Title
The UCSC Genome Browser database: 2017 update.

Permalink
https://escholarship.org/uc/item/6582q1fk

Journal
Nucleic acids research, 45(D1)

ISSN
0305-1048

Authors
Tyner, Cath
Barber, Galt P
Casper, Jonathan
et al.

Publication Date
2017

DOI
10.1093/nar/gkw1134

Peer reviewed
The UCSC Genome Browser database: 2017 update

Cath Tyner1,*, Galt P. Barber1, Jonathan Casper1, Hiram Clawson1, Mark Diekhans1, Christopher Eisenhart1, Clayton M. Fischer1, David Gibson1, Jairo Navarro Gonzalez1, Luvina Guruvadoo1, Maximilian Haeussler1, Steve Heitner2, Angie S. Hinrichs1, Donna Karolchik1, Brian T. Lee1, Christopher M. Lee1, Parisa Nejad1, Brian J. Raney1, Kate R. Rosenbloom1, Matthew L. Speir1, Chris Villarreal1, John Vivian1, Ann S. Zweig1, David Haussler1,3, Robert M. Kuhn1 and W. James Kent1

1Genomics Institute, University of California Santa Cruz, Santa Cruz, CA 95064, USA, 2Emory University School of Medicine, Atlanta, GA 30322, USA and 3Howard Hughes Medical Institute, University of California Santa Cruz, CA 95064, USA

Received September 14, 2016; Revised October 17, 2016; Editorial Decision October 18, 2016; Accepted October 31, 2016

ABSTRACT
Since its 2001 debut, the University of California, Santa Cruz (UCSC) Genome Browser (http://genome.ucsc.edu/) team has provided continuous support to the international genomics and biomedical communities through a web-based, open source platform designed for the fast, scalable display of sequence alignments and annotations landscaped against a vast collection of quality reference genome assemblies. The browser’s publicly accessible databases are the backbone of a rich, integrated bioinformatics tool suite that includes a graphical interface for data queries and downloads, alignment programs, command-line utilities and more. This year’s highlights include newly designed home and gateway pages; a new ‘multi-region’ track display configuration for exon-only, gene-only and custom regions visualization; new genome browsers for three species (brown kiwi, crab-eating macaque, Malayan flying lemur); eight updated genome assemblies; extended support for new data types such as CRAM, RNA-seq expression data and long-range chromatin interaction pairs; and the unveiling of a new supported mirror site in Japan.

INTRODUCTION
With the revolutionary rate at which vast amounts of biological data are now being generated, new technologies and resources that can accommodate computational challenges are constantly in demand. The UCSC Genome Browser (1) continually strives to meet the needs of biological researchers in the face of exponential growth. This update gives a brief overview of the Genome Browser tools and data, and describes what has been added, changed, and improved since the previous 2016 update (2).

Genome assemblies
From its original focus on the early drafts of the human genome, the Genome Browser database now offers genomic data for nearly 100 organisms, many with multiple assemblies. Available as genome browsers and as downloadable data, this year there are three new species assemblies available (brown kiwi, crab-eating macaque, Malayan flying lemur) and new assembly versions for the following eight species: C. elegans, cat, chicken, gray mouse lemur, platypus, rhesus macaque monkey, western clawed frog, and western gorilla (3–8). For a detailed list of new and updated UCSC-hosted genomes, see Table 1.

Genome sequences and annotations for all UCSC-hosted assemblies can be downloaded from http://hgdownload.soe.ucsc.edu/. Downloadable annotations for UCSC-hosted assemblies generally include GenBank sequence alignments (9), repeat annotations using RepeatMasker (Smit et al., RepeatMasker Open-4.0 at http://repeatmasker.org) and Tandem Repeats Finder (10), soft and hard masked assembly sequences, chromosome sizes, and more.

Genome browser annotation data tracks
Assemblies for all species contain a set of basic annotations (‘tracks’), some of which are automatically generated by UCSC (e.g., assembly and gaps, the percentage of guanine and cytosine bases, repeat regions). Popular research species such as human and mouse are much more richly annotated, with tracks showcasing data from international sources.

