


Genome analysis

# Ribbon: intuitive visualization for complex genomic variation

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## Abstract

**Summary:** Ribbon is an alignment visualization tool that shows how alignments are positioned within both the reference and read contexts, giving an intuitive view that enables a better understanding of structural variants and the read evidence supporting them. Ribbon was born out of a need to curate complex structural variant calls and determine whether each was well supported by long-read evidence, and it uses the same intuitive visualization method to shed light on contig alignments from genome-to-genome comparisons.

**Availability and implementation:** Ribbon is freely available online at <http://genomeribbon.com/> and is open-source at <https://github.com/marianattestad/ribbon>.

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**Supplementary information:** [Supplementary data](#) are available at *Bioinformatics* online.

## 1 Introduction

Visualization has played an important role in the genomic revolution, enabling scientists to investigate variants, expression patterns, evolutionary changes and a number of other relationships (Kent *et al.*, 2002; Krzywinski *et al.*, 2009; Robinson *et al.*, 2011). Long-standing genome browsers have taken a very useful reference-based perspective, which is largely sufficient for representing short-read alignments. However, advances in long-read sequencing (Sedlazeck *et al.*, 2018) have shed light on examples where visualizing the portions of reads that maps to different parts of the reference is crucial. When a read is over 10 kb long, it can cover large compound structural variations on several loci, but that context is lost using IGV (Robinson *et al.*, 2011) or other reference-centric viewers without adding a read-based perspective and showing all relevant reference loci. In recent years, other visualization tools have tackled some challenging aspects of validating structural variant calls. In particular, SV-plaudit (Belyeu *et al.*, 2018) and SVCurator (Chapman *et al.*, 2019) have enabled crowd-sourced curation of structural variants, and svviz (Spies *et al.*, 2015) has used realignment against a putative structural variant allele to better evaluate the evidence. Even with these very useful approaches, more complex patterns of structural variation remain hard to represent. A more flexible approach can be helpful by showing the full picture of alignments from both the reference and read perspective, allowing researchers to explore and communicate about complex variation in a more intuitive way. The true power of long reads is hidden when the tools we use are built

for short reads. We have addressed this problem by creating Ribbon (<http://genomeribbon.com>) an interactive online visualization tool that displays alignments along both reference and query sequences, along with variant calls, genes and other genomic features.

## 2 Materials and methods

Ribbon can load alignments from SAM (Li *et al.*, 2009) and BAM files with long, short or paired-end reads. While the main strengths of Ribbon are more apparent with long sequences, short and paired-ends reads are also supported and can benefit from the more structurally focused visualization Ribbon provides (Supplementary Fig. S9). BAM files can be read from either a local file on the user's computer or from a remote server by entering the file's URL (Miller *et al.*, 2014). To support parsing local BAM files, we compiled the command-line utility SAMtools (Li *et al.*, 2009) from C to WebAssembly (Haas *et al.*, 2017), which allows us to process BAM files efficiently inside a web browser.

In addition to long-read alignment, Ribbon also has features for the related field of whole-genome alignment visualization. First, we added support in Ribbon for visualizing genome-to-genome alignments, such as for comparing genomes of two related species or comparing a new genome assembly to an existing reference. This is done by supporting a simple tab-delimited, human-readable coordinate file format that can be created by the whole genome aligner MUMmer (Kurtz *et al.*, 2004; Robinson *et al.*, 2011) or by scripting

