



# tskit: WORKING WITH TREE SEQUENCES

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Code: [github.com/petrelharp/progen-2023](https://github.com/petrelharp/progen-2023).

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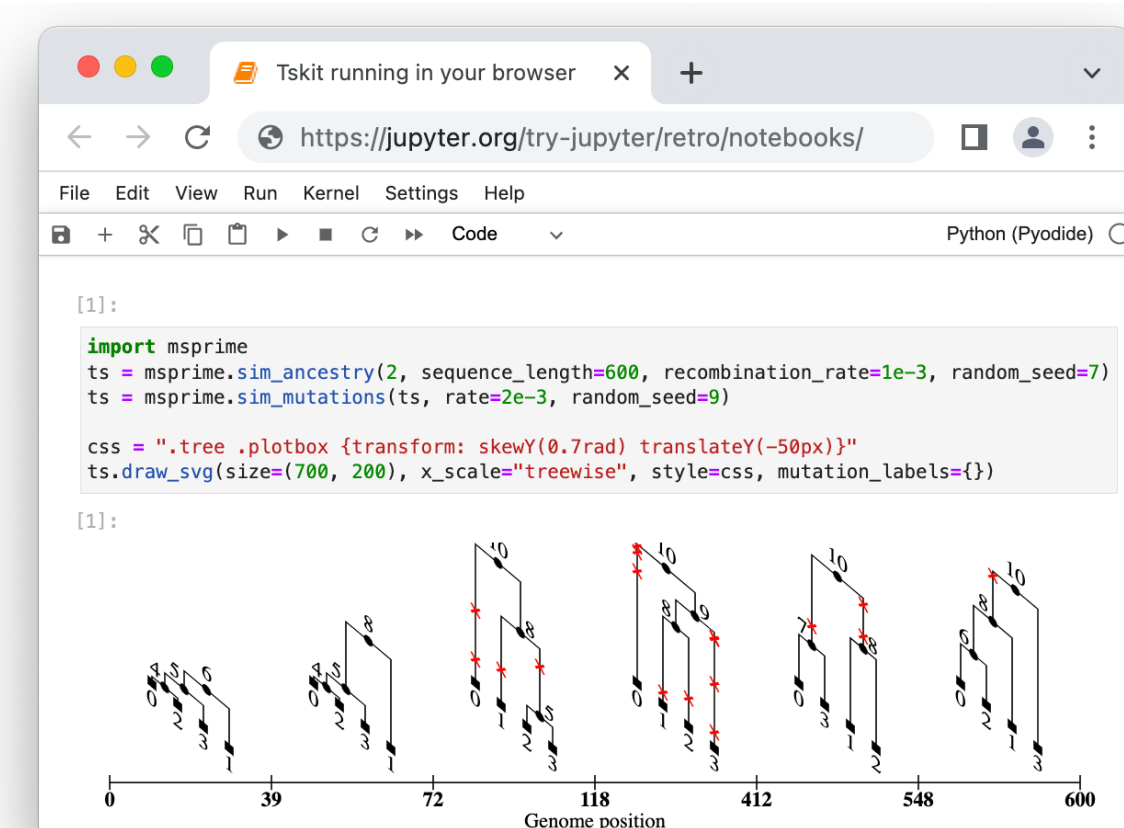
## INTERFACES AND INTEROPERABILITY

We aim to provide *stable*, *well-tested* and *well-documented* software so others can reliably build with it – including a *backwards compatibility* guarantee. **tskit** is already being used in a number of inference and simulation packages. The core functionality is implemented via a C API, and the primary interface is via a Python library, but others are available:

- Well-tested API used by many software packages: SLiM, fwdpy11, msprime, tsinfer/tsdate, Relate, slendr, etc.
- Available in multiple programming languages:



- Runs in-browser (no install required!) for quick demos / teaching (see screenshot).
- Can represent full Ancestral Recombination Graphs; includes ARG likelihood calculations.
- Interoperable with other packages (e.g., VCF output for sequence data, newick/nexus output to Dendropy, numpy arrays to scikit-allele)