The UCSC Genome Browser team regularly updates data tracks and adds new annotations. Although some annota-

*To whom correspondence should be addressed. Tel: +1 831 459 3597; Email: cath@ucsc.edu

© The Author(s) 2016. Published by Oxford University Press on behalf of Nucleic Acids Research. This is an Open Access article distributed under the terms of the Creative Commons Attribution License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted reuse, distribution, and reproduction in any medium, provided the original work is properly cited.
Table 1. New and updated UCSC-hosted genomes, 2015–2016

| Scientific name (common name)                                      | Assembly date | UCSC ID          | Sequencing center, NCBI accession ID, Assembly name |
|-------------------------------------------------------------------|---------------|------------------|-----------------------------------------------------|
| **New Species**                                                   |               |                  |                                                     |
| Apteryx australis mantelli (brown kiwi)                           | June 2015     | aptMan1          | Max-Plank Institute for Evolutionary Anthropology    |
|                                                                  |               |                  | GCF_001039765.1                                      |
|                                                                  |               |                  | AptMant0                                             |
| Macaca fascicularis (crab-eating macaque)                         | June 2013     | macFas5          | Washington University (WashU)                        |
|                                                                  |               |                  | GCA_000364345.1                                      |
|                                                                  |               |                  | Macaca_fascicularis_5.0                              |
| Galeopterus variegatus (malayan flying lemur)                      | June 2014     | galVar1          | Washington University (WashU)                        |
|                                                                  |               |                  | GCF_000696425.1                                      |
|                                                                  |               |                  | G_variegatus-3.0.2                                   |
| **Updated Species**                                               |               |                  |                                                     |
| Felis catus (domestic cat)                                        | November 2014 | felCat8          | International Cat Genome Sequencing Consortium       |
|                                                                  |               |                  | GCA_000181335.3                                      |
|                                                                  |               |                  | Felis_catus_8.0                                      |
| Caenorhabditis elegans (C. elegans)                               | February 2013 | ce11             | C. elegans Sequencing Consortium WBcel235           |
|                                                                  |               |                  | GCA_000002985.3                                      |
|                                                                  |               |                  | WBcel235                                             |
| Xenopus tropicalis (tropical clawed frog)                         | September 2012| xenTro7          | US DOE Joint Genome Institute (JGI-PGF)              |
|                                                                  |               |                  | GCA_000004195.2                                      |
|                                                                  |               |                  | Xtropicalis_v7                                       |
| Gorilla gorilla (western gorilla)                                 | December 2014 | gorGor4          | Wellcome Trust Sanger Institute                      |
|                                                                  |               |                  | GCA_000151905.3                                      |
|                                                                  |               |                  | gorGor4                                              |
| Microcebus murinus (gray mouse lemur)                             | May 2015      | micMur2          | Broad Institute and Baylor College of Medicine       |
|                                                                  |               |                  | GCA_000165445.2                                      |
|                                                                  |               |                  | Mmur_2.0                                             |
| Ornithorhyncus anatinus (platypus)                                | February 2007 | ornAna2          | Washington University Genome Sequencing Center       |
|                                                                  |               |                  | GCA_000002275.2                                      |
|                                                                  |               |                  | Ornithorhyncus_anatinus-5.0.1                        |
| Macaca mulatta (rhesus monkey)                                   | November 2015 | rheMac8          | Baylor College of Medicine Genome Sequencing Center   |
|                                                                  |               |                  | GCF_00072875.2                                       |
|                                                                  |               |                  | Mmul_8.0.1                                           |
| Gallus gallus (chicken)                                           | November 2011 | galGal5          | Chicken Genome Consortium (ICGC)                     |
|                                                                  |               |                  | GCA_000002315.2                                      |
|                                                                  |               |                  | Gallus_gallus-5.0                                     |

**Mapping and sequencing.** Tracks in this group annotate the foundational structure of the assembly and constructed contigs. Beyond automatically updated tracks, most of the prominent new tracks in the Mapping and Sequencing category this year were added to the human GRCh38/hg38 assembly. These include a new fluorescent in situ hybridization (FISH) clones track (14) lifted from NCBI36/hg18, and a new track that annotates STS markers (15,16). A Clone Ends track was added showing mapped clone end libraries for mouse (GRCh38/mm10) and rat (RGSC_6.0/rn6) from NCBI's Clone DB (17).

**Genes and gene predictions.** This group includes computationally predicted gene sets (e.g., Ensembl (13), Augustus), high-quality manually curated and evidence-based automated gene predictions for the human and mouse genomes from the GENCODE project (18), and subsets from NCBI such as manually curated sets from the Reference Sequence collection (RefSeq) (19). This year there were several major gene set additions and updates, primarily to the latest human and mouse assemblies. Existing Ensembl gene annotations were updated to version 81, and new Ensembl tracks were added to several assemblies: *Drosophila* (dm6), rat (rn6) and zebrafish (danRer10). Two major gene sets (GENCODE Genes, UCSC Genes) were updated, as described below.

