**Abstract**

**Motivation:** Genome browsers that support fast navigation through vast data sets and provide interactive visual analytics functions can accelerate research and help scientists achieve deeper insight into data. Previously, we developed Integrated Genome Browser (IGB), a highly configurable, interactive and fast open source desktop genome browser.

**Results:** Here we describe multiple updates to IGB, including all-new capability to display and interact with data from high-throughput sequencing experiments. To demonstrate, we describe example visualizations and analyses of data sets from RNA-Seq, ChIP-Seq, and bisulfite sequencing experiments. Understanding results from genome-scale experiments requires viewing the data in the context of reference genome annotations and other related data sets. To facilitate this, we enhanced IGB’s ability to consume data from diverse sources, including Galaxy, Distributed Annotation, and IGB-specific Quickload servers. To support future visualization needs as new genome-scale assays arise and enter wide use, we transformed the IGB codebase into a modular, extensible platform for developers to create and deploy all-new visualizations of genomic data.

**Availability:** IGB is open source and is freely available from http://bioviz.org/igb.

**Contact:** alorraine@uncc.edu

---

**1 Introduction**

Genome browsers are visualization software tools that display genomic data in interactive, graphical formats. Since the 1990s, genome browsers have played an essential role in genomics, first as tools for building, inspecting, and annotating assemblies and later as tools for distributing data to the public (Durbin and Thierry-Mieg, 1991; Harris, 1997; Kent, et al., 2002). Later, the rise of genome-scale assays created the need for a new generation of genome browsers that could display user’s experimental data alongside reference sequence data and annotations.

Integrated Genome Browser (IGB), first developed in 2001 at Affymetrix, was among the first of this new breed of tools. IGB was first written to support Affymetrix scientists and collaborators who were using whole genome tiling arrays to probe gene expression and transcription factor binding sites as part of the ENCODE project (Kapranov, et al., 2003). As such, IGB was designed from the start to handle and display what we would now “big data” in bioinformatics - millions of probe intensity values per sample. IGB was one of the first genome browsers to support visual analytics, in which interactive visual interfaces augment our natural ability to notice patterns in data. The “thresholding” feature described below is an example.

Because early IGB development was publicly funded, Affymetrix released IGB and its companion graphics library, the Genoviz Software Development Kit (Helt, et al., 2009), as open source software in 2004. Our first article introducing IGB appeared in 2009 (Nicol, et al., 2009) and focused on visualization of tiling array data. Here, we describe new visual analytics and data integration features developed for high-throughput sequencing data. We also introduce a new plug-in application programming interface (API) that makes adding new functionality easier for developers, transforming IGB into an extensible visual analytics platform for genomics.
2 Results

IGB is implemented as a stand-alone, rich client desktop program using the Java programming language. To run IGB, users download and run platform-specific installers; these also support automatic updates. Mac and Windows installers also include a copy of the Java virtual machine, which is installed in an IGB-specific location, making it unnecessary for users to install (and maintain) Java separately.

IGB’s implementation as local application rather than as a Web app means that IGB can access the full processing power of the user’s local computer. IGB is always present on the user’s desktop, regardless of internet connection status, but IGB can also use the Web to consume data, as describe below. Because IGB runs locally, users view their own datasets without uploading them to a server, which can be important when working with confidential data. IGB’s implementation in Java means that developers benefit from Java’s robust support for multi-threaded, object-oriented programming and dozens of well-tested, robust libraries and tools that are available for Java.

3.1 Viewing genomes and annotations

On startup, IGB displays a “home” screen featuring a carousel of images linking to the latest versions of model organism, crop plant, and reference human genome assemblies. More species and genome versions are available via menus in the Current Genome tabbed panel, including more than seventy animal, plant, and microbial genomes. Users can also load and visualize their own genome assemblies, called “custom” genomes in IGB, provided they have a reference sequence file in FASTA or 2bit format.

Once a user selects a species and a genome version, IGB automatically loads the reference gene model annotations for that genome, if available. For any given genome version, additional annotations or high-throughput data sets may also be available via the Data Access Panel interface. These data sets can come from different sites, and to highlight this, IGB can display a favico.ico graphic distinguishing these different data sources. This is why IGB is named “integrated” - it integrates data from different sources into the same view. To illustrate, Figure 1 shows an example view of integrated data sets from the human genome. Reference gene model annotations and other data sets from IGB Quickload are shown, together with tracks loaded from a DAS1 server hosted by the UCSC Genome Bioinformatics group.

