing a graph from a `WorkSpace` or GFA, or works with an already generated one. A count can be added directly from raw reads or from a datastore. The `KmerCounter` is persisted on file with extension `.sdgkc`.

...
• **sdg-workspace**: creates a WorkSpace from a base graph or works with an already generated one. Datastores and KmerCounters can be added. The WorkSpace is persisted on file with extension 'sdgws'.

• **sdg-dbg**: creates a WorkSpace from a PairedReadDatastore by building a deBruijn graph and using this as the base graph. Counts for the k-mers from the graph and raw reads are added too.

• **sdg-mapper**: maps reads within a WorkSpace. An updated WorkSpace is produced and dumped to the specified prefix.

WorkSpaces can also be instantiated with an empty graph, and the graph populated through the add_node and add_link methods. The following example on a python session shows how the simple graph from Figure I can be created from scratch, navigated through a NodeView instance and sequence from its paths extracted.

```python
>>> import pysdg as SDG
version 0.1 master b4d3f02
>>> ws=SDG.WorkSpace()
>>> ws.sdg.add_node("CTACGGA")
1
>>> ws.sdg.add_node("GACCTTA")
2
>>> ws.sdg.add_node("AATACGGTCC")
3
>>> ws.sdg.add_node("TTACGAA")
4
>>> ws.sdg.add_node("CTGATATGA")
5
>>> ws.sdg.add_link(-1, 2, -2)
>>> ws.sdg.add_link(-1, 3, -3)
>>> ws.sdg.add_link(-2, 4, -3)
>>> ws.sdg.add_link(3, 4, -2)
>>> ws.sdg.add_link(-4, 5, 10)
>>> nv=ws.sdg.get_nodeview(1)
>>> nv
<NodeView: Node 1 in SDG>
>>> nv.next()
<Vector: 2 LinkViews>
>>> print(nv.next())
[<LinkView: -3bp to Node -3>,
<LinkView: -2bp to Node 2>]
>>> nv = nv.next()[0].node()
>>> nv
<NodeView: Node -3 in SDG>
>>> print(nv.prev())
[<LinkView: -3bp to Node 1>]
>>> nv.sequence()
'GGACCGTATT'
>>> SDG.SequenceDistanceGraphPath(ws.sdg, [1, -3, 4, 5]).sequence()
'CTACCGGATATTAGAANNNNNNNNNNTGATATGA'
```

Typically, as shown in Figure 2, the API is used to explore a larger WorkSpace, with the methods accessing both in-memory and on-disk data, and modifying the status of the WorkSpace.

**Example use cases**

To illustrate the use of SDG, we have reproduced a short version of two examples from [http://bioinfologics.github.io/sdg_examples](http://bioinfologics.github.io/sdg_examples).
All paired end datasets are available on https://zenodo.org/record/3363871#.XUwyVy2ZN24, and the PacBio reads are from NCBI accession PRJNA194437. For simplicity, we have also made the datasets available on https://opendata.earlham.ac.uk/opendata/data/sdg_datasets/ as ready-to-use 'fastq.gz' files.

Hybrid assembly of short and long reads
This example is based on an E. coli dataset combining PacBio reads from 17 and Illumina Miseq 2x300bp reads subsampled from a test run. It uses the long reads to scaffold a short read based graph produced by sdg-dbg. Graphs are dumped to GFA files at different stages, and visualised using Bandage v0.8.1.

First, we use the command line tools to create datastores for both long and short reads and an initial WorkSpace containing a DBG assembly:

```bash
sdg-datastore make -t paired -o ecoli_pe ../ecoli_pe_r1.fastq.gz -2 ../ecoli_pe_r2.fastq.gz
sdg-datastore make -t long -o ecoli_pb -L ../ecoli_pb_all.fastq.gz
sdg-dbg -p ecoli_pe.prseq -o ecoli_assm
```

From this point on, we use the python SDG library. First, we load the workspace, add a long read datastore and map its reads using a k=11 index.

```python
import pysdg as SDG
# Load sdg-dbg's workspace from disk, add the pacbio datastore
ws = SDG.WorkSpace('ecoli_assm.sdgws')
lords = ws.add_long_reads_datastore('ecoli_pb.loseq')
```