**GENCODE genes.** The GENCODE Genes set is now the default gene set for the latest GRCh38/hg38 human assembly, replacing the previous default set, UCSC Genes. This transition was made to reduce the number of competing gene transcript sets used by the bioinformatics community, which have caused confusion in downstream analyses due to subtle differences among the sets. As of September 2016, the browser displays the v24 version, released by GENCODE in August 2015. The v24 gene set has increased by 2604 new transcripts to a total of 197,782 total transcripts. Only the Basic subset is displayed in the Genome Browser by default; the underlying database tables contain the Comprehensive
set, which can be viewed by clicking the ‘show comprehensive set’ checkbox on the track controls page. The database schema retains most of the same table names (starting with ‘known’ or ‘kg’). Every transcript has a UCSC identifier (e.g., uc001abz.5) in addition to the GENCODE identifier (e.g., ENST00000327044.6).

**UCSC genes.** The UCSC Genes set (20) has been updated for the mm10 mouse assembly. The new release increased by 515 new transcripts to a total of 63,759 total transcripts, 95% of which remained the same between versions. The total number of canonical genes has increased by 121 to a total of 33,079.

**Phenotype and literature.** This group contains tracks linking to phenotype references and literature (e.g., published articles) in which referenced sequences or identifiers are mapped to the assembly. The GRCh37/hg19 human assembly is currently the most comprehensively annotated with 21 major tracks, displaying data related to genetic phenotypes such as Online Mendelian Inheritance in Man (OMIM) allelic variants and positions of OMIM genes (21) as well as variants and copy number variants from the National Institutes of Health (NIH) funded ClinVar (22) and ClinGen (23) datasets. One highlight in this category is the addition of a super-track showing the genomic mapping of biomedic sequences submitted as part of patent application documents worldwide, obtained from the Cambia Lens Patent database (24) and aligned to the human (hg19), mouse (mm10) and Ebola virus (eboVir3) genome assemblies (25).

Other new track additions for the recent human assemblies include new copy number variant tracks for CNV Developmental Delay and ClinGen Benign Aggregate, both for hg38 and hg19. In hg38’s new non-coding RNA track (lifted from hg19), four different types of RNA data are positioned in three different sub-tracks: sno/miRNAs, tRNAs (also added as a new track in hg19), and lincRNAs (26-29).

As of 2016, proprietary OMIM data can now be accessed on a per-chromosome basis from the Table Browser and the Data Integrator; whole genome downloads are not available, per agreement with OMIM.

**Expression.** This category contains tracks related to the synthesis of functional gene products. A new gene expression track based on data from the NIH Genotype-Tissue Expression (GTEx) project (30,31) was released for the most recent human assemblies, hg38 and hg19. The GTEx track displays tissue-specific gene expression based on RNA-seq in 53 tissues from 570 donors obtained from the GTEx midpoint data release (V6, October 2015). This browser track uses a new bar graph display with a graph shown for each gene, and a colored bar indicating each tissue assayed (Figure 1). The ‘squish’ view colors the gene by most highly expressed tissue. A box plot showing the range of expression levels across samples is displayed on the details page for each gene in the GTEx track, and also on the details page for that gene on the browser’s default gene set.

Additional GTEx expression data are available in two UCSC-hosted track hubs. The first is the GTEx RNA-Seq Signal hub, hosting ~7000 tracks, one for each tissue sample in the midpoint data release. The second is the GTEx Analysis hub, which currently hosts summary tracks of allele-specific expression (per-tissue and cross-tissue). GTEx gene expression has also been incorporated in the Gene Sorter tool, replacing GNF Atlas2 as the default gene expression set.

**Regulation.** This group contains tracks related to regulation of gene expression, including a wide range of mechanisms used to increase or decrease the production of gene products. The most notable tracks in this category are found primarily on the hg19 assembly, representing data from the ENCyclopedia Of DNA Elements (ENCODE) project (32). The UCSC Genome Browser offers ENCODE data through 2012 (33). The entire collection of ENCODE results from 2007 until present day are available from the ENCODE Project Portal (http://encodeproject.org) (34) hosted by Stanford University. The major integrated ENCODE tracks have been mapped to hg38; however, the bulk of annotations remain on hg19.

**Comparative genomics.** Conservation tracks combine multi-species genome-wide alignments and estimated evolutionary conservation and acceleration scores to enable visualization and analysis of evolutionary processes. In September 2015, a new 100-vertebrate conservation track was released on the hg38 human assembly that compares pairwise alignments for 100 species from a variety of clades. Another addition is the C. elegans (ce11) 26-species conservation track, which shows the multiple alignments and measurements of evolutionary conservation for 26 nematode species.