3.2 Navigating and interacting

Genomic data sets span many scales, and fast navigation through these different scales is a key feature for a genome browser. Users need to be able to quickly travel between base-level views depicting sequence details like splice sites, gene-level views depicting the exon-intron organization of genes, and chromosome-level views showing larger-scale structures, like centromeres and chromosome bands. Tools that support fast navigation through the data accelerate the discovery process.

For this reason, IGB implements a visualization technique called one-dimensional, animated semantic zooming, in which objects change their appearance in an animated fashion around a central line, called the “zoom focus” (Lorraine and Helt, 2002). Animation helps users stay oriented during zooming and can create the pleasant impression of flying through one’s data (Bederson and Boltman, 1999; Cockburn, et al., 2009). To implement animated zooming, IGB uses graphical components from the BioViews Software Development Kit, an updated and open source version of the BioViews library (Helt, et al., 1998; Helt, et al., 2009).

To better simulate the experience of flying through a genome, IGB introduces a simple but innovative technique in which users set the zoom focus by clicking locations in the display. A vertical, semi-opaque line called a “zoom stripe” indicates the zoom focus position and also serves as a pointer and guideline tool when viewing sequence or exon boundaries. When users zoom, the display appears to contract or expand around this central zoom stripe, which remains in place, thus creating a feeling of stability and control even while the virtual genomic landscape is rapidly changing. Animated zooming combined with settable zoom focus help users to move quickly across multiple scales without getting lost, saving time and enhancing their ability to

![Image of IGB viewing multiple human genome data sets](image.jpg)

**Fig. 1.** Viewing multiple human genome data sets in IGB. IGB screen showing human genome build 38, released in December 2013. Gene annotations load by default, with additional data available in the section labeled Available Data (lower left). Each dataset occupies a separate track within the main window, and can be colored according to user preference. Right-clicking items in the main window activates a context menu with options to search Google, run BLAST, or view the underlying genomic sequence.
Fig. 2. RNA-Seq data from human lung adenocarcinomas bearing mutant or wild-type (WT) alleles of the KRAS oncogene. (a) Coverage depth graphs show transcript abundance across a 250 kb region. Mutant samples contain a peak indicating higher expression in the mutant sample. (b) Overlaid depth graphs showing a discontinuity in coverage indicating differential splicing in PFN2. Quantification of split reads by FindJunctions further supports differential splicing. (c) Zoomed in view of (b), showing aligned reads.

notice meaningful patterns in data. To our knowledge, no other genome browser offers this fine-tuned control over zooming.

IGB also supports “jump zooming”, a more common implementation of zooming in genome browsers. In jump zooming, a request to zoom triggers what feels like instantaneous teleportation to a new location. To jump-zoom to a gene model or other feature, users double-click it. To jump-zoom to a region, a users click-drag across the region in the IGB coordinates axis. Users can also quickly move to a gene or feature by searching for it using the Advanced Search tab or the quick search box in the top left of the display.

Panning interactions are also important for navigation, and IGB supports multiple ways to move (pan) without changing the scale. Borrowing from many other graphical tools, IGB implements a move tool cursor that users click-drag to move the display in any direction. Clicking arrows in the toolbar move the display from left to right, and vertical scrollbars offer ways to rapidly move in vertical or horizontal directions. Users can also pan by click-dragging the selection (arrow) cursor into the left or right border of the main display window.

IGB helps users ask and answer questions about their data by supporting multiple ways for users to interact with what they see. Selecting data display elements (Glyphs) triggers display of meta-data about the selected item and mouse-over causes a tooltip to appear. Right-clicking items within tracks activates a context menu with options to search Google, run a BLAST search at NCBI, or open a sequence viewer (Fig. 1). Right-clicking track labels activates context menus showing a rich suite of visual analytics tools and functions, some of which we discuss in more detail below.