**Figure 2. Structure of a WorkSpace and access via an Python interactive session.** The WorkSpace holds the information for a project and contains the graphs, the mappers and k-mer counts. From Python, a previously saved WorkSpace is loaded from disk (1). The NodeView object is centred on a specific node and can be used to access node characteristics (ie. size and sequence), graph topology from the perspective of the node you are on (i.e. neighbours in both directions (2)) and can also retrieve information projected onto the selected node (ie. mappings (3) and k-mer coverage (4)). Operations such as adding a KmerCounter to the WorkSpace and adding a count (5) can be performed, and the WorkSpace can be saved back to disk (6). Once loaded, the bulk of the WorkSpace is held in memory for fast access with the raw read data from the DataStores remaining on disk accessible through random access.
# Map long reads
lords.mapper.k = 11
lords.mapper.map_reads()

ws.sdg.write_to_gfal('initial_graph.gfa')

The graph, as shown in Figure 3A contains multiple unresolved repeats.

We can use the LinkageMaker to create linkage using the long reads datastore. We do this by selecting the nodes between which to analyse possible linkage, in this case all nodes of 1100bp or more, and then calling the make_longreads_multilinkage method, with alignment filtering parameters of 1000bp and 10% id.

```python
lm = SDG.LinkageMaker(ws.sdg)
mlg = lm.select_by_size(1100)
mldg = lm.make_longreads_multilinkage(ws.long_reads_datastores[0].mapper, 1000, 10)
```

This multi-linkage can be collapsed using the LinkageUntangler. The make_nextselected_linkage method links every selected node to its closest selected neighbours on each direction, aggregating the distances via a simple median calculation:

```python
lu = SDG.LinkageUntangler(mldg)
lm.select_by_size(1100)
ns_dg = lu.make_nextselected_linkage()
ns_dg.write_to_gfal('ns_collapsed.gfa')
```

The new graph we dumped, as shown in Figure 3B, has disconnected the repeats and introduced long read linkage which skips over them, but it is still not fully solved. We can improve this further by getting rid of repetitive nodes that will be connected to multiple neighbours, as each of them belongs in more than one place. We do that by just turning these nodes’ selection off in the LinkageUntangler, which will then skip them in the solution.

```python
for nv in ns_dg.get_all_nodeviews():
    if len(nv.prev()) > 1 or len(nv.next()) > 1:
        lu.selected_nodes[nv.node_id()] = False
ns_nr_dg = lu.make_nextselected_linkage()
ns_nr_dg.write_to_gfal('ns_nr_final.gfa')
```

The last graph is now a circle, with all the repeats disconnected from any linkage.
Analysing a simulation of heterozygous parent-child trio with short reads

We created a simulation of a trio dataset for this example using the synthetic genome creation and sequencing package Pseudoseq.j\textsuperscript{19} Chromosomes 4 and 5 of the reference genome of the yeast strain S288C were used as templates to create a diploid, genome for each parent with 1% heterozygous sites. Each homologous pair of chromosomes was crossed over and recombined and the child inherited one chromosome from the first parent at random, and one chromosome from the second parent at random. Simulated paired end reads were generated for each genome, using an average fragment length of 700bp and a read length of 250bp, and an expected coverage of 70x with error rate was set to 0.1%.

First we used the command line tools to create a graph from the child reads using sdg-dbg, and add \textit{k-mer} counts for both parents into the datastore.

```bash
sdg-datastore make -t paired -1 child/child-pe-reads_R1.fastq.gz -2 child/child-pe-reads_R2.fastq.gz -o child_pe
sdg-dbg sdg-dbg -p child_pe.prseq -o sdg_child
sdg-kmercounter add -c main.sdgkc -n p1 -f p1/pl-pe-reads_R1.fastq.gz -f pl1/pl-pe-reads_R2.fastq.gz -o main
sdg-kmercounter add -c main.sdgkc -n p2 -f p2/p2-pe-reads_R1.fastq.gz -f p2/p2-pe-reads_R2.fastq.gz -o main
```

We now open the WorkSpace and use the \texttt{NodeView::parallels} method to look for the largest bubble structure in the graph, which should be formed by two parallel nodes with haplotypes coming from each parent.

```python
import pysdg as SDG
ws = SDG.WorkSpace('sdg_child.sdgws')
# Largest node with one parallel node, and its parallel
maxbubble = 0
for nv in ws.sdg.get_all_nodeviews():
    if nv.size() > maxbubble and len(nv.parallels()) == 1:
        maxbubble = nv.size()
        bubble_nvs = (nv, nv.parallels()[0])
```