**Variation.** Major variant annotation tracks include small variations from NCBI’s dbSNP database (35), including tracks for older dbSNP releases in human assemblies, along with variant calls and high-confidence regions from the 1000 Genomes Consortium (36).

**New dbSNP Build 146 and Build 147.** Data from NCBI’s dbSNP builds 146 and 147 have been added to the two most recent human assemblies (hg38, hg19), and Build 146 annotations were also added to the mm10 mouse assembly. Both builds 146 and 147 contain >150 million variants for human.

**New phase 3 data of the 1000 Genomes Project.** Data from Phase 3 of the 1000 Genomes Project (representing 2504 individuals from 26 populations worldwide) are now available for the hg19 human assembly. These data include nearly 90 million single nucleotide variants (SNVs), insertions/deletions (indels), and structural variants (SVs). For each variant, a detail page incorporates links to the variant in the dbSNP database, quality scores and allele frequency information for various populations. In addition to variant calls, the ‘Paired-end Accessible Regions’ tracks encompass genomic regions estimated to be accessible to next generation sequencing technologies using short paired-end reads, based on depth and quality of mapped reads.
Figure 1. GTEx tracks in the UCSC Genome Browser. This image shows a 31 kb region of chromosome 9 where GENCODE annotates four protein-coding genes showing different patterns of tissue-specific gene expression in GTEx tissue samples. The GTEx Gene Expression track displays a bar graph for each gene, where bar color indicates the tissue assayed and bar height depicts the median expression level across all samples in reads per kilobase of transcript per million mapped reads (RPKM). Tissues are identified on mouseover and the full legend is shown on the track description and details pages. The GTEx transcript model used to quantify expression level is displayed below the graph, colored by gene class using GENCODE conventions. The GTEx RNA-seq tracks show per sample read density; in this image, eight tissues from a single GTEx donor. The GTEx Analysis Hub is also connected and therefore is visible in the track menu, but no tracks are selected for display. Tissue colors in all GTEx tracks are assigned using GTEx project conventions.

Home and gateway pages

While the home page of the UCSC Genome Browser is the front door to the online browser, the gateway page is the entrance to visualizing all genomes hosted by the browser, providing opportunities to access 96 unique species (all eukaryotic, except for one virus), whose assembly versions combine to a total of 175.

Both the home page and the gateway page have been redesigned this year to be more interactive and intuitive. The new home page (Supplementary Figure S1) includes more graphics, less text, and better access to Genome Browser tools and content, such as the ability to access popular organisms from the site-wide menu. The newly designed gateway page (Supplementary Figure S2) features an autocomplete search for both UCSC-hosted and public assembly hubs, an interactive phylogenetic tree menu for selecting species, quick access shortcuts to popular browsers, and a new style and color scheme. From both the gateway page and the genome browsers, the ability to search genomes by submitting a chromosomal position, gene symbol, or search term remains a much-used feature. For the human genome, the hg38 assembly is now the default; the previous default assembly, hg19, can be quickly selected from the Genomes menu in the site-wide top menu bar.

Visualizing genomes: genome browser tracks display

The Genomes Browser tracks display can be seen as the ‘heart’ of the browser. The selected reference genome assembly provides navigational structure for browsing sequence alignments and annotations. The fast, responsive interface shows details from the nucleotide base level out to whole-chromosome perspectives. Major features of the browser display include a powerful search for all data tracks; track configuration and reordering; reverse strand view; and a variety of navigation options to zoom, pan, or quickly jump between features.

New multi-region display feature. The multi-region display configuration was developed to omit introns or intergenic
regions from the display, resulting in a Genome Browser display with exon-only or gene-only views (Figure 2), much like a handheld paper fan where portions of the design are hidden behind creased folds. This custom viewing mode can be launched from the browser multi-region button, from the ‘View’ menu on the navigation bar, or (in the case of the exon-only view) by using the keyboard shortcut ‘e v’ while on the graphic browser window.

From the multi-region configuration window, clicking the checkbox to turn on alternating region highlights will help differentiate alternating regions; every other region will appear with a tinted hue, thus creating a series of vertical stripes in the graphic browser window as the genome is ‘sliced’ to display only the defined regions.

The multi-region configuration window offers several display choices: exon-only, gene-only, alternate haplotype view, or custom BED regions. For the exon-only and gene-only modes, the browser automatically chooses which gene track to use as the source for the region coordinates, and the user may configure the padding between features. Available on human assemblies hg17 and newer is the option to display an alternate haplotype within the context of the reference genome (Supplementary Figure S3). A particularly powerful feature of the multi-region configuration is the option to specify the URL of a remotely hosted BED (Browser Extensible Data) file containing coordinates that define custom genomic regions, even those that span chromosomes.