3.3 Loading data from files or URLs

IGB can consume data from more than 30 different file formats, and users can open and view data from local files or URLs. When users open a data set, a new track is added to the main display. The user then chooses how much will load into the new track by operating data loading controls. Larger data sets, such as RNA-Seq data, should be loaded on a region-by-region basis, while others that are small enough to fit into memory can be loaded in their entirety, e.g., a BED file containing ChIP-Seq peaks. To load data into a region, users zoom and pan to the region of interest and click a button labeled “Load Data.” To load all data in an opened data set, they can change the data sets loading method by selecting a “genome” load mode setting in the Data Access Panel.

This behavior differs from other genome browsers in that most tools link navigation and data loading. In other tools, a request to navigate to a new location or zoom level both redraws the display and also triggers a data loading operation. Although sometimes convenient...
for users, this limits the types of navigation interactions a tool can support. This trade-off can be seen in the Integrative Genomics Viewer (IGV), a desktop Java application developed after IGB (Thorvaldsdottir, et al., 2013). IGV auto-loads data but restricts movement; for example, it lacks panning scrollbars and does not support fast, animated zooming. IGB, by contrast, prioritizes navigation speed and gives users total control over when data load, which can be important when loading data from distant locations, which can sometimes feel slow. IGB gives users control over when delays may occur, thus making waiting more palatable.

3.4 Sharing and integrating data
IGB aims to support the scientific discovery process by making it easy for users to document and share results. Taking a cue from Web browsers, IGB for many years has supported bookmarking genomic scenes. In IGB, genomic scene bookmarks record the location, genome version, and data sets loaded into the current view. Users can also add free text notes and thus record conclusions about what they see. Selecting a bookmark causes IGB to zoom and pan to the bookmarked location and load all associated data sets, thus enabling users to quickly return to a region of interest. Users can sort, edit, import and export bookmarks using the Bookmarks tab. Exporting bookmarks creates an HTML file which users can re-import into IGB or open in a Web browser. If IGB is running, clicking an IGB bookmark in a Web browser causes IGB to zoom to the bookmarked location and load data sets associated with the bookmark.

IGB loads bookmarks through a ReST-style endpoint implemented within IGB itself, using a port on the user’s computer. This endpoint allows bookmarks to be loaded from HTML hyperlinks embedded in web pages, spreadsheets, or any other document type that supports hyperlinks. IGB was the first desktop genome browser to implement this technique; for many years, Affymetrix used it to display probe set alignments on their NetAffx Web site (Liu, et al., 2003).

More recently, we used IGB’s ReST-style bookmarking system to implement a Javascript bridge between IGB and the Galaxy bioinformatics workflow system (Goecks, et al., 2010). When users generate IGB-compatible data files within Galaxy, they can now click a ‘display in IGB’ hyperlink. Clicking this link opens a BioViz.org Web page containing a javascript program that forwards the hyperlink to IGB, causing IGB to open and display the data set. If IGB is not running, the javascript instead invites the user to launch IGB. Once they do, the data set loads.

IGB supports multiple formats and protocols for sharing data set collections and integrating across data sources. The most lightweight and easy to use of these is the IGB specific “Quickload” format, which consists of a simple directory structure containing plain text, metadata files. The metadata files can reference data sets stored in the same Quickload directory or in other locations, including Web, ftp sites, or cloud storage resources like iPlant or Dropbox (Goff, et al., 2011). This flexibility makes it possible for a Quickload site to aggregate data sets from multiple locations. The metadata for a Quickload site also includes styling and data access directives controlling the way IGB loads each file, and how the file will appear when loaded.

Sharing a Quickload site is straightforward. Users simply copy the contents to a publicly accessible location and then publicize the URL, which IGB users then add as a new Quickload site to their copy of IGB. Typically, users copy their Quickload sites to the content directories of Web sites. IGB supports secure access via the HTTP basic authentication protocol, making it easy to keep data sets private if desired. If users do not require password-protection for their data, they can also use Public Dropbox folders to share Quickload sites.

3.5 Visualizing RNA-seq data
Ultra high-throughput sequencing of cDNA (RNA-Seq) has largely replaced microarrays as a method for surveying gene expression, and so we added many new features to IGB to support visualization of RNA-Seq data, mostly focused on loading and manipulating BAM (binary alignment) and bedGraph/wiggle files.

RNA-Seq analysis data processing workflows typically produce large read alignment files (BAM) files, which IGB can open and display. However, viewing these “raw” read alignments is only the first step toward exploiting the depth and richness of these data sets. To help users better understand their data, IGB implements visual analytics functions developed to highlight and expose underlying patterns in RNA-Seq data. We describe a subset of these analytics functions below.