## METADATA

**tskit** now has integrated metadata for all objects (genomes, mutations, sites, etc). For instance (*spoiler alert*), the complete ARG for 1.26 million SARS-Cov-2 genomes: fits in 57MB! Loads in under 1 second!

```
ls -lh SARS-Cov-2-ARG.ts.tsz

-rw-rw-r-- 1 jk jk 57M Mar  2 13:32 SARS-Cov-2-ARG.ts.tsz

ts = tszip.decompress("SARS-Cov-2-ARG.ts.tsz")

CPU times: user 775 ms, sys: 533 ms, total: 1.31 s
Wall time: 842 ms
```

ts kit	Tree Sequence	Table	Rows	Size	Has Metadata
Trees	1496	Edges	1458146	44.5 MB	
Sequence Length	29904.0	Individuals	0	24 Bytes	
Time Units	days	Migrations	0	8 Bytes	
Sample Nodes	1265685	Mutations	1213193	45.8 MB	✓
Total Size	819.3 MB	Nodes	1453347	716.5 MB	✓
Metadata	No Metadata	Provenances	1	874 Bytes	
		Sites	29422	1.4 MB	✓

**The integrated data model** links nodes, edges, sites and mutations, and now allows annotation of all objects with arbitrary external metadata. For instance, here's the first five sites in the SARS-Cov-2 ARG, and all three mutations at the first site:

ts.tables.sites[:5]					ts.tables.mutations[ts.mutations_site == 0]						
id	position	ancestral_state	metadata		id	site	node	time	derived_state	parent	metadata
0	56	G	{masked_samples: 727232}		0	0	171200	181	A	-1	{}
1	57	A	{masked_samples: 726137}		1	0	331354	143	C	-1	{}
2	58	T	{masked_samples: 725063}		2	0	1026732	60	T	-1	{}
3	59	C	{masked_samples: 724533}								
4	60	T	{masked_samples: 721663}								

## OVERVIEW

**tskit** is the C and python library providing tools for working with *succinct tree sequences*. We provide solid, stable, well-tested software for you to use and build on. Why might you want to use tree sequences?

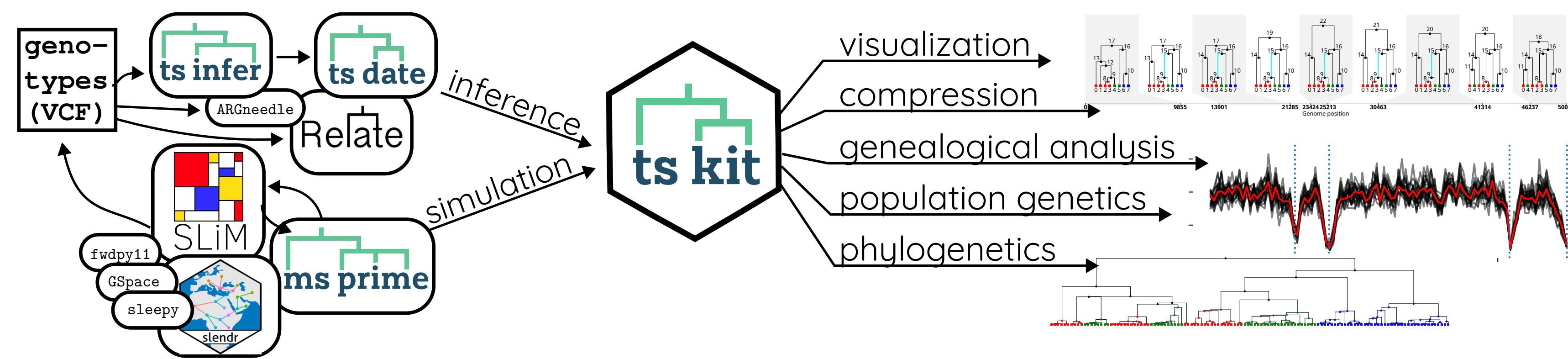
- For large samples, stores genotypes losslessly using (estimated) underlying genealogical relationships in orders of magnitude less space,
- ... and allows fast processing and exploration, in seconds, not hours.
- Genealogical relationships – “the trees” – are often closer to things we want to learn about
- ... and explicitly include a *time dimension*.
- History of a process can be recorded in a simulation, not just the genotypic outcome,
- ... and simulations can be much faster/more efficient.