New density graph feature. Another new visualization option for tracks is a density graph track display, useful for viewing tracks with many items in a large region (e.g., short-read sequence alignments from a BAM file). Selecting this option from the track settings will replace the visualization of dense data tracks with bar charts that show data coverage. This graph can be seen in tracks such as gene prediction tracks (e.g., RefSeq Genes and UCSC Genes), alignment tracks (e.g., GenBank mRNAs), and cross-species chained alignment tracks. The density coverage graph can also be configured for visibility in custom tracks, track hubs and assembly hubs with supported formats.

New keyboard shortcuts. From the Genome Browser data tracks webpage, pressing the question mark “?” key will open a menu of keyboard shortcuts, which has been expanded to include several new options (e.g., toggling zoom views, navigating upstream or downstream in a chromosome, hiding all tracks and launching the ‘view DNA sequence in external tools’ feature).

New browser menu options. There have been three major changes to the browser’s site-wide top menu bar. First, the ‘Genomes’ menu option now allows quick access to the latest two human and mouse assemblies. Secondly, the ‘View > In External Tools’ menu provides an option to export the current browser region to external tools (e.g., Ensembl, NCBIMapView (37), CRISPOR (38)). Lastly, the ‘Genome Browser’ menu now features a ‘Configure’ link to set visibilities for all tracks on the currently browsed assembly. The ‘Reset All User Settings’ option in the same menu will immediately restore the UCSC Genome Browser on the current Internet browser to a default state by clearing connections to external hubs, removing loaded custom tracks, and reverting to site-wide default configurations (such as default track views for the hg38 assembly).

New features in gene description page. In the Genome Browser tracks display, clicking on a gene from a browser’s default gene set will display a gene description page containing links to other tools and databases. From this description page, a chart showing RNA-Seq Expression Data from GTEx (53 Tissues, 570 Donors) is now displayed on the two latest human assemblies. Other changes for the gene description page include updated data and links to the Malacards human disease database (39) and updated Wikipedia links.

New sessions gallery and public sessions
One widely used feature of the browser is the ‘Sessions’ tool, which allows the user to save the configuration settings of the current browser view for later viewing and sharing (40). Sessions are now arranged in a table with filtering and ordering options based on session name, primary assembly, and creation date. From the session list, clicking on the session name displays that session in the browser. Session expirations have been eliminated to assist with long-term data sharing.

A new Session Gallery has been added, featuring sample sessions created by the UCSC Genome Browser team, which highlight topics of interest to the genomics community. To date, sessions in the gallery include the display of coding and wobble bases, alt-splicing, evolution, variation and disease, topics derived from commonly asked questions on the browser user support mailing list, and sessions used in browser training workshops.

A new ‘Public Sessions’ page has been released, displaying Genome Browser sessions that users have opted to share to this publicly accessed list. In the Session tool’s listing of saved sessions for a user’s account, a checkbox can be marked to enable public viewing permissions, thus adding the session to the public list. From the Public Sessions page, sessions can be filtered by assembly, name, popularity, creation date, or a phrase from the description. Public Sessions can be reached from the site-wide menu (My Data > Public Sessions).

Table-based browsing and querying
There are several major interactive web-based tools purpose for browsing and querying underlying tables in the UCSC Genome Browser, including most track data.

First, and most widely used, the web-based Table Browser tool (41) can be accessed directly from the Genome Browser menu, pre-loading current browser configurations for the querying, intersecting, filtering and downloading of genomic data. Another tool is the web-based Data Integrator (VAI) (42), specializing in combining and exporting data from multiple tracks simultaneously. The Variant Annotation Integrator (VAI) (42) annotates user-provided variant calls with predicted functional effects and data from other tracks.

Related database tables in the data integrator. Several UCSC-hosted tracks use multiple related tables in the SQL
Figure 2. Multi-Region display in the UCSC Genome Browser. This figure displays the MYPN gene in the UCSC Genome Browser using the new multi-region display mode with the ‘exon-only’ option and ‘highlight alternating regions’ enabled. Exonic regions are calculated using the GENCODE v24 Comprehensive Transcript Set on the GRCh38/hg38 human genome assembly and are highlighted in alternating white and blue backgrounds. Intergenic and intronic regions that do not overlap with exons in other transcripts are removed from the display.

database to store track data. For many years, the Table Browser has allowed related tables to be joined with the main track table so that their fields can be included in the output. For example, protein product accessions and descriptions associated with RefSeq Genes can be joined by selecting the protAcc and product fields from the refLink table. The Data Integrator now offers the same functionality via the ‘Choose fields...’ button and dialog box.