Right-clicking a BAM track label opens a context menu listing options to create coverage graphs, called “depth” graphs in IGB. At present, IGB supports two types of depth graphs: “depth graph start” that counts the first mapped nucleotide of a read and “depth graph all” that counts the number of reads overlapping each position. The latter is useful when investigating overall expression at a locus, while the former is useful when investigating sequencing bias. Both graph types are implemented using IGB’s track operations API, which developers can use to add all-new graph generation algorithms as plug-ins, discussed below.

By comparing depth graphs for multiple samples, users can identify differences in transcript levels. Figure 3a shows “depth graph all” graphs created from lung cancer samples bearing wild-type and mutant copies of the KRAS oncogene (Kalari, et al., 2012). Using the Graph tab, users can adjust the scale of both graphs; here both are scaled the same. Sequencing depth was similar in both samples, peaks that are the same height and shape in both samples reflect similar expression. However, one peak covering AQP3, encoding a water/glycerol-transporting protein (Hara-Chikuma and Verkman, 2005), is much taller in the mutant sample, indicating higher expression.

Visual analysis of RNA-seq data can also highlight alternative splicing differences between samples. Figure 3b shows PFN2, which produces two isoforms due to alternative splicing. Here, the depth graphs from Figure 2a were merged into a single track and again set to the same scale. Comparing the discontinuity in the peaks to the gene models shows that the mutant sample favors the shorter isoform.

To aid splicing analysis, we developed FindJunctions, a visual analytics tool that quantifies and displays read support for exon-exon junctions (i.e., introns) in RNA-Seq data. FindJunctions identifies split read alignments in an RNA-Seq track, uses them to identify
Fig. 3. Visualizing ChIP-seq data. (a) MACS BED file with peak regions from mouse ChIP-Seq data investigating binding sites for transcription factor SOX9 (Kadaja, et al., 2014). MACS peak regions are colored by score, with higher-scoring regions appearing red. (b) Zoomed in view of (a). MACS identify four significantly enriched peaks near Htral (bottom), but only two peaks exceed a user-defined coverage threshold, visible as a thin horizontal line, applied to the ChIP-Seq WIG track (top). (c) Zoomed in view of (b). Advanced residue search for the SOX9 binding motif AGCCGYG identified two instances, one in each of the two significant peaks on the positive strand.

3.6 Visualizing ChIP-Seq data

Knowing where a transcription factor binds DNA in relation to nearby genes is tantamount to understanding its function, as genes whose promoters are bound by a given transcription factor are likely to be regulated by it. Identifying binding sites of DNA-binding proteins is now routinely done using whole-genome ChIP-Seq, in which DNA cross-linked to protein is immunoprecipitated using antibodies against the protein of interest and then sequenced. Subsequent data analysis typically involves mapping reads onto a reference genome, identifying regions with large numbers of immunoprecipitated reads, and then performing statistical analysis to assess significance of enrichment. Each step requires users to choose analysis parameters whose effects and importance may be hard to predict. To validate these choices, it is important to view the “raw” alignments and statistical analysis results in a genome browser. As an example, we describe using IGB to view results from a ChIP-Seq analysis done using MACS, a widely used tool (Zhang, et al., 2008).

MACS produces a BED format file containing peak locations and significance scores indicating which peaks likely contain a binding site. Opening and loading this BED file in IGB creates a new track with single-span annotations representing the extent of each peak. At first, they look identical, varying only by length, and it is difficult to distinguish them. To highlight the highly scoring peak regions, IGB offers a powerful “Color by” visual analytics feature that can assign colors from a heatmap using quantitative variables associated with features. For this, users right-click a track, select “Color by” from the context menu, and then operate a heatmap editor adopted from the Cytoscape codebase to color-code by score (Shannon, et al., 2003). Doing this makes it easy to identify the highest scoring region most
likely to contain a binding site (Fig. 4a).

ChIP-Seq analysis tools also typically produce WIG-format depth-graph files that report where immunoprecipitated sequences have “piled up,” forming peaks. Loading this WIG file into IGB creates a new graph track that shows how these peaks coincide with regions from the BED file. In IGB, users can move tracks to new locations in the display. As shown in Figure 4b, placing the WIG track above the color-coded BED track makes it easy to observe how taller peaks typically have higher significance values.

