PyBamView: a browser-based application for viewing short read alignments

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Abstract

Summary: Current sequence alignment browsers allow visualization of large and complex next-generation sequencing datasets. However, many of these tools provide inadequate display of insertions and can be cumbersome to use on large datasets. I implemented PyBamView, a lightweight Web application for visualizing short read alignments. It provides an easy-to-use Web interface for viewing alignments across multiple samples, with a focus on accurate visualization of insertions.

Availability and Implementation: PyBamView is available as a standard python package. The source code is freely available under the MIT license at https://mgymrek.github.io/pybamview.

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Supplementary information: Supplementary data are available at Bioinformatics online.

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1 INTRODUCTION

The rapid growth of next-generation sequencing (NGS) technologies has led to a wide variety of short read DNA datasets. Manual inspection of sequence alignments is an important aspect of quality control. While the majority of NGS analyses have focused on single nucleotide polymorphisms (SNPs), recent bioinformatics advances allow analysis of more complicated variants, such as small insertions or deletions (Montgomery et al., 2013), larger structural variants (Ye et al., 2009) and short tandem repeats (Gymrek et al., 2012; Highnam et al., 2013). Furthermore, widely used genome engineering techniques, such as the CRISPR-Cas9 system (Cong et al., 2013) can often produce a wide range of complex variants. In these cases, visualization of insertion and deletion events is a particularly critical analysis step.

Current genome browsers, such as the UCSC Genome Browser (Kent et al., 2002) and the Integrative Genomics Viewer (IGV) (Robinson et al., 2011), offer visualization of alignments from SAM/BAM files across multiple samples and integration of many layers of genomics datasets. However, most existing tools have two important limitations. First, most are based on alignments to an ungapped reference sequence, which provides inadequate visualization of insertions. The SAM specification supports a padded reference, which captures multiple sequence alignment information and results in accurate insertion display by most browsers. However, most BAM files consist of pairwise alignments of short reads to a reference and do not use this feature. As a result, insertions are represented by an icon such as a vertical bar, which does not provide any visual information about the size or sequence of the inserted nucleotides. Second, the majority of alignment browsers are cumbersome to use, especially to visualize the large datasets typical of NGS experiments. They either require that the user upload large data files to a remote server or involve complicated installation and large resource requirements to run locally.

Several alignment browsers, such as Bambino (Edmonson et al., 2011), Consed (Gordon and Green, 2013) and the text-based SAMtools (Li et al., 2009) tview, overcome these limitations: they display the sequence of insertions even when using the standard ungapped reference, and are run locally with relatively low system requirements. However, tview does not allow the user to view multiple BAM files at once, and none of these tools allow for exporting alignments as snapshots or for sharing alignments remotely through a Web browser.

Here, I present PyBamView, a lightweight Web application for viewing alignments from BAM files. PyBamView provides alignment visualizations that accurately represent SNP, insertion and deletion events that can easily be exported to create publication-ready figures. It runs locally from the command line with minimal resource requirements and displays alignments in a Web browser. This interface allows users to quickly view alignments locally and to easily share alignments with local or remote collaborators.

2 BASIC USAGE AND FEATURES

PyBamView is a Python-based Web application that is run from the command line. Users provide PyBamView with a directory containing indexed BAM files and an optional reference genome in fasta format:

```
pybamview --bamdir DIRECTORY/WITH/BAMS --ref REF.fa
```

PyBamView will start a small Web server that can be accessed locally in a Web browser. Optional arguments can serve the application over a different address, for instance, for sharing...
Additional features are described in the Supplementary Text.

**3 EXAMPLE USE CASES**

Alignment visualization is a critical step of any sequencing experiment. Here, I show three examples where PyBamView provides useful visualization of sequence variants. Use cases are not limited to these examples and can theoretically include any ‘-seq’ experiment that can be represented by a BAM file.

First, it provides accurate visualizations of different length insertions, such as different alleles of a tandem repeat (Supplementary Fig. S2a). Furthermore, zooming out allows for visualization of large repeat expansions, such as a 60bp CAG expansion in Huntington’s Disease (Supplementary Fig. S2b, simulated 250bp reads).

Second, it can be used to analyze variation across samples. This is useful in such analyses as comparing matched tumor versus normal samples or looking for mutations in affected versus non-affected individuals in disease genetic studies. Supplementary Figure S3 shows example comparisons of individuals at a SNP, small insertion and a large deletion spanning several kb.

Third, it can visualize complex mutations generated by genome engineering technologies such as CRISPR-Cas9 (Cong et al., 2013). Dissecting these mutations requires adequate visualization of indels. An example alignment from a CRISPR library is shown in Supplementary Figure S4.

**4 IMPLEMENTATION**

PyBamView is implemented as a Python-based Web application using the Flask library. Alignments are processed using a Python backend, which then generates HTML, Cascading Style Sheets (CSS) and JavaScript files that are displayed in the Web browser. PyBamView takes advantage of BAM and fasta indexing to avoid loading large files into memory. It uses the pysam and pyfasta libraries for parsing BAM and fasta files, respectively. Both libraries use efficient index data structures, which allow them to quickly fetch data from specific genomic regions of interest. Read alignments are parsed from the CIGAR strings in the BAM file and are displayed as simple HTML tables as Scalable Vector Graphics (SVG) elements using Javascript. All CIGAR options reported in the SAM specification, including the padded reference option, are supported (Supplementary Fig. S5).

**5 CONCLUSION**

As the use of NGS to analyze complex genomic events grows, there is a critical need for accurate and easy-to-use visualization tools. PyBamView provides a simple, yet powerful, interface for alignment visualization that facilitates collaborative data analysis.

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