Updated dbNSFP data in hgVai. For hg38, the VAI now offers selected annotations from version 3.1a of the Database of Non-Synonymous Functional Predictions (dbNSFP), which provides pre-computed scores and predictions of functional significance from a variety of tools.

Custom tracks, track hubs, assembly hubs and public hubs

The UCSC Genome Browser offers three primary options for viewing custom genomic data in the form of custom tracks, track hubs and assembly hubs. These tools provide methods for publicly sharing or privately viewing custom genome assemblies, large-scale genomic annotations, and data tracks.

The collection of data types supported by the custom track, track hub, and assembly hub tools continues to expand. New supported data types include bigPsl, bigChain and bigMaf (which store alignments, pairwise alignments, and multiple alignments, respectively), and CRAM. CRAM is similar to BAM, but denser, using an external reference sequence index. Another new UCSC-specific file type, longTabix, is now supported as a custom track or track hub. This format stores chromatin interactions of paired chromosomal regions. Regional pairs are visualized within the browser with an ‘arc’ representation, similar to chromatin interactions displayed in the WashU EpiGenome Browser (43). For a complete listing of data types supported by track hubs and custom tracks, see Supplementary Table S2.

Custom tracks. The simplest way to view custom data against a UCSC-hosted genome assembly is by uploading a file to the browser as a custom track in one of many supported file formats, some of which are new as of this last year (Supplementary Table S2). For most big* compressed formats (e.g., bigWig), the URL to the hosted files alone can be pasted into the custom tracks data-loading dialog box, without additional track information needed.

As with other major tools such as the Table Browser, custom tracks are now integrated with the Data Integrator tool. The new option to view dense data as a bar graph is available for supported custom track data types.

Track hubs. Track hubs (44) are web-accessible directories of genomic data that can be viewed as a data portal through the UCSC Browser. The new option to view dense data as a bar graph is available for supported data types.

Public UCSC-reviewed track and assembly hubs. The UCSC Genome Browser team continues to promote the use of public track and assembly hubs to display large data sets
from consortia and external labs. As of September 2016, there are over 45 public hubs linked for display in the UCSC Genome Browser. Of these, thirteen were added this last year. For a complete listing of recently added public hubs, see Table 2.

### Alternative UCSC genome browsers: GBiB, mirrors

The UCSC Genome Browser is available as a stand-alone virtual machine, the Genome Browser in a Box (GBiB) (45), that can be used on a personal computer. GBiB provides a complete solution for browsing confidential data, while requiring a small memory footprint. Alternatively, the Genome Browser can be fully or partially mirrored at a third-party site, thus allowing customizations of any browser component or data. The ‘My Data’ menu now includes a new GBiB Shared Data Folder, providing navigation to locally shared files that can be entered as URLs in the custom tracks or track hubs tools. The GBiB custom track tool provides local file-loading capabilities by pointing to the ‘folders’ directory with the bigDataUrl parameter.

A new supported Genome Browser mirror (genom asia.ucsc.edu) has been released to provide an access point of faster functionality for those users who are geographically closer to Asia than to the United States. This UCSC-administered mirror site is physically located at the RIKEN Yokohama Campus in Japan. Technical support continues to be provided for full and partial mirrors of the Genome Browser offered by academic and noncommercial institutions. Mirror sites can now specify the location of a Galaxy (46) instance to the Table Browser.

### Public MySQL server access

As an alternative to the graphical Table Browser tool, the UCSC-hosted MySQL database is publicly accessible and synchronized on a weekly basis from the main databases on the UCSC Genome Browser website. Many of the command-line utilities from the source tree can be used with the public MySQL databases. These data are stored on a central file server and MySQL server, totaling approximately 20 terabytes of disk space, and growing daily.

This past year, additional security was implemented in the form of SSL (Secure Sockets Layer) connectivity between the browser hosts and MySQL servers. Also, the location of related tables that support the GenBank and RefSeq tracks (e.g., RefSeq, Other RefSeq, mRNA, EST) have moved from individual assembly databases to one shared metadata database, hgFixed.

### Utilities and source code

The UCSC Genome Browser team provides ~250 application binaries built for standalone command-line use on Linux and Mac OSX platforms. There are several new additions to the command-line utility collection, including file conversion tools (pslToBigPsl, bigPslToPsl, chainToPslBasic) and utilities for loading specific file types into a database (hgLoadChain, hgLoadMaf, hgLoadNet, hgLoadMafSummary).