In summary: by representing genomes using the genealogical process that generated the data, we get both a huge advantage storing and manipulating genomic data, as well as a more direct look at the processes that generated the data.

Here's some **silly slogans**, care to suggest any more?

- “**tskit**: launching your genomes into the time dimension!”
- “**tskit**: tree thinking, for popgen”
- “**tskit**: stable software for genome-scale trees”
- “**tskit**: all your insights, much faster!”

Documentation and examples: <https://tskit.dev/>

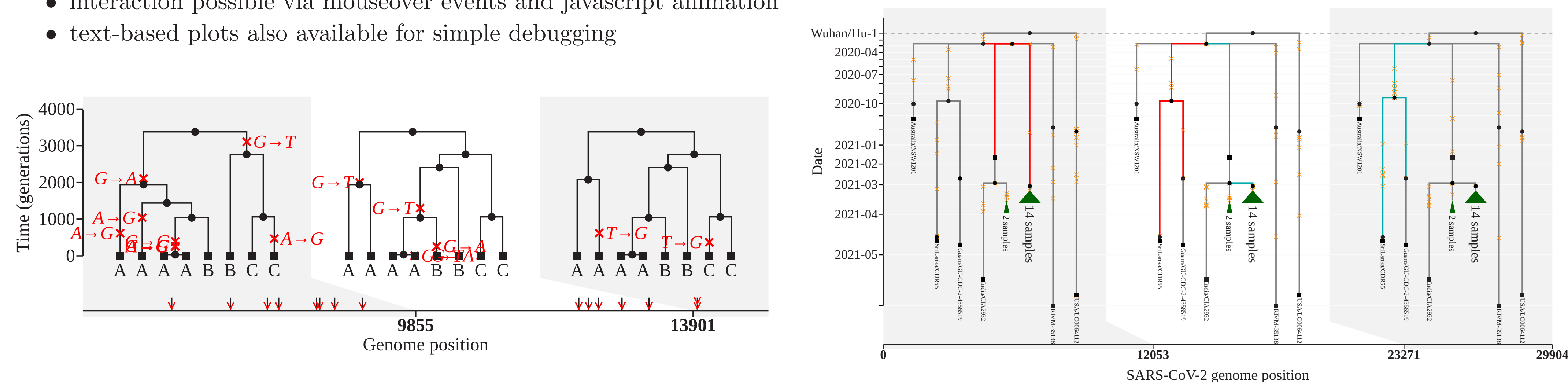


## VISUALIZATION – see more at <https://tskit.dev/tutorials/viz.html>

SVG-based visualization allows flexible styling of local trees.

Example:

```
simpl_ts.draw_svg(
    size=(800, 400), canvas_size=(850, 405),
    style=style + ".join(node_styles)",
    y_axis=True, time_scale="log_time",
    symbol_size=4.5, y_label="Date",
    x_label="SARS-CoV-2 genome position",
    y_ticks=y_ticks, mutation_labels={},
    y_gridlines=True, node_labels=node_labels,
    root_svg_attributes={"id": "ns_rec"},
)
```



## CONTRIBUTORS

**tskit** is developed by an open and inclusive community. Want to get involved? All skill levels welcome – email us at [admin@tskit.dev](mailto:admin@tskit.dev).