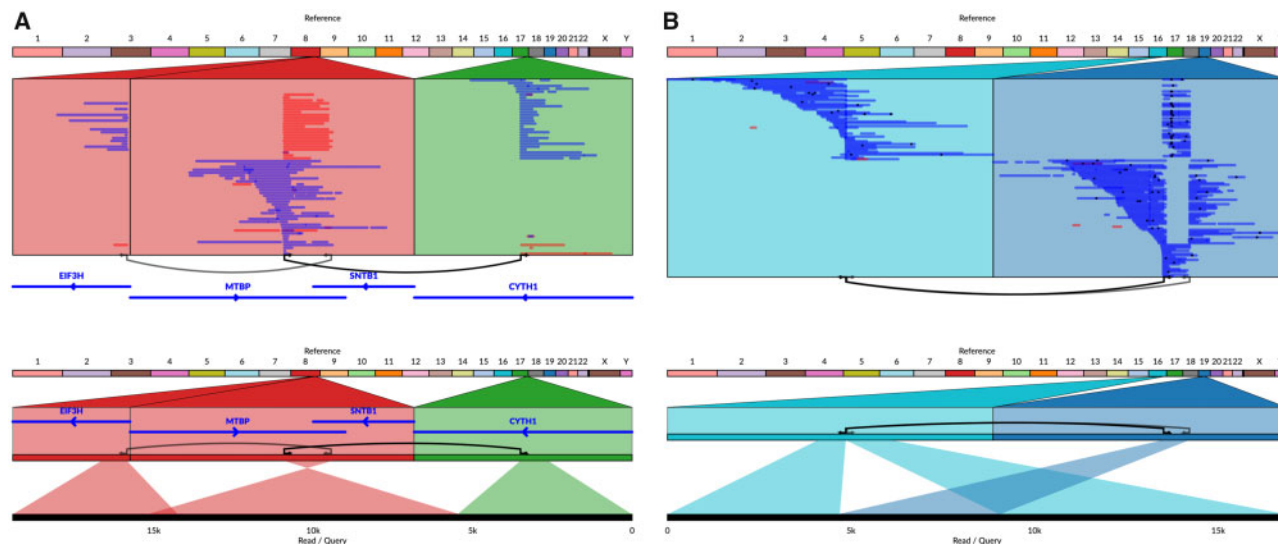


Fig. 1. Ribbon shows all alignments from the perspective of both the reference and the read or query. Here, long reads from SMRT sequencing show evidence for complex variants in the SK-BR-3 breast cancer genome. (A) The CYTH1-EIF3H gene fusion takes place through a series of two events that are captured within some of the long reads. (B) Long reads from SMRT sequencing show evidence of a full interchromosomal translocation where a sequence is homozygously deleted from chromosome 19 and homozygously inserted into chromosome 16

from another file format. Second, we adopted a version of the dot plot visualization as an optional alternative to the classic ribbon visualization, since dot plots are often used in the genome assembly/comparison field. The ribbon and dot plot visualizations show the same coordinates in different ways, but most users have found the ribbon visualization more intuitive, while the dot plot remains a common visualization method in the field of whole-genome alignment.

Finally, in addition to alignments with SAM, BAM and general coordinate files, Ribbon can also show variants in VCF or BEDPE format, and genes or any other features in BED format. The user can select a variant to jump to that locus (or both loci for two-breakpoint variants) in the genome, and all the reads with alignments in the given region(s) will show up, including all their alignments to other places in the genome. This is particularly powerful because it pulls other relevant regions into focus; for example, in Figure 1A, the left-most locus is included automatically because it contains alignments of reads also found at the other two loci.

### 3 Results

By showing a synchronized read and reference perspective, Ribbon shows patterns in alignments of many reads across multiple chromosomes, while allowing detailed inspection of individual reads (Supplementary Note S1). For example, here we show a gene fusion in the SK-BR-3 breast cancer cell line linking the genes CYTH1 and EIF3H revealed by PacBio long-read sequencing (Nattestad et al., 2018). While it has been found in the transcriptome previously (Chen et al., 2013; Edgren et al., 2011; Kim and Salzberg, 2011), genome sequencing did not identify a direct chromosomal fusion between these two genes. Using long-read sequencing, Ribbon shows that there are indeed reads that span from one gene to the other, going through not one but two variants, for the first time showing the genomic link between these two genes (Nattestad et al., 2018) (Fig. 1A). More gene fusions of this cancer cell line are investigated with Ribbon in Supplementary Note S2. Figure 1B shows another complex event in this sample made simple in Ribbon: the translocation of a 4.4-kb sequence deleted from chr19 and inserted into chr16.

With the support for genome-genome alignments, Ribbon can also be used to test assembly algorithms or inspect the similarity between species. Supplementary Note S4 shows a comparison of

gorilla (Gordon et al., 2016) and human genomes using Ribbon, highlighting major structural differences.

### 4 Discussion

Ribbon enables understanding of complex variants, and it may also help in the detection of sequencing and sample preparation issues, testing of aligners and variant-callers and rapid curation of structural variant candidates (Supplementary Note S3). Ribbon is a powerful and versatile visualization tool for investigating complex structural differences between any two genomes, using intuitive methods that take advantage of the rich context of long reads and contigs.

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*Conflict of Interest:* During the initial work, M.N. was a contractor and C.-S.C. was an employee and stockholder of Pacific Biosciences, a company commercializing DNA sequencing technologies. M.N. is currently an employee and stockholder of Google. C.-S.C. is currently an employee and stockholder of DNAnexus. R.A. is an employee and stockholder of Invitae. The other author declared no conflict of interest.

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