---

**Table 2.** New track and assembly public hubs hosted by UCSC, 2015–2016

| Hub name & assemblies | Source |
|-----------------------|--------|
| GTEX RNA-Seq Signal Hub: RNA-seq read coverage in 53 tissues from GTEX V6 (7572 samples). *Assemblies*: hg38, hg19 | GTEX Consortium, UCSC Computational Genomics Lab, UCSC Genome Browser group |
| GTEX Analysis Hub: Allele-Specific Expression in 53 Tissues. *Assemblies*: hg38, hg19 | GTEX Analysis Working Group (Lappalainen lab, NY Genome Center) |
| Peptide evidences CNIO: Peptides detected from a re-analysis of multiple experiments and databases. *Assemblies*: hg38, hg19 | Spanish National Cancer Research Centre (CNIO), Spanish Institute of Bioinformatics |
| Principal Splice Isoforms APPRIS: Selects a single CDS variant for each gene as the ‘PRINCIPAL’ isoform based on the range of protein features. *Assemblies*: hg38, hg19, mm10, danRer10, rn6, susScr3, panTro4, d0117, ce10 | Spanish National Cancer Research Centre (CNIO), Spanish Institute of Bioinformatics |
| DASHR small ncRNA: DASHR Human non-coding RNA annotation. *Assemblies*: hg19 | Li-San Wang Lab, University of Pennsylvania |
| Cancer Genomics Tracks: TCGA and ICGC Cancer Mutations, TCGA Expression, Immune Epitopes Database (IEDB), Cancer Immunity Peptides Database, Dienstmann Variant/Cancer database, CIVIC and MyCancerGenome.org. *Assemblies*: hg19 | UCSC Genome Browser group |
| lncRNA in Breast Cancer: long non-coding RNAs in Breast Cancer. *Assemblies*: hg19 | Kraus Lab at UT Southwestern |
| mm9.SMC1.ChIAPET: Cohesin(Smc1)-associated chromatin interactions in murine embryonic stem cells. *Assemblies*: mm9 | Young Lab, Whitehead Institute, MIT |
| Promoterome CAGE and nucleosome positioning: Zebrafish promoterome based on Haberle et al. and Nepal et al. *Assemblies*: danRer7 | Computational Regulatory Genomics Group, MRC Clinical Sciences Centre |
| Peterhof Yeasts: Assemblies, SNV and CNV data for Saccharomyces cerevisiae strains of the Peterhof Genetic Collection. *Assemblies*: sacCer3 & strains | Saint Petersburg State University |
| CESAR Gene Mappings to 99 vertebrates: Human Exons mapped by CESAR. *Assemblies*: hg38, hg19 > many | Max Planck Institute of Molecular Cell Biology and Genetics |
| ChIP-seq data track HUBs fromMSC cells from GSE79815. *Assemblies*: mm9 | NCBI: Gene Expression Omnibus |
| UniProt Features: UniProtKB reviewed protein features mapped to the Ensembl genome assembly. *Assemblies*: hg38 | The UniProt Consortium |
The entire source code tree, which is written primarily in C, HTML, and Javascript, is publicly available. To install the package, consult the detailed README files in the source tree (http://genome-source.cse.ucsc.edu/gitweb/).

Both the binaries and source code are available for download. Licenses are required for commercial use (https://genome-store.ucsc.edu/), with several exceptions (see http://genome.ucsc.edu/license/).

Support, training and documentation
This last year, in-person trainings, workshops and presentations were conducted in the United States and international locations, including Germany, Brazil, South Korea, Japan, Sweden, Spain and Denmark. More information regarding workshop requests can be found online (http://bit.ly/ucscTraining).

Multi-media learning materials continue to be developed, such as user guides (e.g., the new multi-region display), online training videos (e.g., how to view exons only and export exon coordinates), announcements of major new data releases, and new blog posts.

FUTURE PLANS
The UCSC Genome Browser team will continue to release new genome assemblies for species not yet hosted at UCSC, for updates for existing genomes, new and updated annotations, new software and overall site improvements. In the coming year, the new Genome Browser in the Cloud (GBiC) will be released. The GBiC is a cloud-based solution that is similar to the stand-alone virtual machine, GBiB.

The Genome Browser team has been working to provide an NCBI RefSeq Gene composite track that will provide alignments and annotations directly from NCBI in addition to providing UCSC’s BLAT (47) alignments of RefSeq sequences and resulting transcript models.

Other planned software changes include new features for the multi-region display, improved track sorting, improved track hub searching, extensions to the GTEX track display, a publications track for the human assembly (hg38), and a CRISPR/cas9 track.