Since each side should be a haplotype from a different parent, we should see a loss of \textit{k-mer} coverage on the parent that didn’t contribute that haplotype. To check this, we create a plotting function to plot the output from the \texttt{NodeView::kmer_coverage} method.

```python
def plot_kcov(nv):
    '''Plot kmer coverage across the three read sets. Requires pylab.'''
    figure(); suptitle("Coverage for " + str(nv));
    subplot(3,1,1); ylim((0,120))
    plot(nv.kmer_coverage("main","PE"), label="child"); legend(loc=1);
    subplot(3,1,2); ylim((0,120))
    plot(nv.kmer_coverage("main","p1"), "red", label="parent 1"); legend(loc=1);
    subplot(3,1,3); ylim((0,120))
    plot(nv.kmer_coverage("main","p2"), "blue", label="parent 2"); legend(loc=1);

    plot_kcov(bubble_nvs[0])
    plot_kcov(bubble_nvs[1])
```

The plots, shown in Figure 4, reflect how Node 4775 contains content inherited from parent 2 and its parallel node 11414 contains content inherited from parent 1. We can create a function to extend these parent-specific regions by walking forward and backward as long as only one link takes us to a node that is fully covered by the content of the parent we are following.
def extend_parent_covered_path(starting_node, target_parent):
    if ws.sdg.get_nodeview(starting_node).kmer_coverage("main", target_parent).count(0) != 0:
        return SDG.SequenceDistanceGraphPath(ws.sdg, [])
    p = SDG.SequenceDistanceGraphPath(ws.sdg, [starting_node])
    for x in [0, 1]:
        nv = ws.sdg.get_nodeview(p.nodes[-1])
        while nv.next():
            next_node = 0
            for nl in nv.next():
                if nl.node().kmer_coverage("main", target_parent).count(0) == 0:
                    if next_node or nl.node().node_id() in p.nodes:
                        next_node = 0
                        break
                else:
                    next_node = nl.node().node_id()
            if next_node == 0:
                break
        p.nodes.append(next_node)
        nv = ws.sdg.get_nodeview(next_node)
    p.reverse()
    return p

path1 = extend_parent_covered_path(11414, "p1")
path2 = extend_parent_covered_path(4775, "p2")

After using this function, path1 contains 49 nodes yielding 8672bp of sequence inherited from parent 1, and path2 contains 139 nodes yielding 26351bp of sequence inherited from parent 2. It is important to note that the difference in node count and sequence length arises because the extension function is haplotype-specific and its results depend in the topology of each haplotype graph.

Summary
The Sequence Distance Graph framework provides a unified workspace for different sequencing technologies using the genome graph as the basis of integration. It enables analyses across the graph topology, the raw data and its projections to the graph. We have shown how the NodeView class can be used through the Python API to produce interactive analyses that are both powerful and easy to follow. We expect this will be a useful codebase for all levels of users, not only for the construction of graph-based analysis but also for their teaching and dissemination.

Data availability
Source data
The PacBio, *E. coli* reads are deposited on NCBI accession PRJNA194437 from Koren et al. 17

*E. coli* K12 Re-sequencing with PacBio RS and 454: Accession number PRJNA194437, https://identifiers.org/ncbi/bioproject:PRJNA194437

Figure 4. Trio analysis: k-mer coverage for each side of the largest bubble structure in the child’s assembly by each of the three read sets. Coverage drops to 0 on the opposite parent for k-mers that are unique to a parent.
Underlying data
The datasets used in the examples are available from: https://opendata.earlham.ac.uk/opendata/data/sdg_datasets/ and archived in Zenodo Zenodo: SDG Paper Datasets. http://doi.org/10.5281/zenodo.3363871

Data are available under the terms of the Creative Commons Zero “No rights reserved” data waiver (CC0 1.0 Public domain dedication).

Software availability
Software documentation: https://bioinfologics.github.io/sdg

Source archived source code at time of publication: 

Archived source code available from: http://github.com/bioinfologics/sdg

License: MIT License

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We would like to thank Richard Harrison for helpful discussions about SDG’s results and continued support through the OctoSeq project. We thank James Cuff for input about design principles and continuous encouragement. We thank Kat Hodgkinson for her feedback and patience as an early user of the rough alpha version of SDG. We thank Camilla Ryan for enduring and joining never-ending discussions about graph representations and the design of the framework.