IGB’s graph thresholding function, originally developed for tiling arrays, is useful for exploring the relationship between coverage depth and peak score. Available from the IGB Graph tab, the thresholding function identifies regions in a graph track where consecutive y-values exceed a user-defined threshold. Users can change the threshold dynamically using a slider and observe in real-time how changing the threshold value affects the number and extent of identified regions. Users can promote regions to new tracks and save them in BED format. As an example, Figure 4b shows four MACS-detected peaks near the H6601 gene, a regulatory target for the SOX9 transcription factor in mouse (Kadaja, et al., 2014). Two exceed the current threshold setting, providing a visual cue that these two peaks may be most important for regulation.

Sometimes the recognition sequence for transcription factor being studied is known or can be deduced from the data. IGB offers a way for users to visualize instances of binding site motifs. Using the Advanced Search tab, users can search the genomic region in view using regular expressions. For instance, to search for the SOX9 binding motif AGCCGTYG (where Y can be C or T) a user would enter “AGCCG[T]YG”. In this example, the search found several instances of this motif, with one located in each of the two user-defined significant peaks near the H6601 gene (Fig. 3c).

### 3.7 Visualizing bisulfite sequencing data

Whole genome bisulfite sequencing (WGBS) refers to bisulfite conversion of unmethylated cytosines to thymines followed by sequencing. This technique can identify methylated sites throughout a genome and expose epigenetic regulation of individual genes. Analysis of bisulfite involves mapping the reads onto the genome using tools that can accommodate reads where many but not all C residues have been converted. Several such tools are available; here we describe using IGB to visualize output of Bismark (Krueger and Andrews, 2011).

Bismark produces a BAM file containing read alignments and a depth-graph file (bedGraph format) reporting percent methylation calculated from sliding windows along the genome. Figure 5a shows a Bismark bedGraph file from an experiment investigating methylation in the model plant Arabidopsis thaliana. The entirety of chromosome one is shown, and peaks indicate regions of high methylation. This
3.8 Extending IGB using plug-ins

Developers have created dozens of genome browser tools, each one aiming to meet a need not met by the tools that preceded it. And yet, each new tool has faced similar problems, such as how to consume data from files, how to lay out genomic features and graphs into tracks, and how to support zooming through vast differences in scale. As described above, IGB has solved many of these problems and offers a flexible and fast environment for users to explore the genomic landscape. If it were easy to add new features and tools to IGB, they could spend more time creating new visualizations and less time re-implementing features that already exist. Toward this end, we transformed IGB into a modular, extensible platform for developers to create and deploy all-new visualizations of genomic data.

The IGB software architecture now resembles other popular open source java projects that support adding new functionality via plug-ins, including Eclipse, the Netbeans Rich Client Platform, and Cytoscape (Shannon, et al., 2003). This similarity comes from our common use of OSGi, a services based architectural framework and community standard for building modular software. By adopting this framework, we increased IGB extensibility, reduced the difficulty of adding new features, and created a simple plug-in API that empowers community developers to contribute new functionality without needing a deep understanding of IGB internal systems. This plug-in API is still new, but community developers are already using it to create novel visualizations using IGB as a platform (Céol and Müller, 2015). Documentation describing how to create plug-ins for IGB is available from the IGB developer’s guide.

3 Future directions

IGB offers users powerful utilities for viewing, analyzing, and interacting within an environment that feels fast, flexible, and highly interactive. In future, we plan to make the IGB’s Quickload data sharing system easier to use by providing tools for building Quickload sites from within IGB, along with a Quickload registry for users to publicize their sites. In addition, we will continue developing and improving the IGB plug-in APIs, providing documentation and example plug-ins to demonstrate how developers can use IGB as a platform to support genomics research.

Acknowledgements

We thank developers, testers, and designers who contributed to IGB, some of whom include Michael Lawrence, Lance Frohman, Gregg Helt, John Nicol, Hiral Vora, Alyssa Gulledge, Fuquan Wang, David Nix, Ido Tamir, Zhong Ren, Tarun Kanaparthi, Kyle Suttlemyre, and Tarun Mall.

Funding

This work received support the National Institutes of Health [R01GM103463 to AL].

References