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## STATISTICS

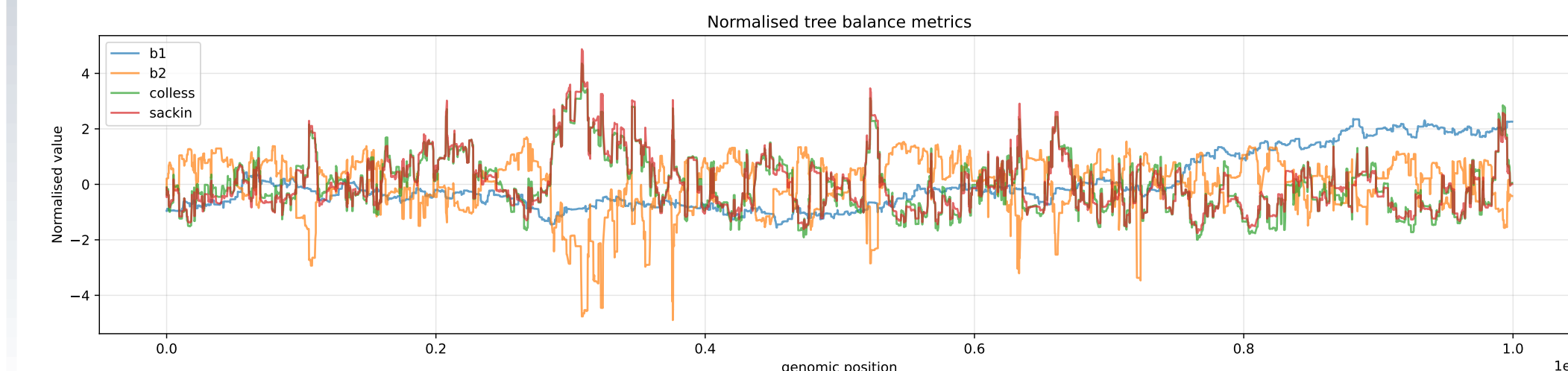
**tskit** lets you perform efficient calculations of statistics along the genome, often many times quicker than other software! You may be interested in calculating:

- the allele frequency spectrum or statistics derived from it, like nucleotide diversity, Tajima's  $D$ ,  $f_4$ , ...
- IBD-based quantities,
- summaries of tree topology, e.g., genealogical nearest neighbours and tree balance metrics,
- cross-coalescence rates (coming soon!)

**Example:** newly-implemented tree balance statistics. A balanced (binary) tree is perfectly symmetric in some way: each node has an equally sized subtree descending from its left- and right- branches, where ‘size’ is determined by some metric involving the tree's nodes and edges. **tskit** now implements several different metrics to calculate how unbalanced each tree is:

```
imb = pd.DataFrame({
    "genomic_position": [t.interval[0] for t in ts.trees()],
    "b1": [t.b1_index() for t in ts.trees()],
    "b2": [t.b2_index() for t in ts.trees()],
    "colless": [t.colless_index() for t in ts.trees()],
    "sackin": [t.sackin_index() for t in ts.trees()]
}).set_index("genomic_position")
```

```
imb = ((imb - imb.mean()) / imb.std())
imb.plot(figsize=(16, 4), alpha=0.7)
```



## NOTABLE NEW FEATURES

**tskit**'s contributors are actively working on new features, bug fixes, and improvements to the usability of existing features. Here's a shortlist of some recent additions:

**Reference sequences** By default, the sites in a tree sequence only define ancestral nucleotides at polymorphism sites. Remaining positions can now be specified using the `TreeSequence.reference_sequence`, and individual sample alignments can be obtained with the new `TreeSequence.alignments()` iterator.

**Structural operations** We've expanded the set of utility functions for large edits on tree sequences. For instance, the `TreeSequence.decapitate` method removes all parts of a tree sequence that are older than some user-specified time, and `TreeSequence.union` allows joining together of separate tree sequences to allow parallel simulation across different branches of a phylogenetic tree.

**Efficient array access** The relationships between nodes in each tree can now be extracted as numpy arrays. When used in conjunction with numba, Python-based calculations on the trees can act as speedily as machine-level code. Here is pure python computing total branch length just as fast as the “built-in” method (implemented in C):