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Max Alekseyev
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The authors describe a software package aimed at construction, storage, and manipulation of sequence graphs. The software is publicly available on Github. While the core functionality is developed in C++, the package also provides Python wrapper library and a command-line tool. The authors outline the software design and provide pipelines and code examples for two different types of data.

Overall, the paper is well-written and describes potentially useful software. At the same time, the paper lacks:

1. Comparison of (features of) the developed software and existing software such as VG toolkit based on variation graphs (reference 5);

2. Discussion of the software applicability (e.g., will it work on large or repeat-rich genomes?), or estimation of its running time/space complexity.

Some minor comments:

1. The use of bold font is not explained. For example, Datastores first appears in the sentence “Datastores allow random access...“ describing its features, but WHAT is Datastores?

2. Orthogonal edge routing in Fig. 1 is somewhat confusing, why not make edges curved?

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Publisher Full Text

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

Is the description of the software tool technically sound?
Yes

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?
Partly

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

**Reviewer Expertise:** genome assembly, comparative genomics

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

Reviewer Report 04 November 2019

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The authors describe a framework for constructing sequence graphs, aligning reads, manipulating graph structures, and extracting them into standard formats (e.g., GFA). This framework is available as both a set of command line tools and a python library which wraps much of the underlying functionality. Their implementation unifies the representation of gaps and overlaps as a single linkage type within the graph. This is the primary theoretical advance of the work. This work is scientifically sound but its description as written could benefit from some minor additions.

The software is freely available on GitHub and binary releases of the command line tools are provided. These are functional on a modern linux laptop and clear examples with data are
provided. The paper includes the outputs of these examples as figures.

The python libraries rely on SWIG and are not included in the binaries. While not requisite for publication, providing the python libraries through pip, conda, or another package manager would increase the reach of the framework. This would match the authors' conclusion that the sdg package provides "a useful codebase for all levels of users."

The examples provided are clear and scientifically relevant. The graph mapping and manipulation example using *E. coli* data (Figure 3) and the description of genotyping a simulated yeast trio (Figure 4) are both realistic.

However, the authors should provide run times and machine details for these examples. Both are relatively fast as the datasets are small. There is no need for extensive benchmarking; a footnote for each example would address this adequately.

A brief 1-2 sentence discussion of a larger scale example the authors have attempted should also be included.

In addition, the phrasing "simulated parent-offspring trio" in the abstract should be modified to make it clear that the data is from yeast. As it is written the phrasing implies the framework may work on human/animal-scale data, though no evidence of this has been provided in this version of the paper.

Lastly, a brief description of the similarities and differences between the sequence (distance) graph, the variation graph, and the de Bruijn graph from an assembler such as ABySS should be included in the introduction or provided by a reference. This description need not be longer than two to four sentences in length. This should highlight the similar representations of the graphs (e.g., sequences stored in nodes and linkages/paths described by edges) and the different amounts of information content within the graph types. This would strengthen the critical need for the software and is partially highlighted by the example in Figure 3.

As it stands the paper is deserving of indexing. These additions would further strengthen what is already an excellent tool description, I hope without adding too much additional work for the authors.

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

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

**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: computational biology; graph genomes; structural variant calling; bioinformatics; cancer biology

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.

Reviewer Report 24 September 2019

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The authors demonstrate a new toolchain and data model for working with sequence graphs. This method allows the user to dynamically interact with sequence graphs made in the process of assembly. They provide a number of examples of the use of the method as well as code snippets to demonstrate its functionality. The library is written in C++, but wrapped in python with SWIG, which should make it useful to many researchers for whom C++ is difficult to use.

I find only one thing strange about the work. In the beginning, the authors indicate that there are not interoperable methods for working with sequence graphs and alignments to them, but they have in effect created another competing standard. Are there particular limitations with existing data models that they hope to address with the Sequence Distance Graph framework? How is their model different than the variation graph model, in which distances are provided by a collection of paths (or equivalently alignments) embedded within the sequence graph?

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

Is the description of the software tool technically sound?
Yes

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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.

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.

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