CONTACT US
All UCSC Genome Browser contact information can be found on the website at http://genome.ucsc.edu/contacts.html.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENTS
Sincere gratitude is extended to the worldwide users of the UCSC Genome Browser, from college students who are just beginning their journey to experienced researchers. Special appreciation is extended to funders, collaborators, and contributors who give value beyond measure to the research possibilities within the browser. Much thanks is given to the exceptional group of system administrators, the management team, and all staff who support the UCSC Genome Browser.

FUNDING
National Human Genome Research Institute [5U41HG002371 to C.T., G.P.B., J.C., H.C., M.D., C.E., C.M.L., C.M.F., D.G., J.N.G., L.G., M.H., S.H., A.S.H., D.K., B.T.L., P.N., B.J.R., K.R.R., M.L.S., C.V., A.S.Z., D.H., R.M.K., W.J.K.; 5U41HG006992 to W.J.K., B.T.L., B.J.R.; 5U41HG007234 to M.D.; 5U54HG007990 to B.P., J.V.]; National Cancer Institute [5U54HG007990 to M.D., H.M., R.M.K.; 5U24CA180951 to M.D.; 5417071–550000716 to J.V.]; California Institute for Regenerative Medicine [GC1R-06673-C to G.P.B., C.E., C.M.F., M.H., S.H., D.K., W.J.K., B.T.L., K.R.R., A.S.Z.]; Howard Hughes Medical Institute [090100 to D.H.]. Funding for open access charge: National Human Genome Research Institute [5U41HG002371].

Conflict of interest statement. G.P.B., J.C., H.C., M.D., L.G., M.H., S.H., A.S.H., D.K., B.T.L., B.J.R., K.R.R., M.L.S., A.S.Z. D.H., R.M.K. and W.J.K receive royalties from the sale of UCSC Genome Browser source code, LiftOver, and GBiB licenses to commercial entities. W.J.K. owns Kent Informatics.

REFERENCES
1. Kent, W.J., Sugnet, C.W., Furey, T.S., Roskin, K.M., Pringle, T.H., Zahler, A.M. and Haussler, D. (2002) The human genome browser at UCSC. Genome Res., 12, 996–1006.
2. Speir, M.L., Zweig, A.S., Rosenbloom, K.R., Raney, B.J., Paten, B., Nejad, P., Lee, B.T., Learned, K., Karolchik, D., Hinrichs, A.S. et al. (2016) The UCSC genome browser database: 2016 update. Nucleic Acids Res., 44, D717–D725.
3. Zimin, A.V., Cornish, A.S., Maudhoo, M.D., Gibbs, R.M., Zhang, X., Pandey, S., Meehan, D.T., Wippler, K., Bosinger, S.E., Johnson, Z.P. et al. (2014) A new rhesus macaque assembly and annotation for next-generation sequencing analyses. Biol. Direct, 9, 20.
4. Scally, A., Dutheil, J.Y., Hillier, L.W., Jordan, G.E., Goodhead, I., Herrero, J., Hobolt, A., Lappalainen, T., Mailund, T., Marques-Bonet, T. et al. (2012) Insights into hominid evolution from the gorilla genome sequence. Nature, 483, 169–175.
5. Pontius, J.U., Mullikin, J.C., Smith, D.R., Lindblad-Toh, K., Gnerre, S., Clamp, M., Chang, J., Stephens, R., Neeham, A., Agencourt Sequencing Team et al. (2007) Initial sequence and comparative analysis of the cat genome. Nature, 17, 1675–1689.
6. Hellsten, U., Harland, R.M., Gilchrist, M.J., Hendrix, D., Jurka, J., Kapitonov, V., Ovcharenko, I., Putnam, N.H., Shu, S., Tahe, L. et al. (2010) The genome of the western clawed frog Xenopus tropicalis. Science, 328, 633–636.
7. Warren, W.C., Hillier, L.W., Marshall, Graves, L.A., Birney, E., Ponting, C.P., Grützner, F., Belov, K., Miller, W., Clarke, L., Chinwalla, A.T. et al. (2008) Genome analysis of the platypus reveals unique signatures of evolution. Nature, 453, 175–183.
8. Howe, K.L., Bolt, B.J., Cain, S., Chan, J., Chen, W.J., Davis, P., Done, J., Down, T., Gao, S., Grove, C. et al. (2016) WormBase 2016: expanding to enable helminth genomic research. Nucleic Acids Res., 44, D774–D780.
9. Clark, K., Karchs-Mizrachi, I., Lipman, D.J., Ostell, J. and Sayers, E.W. (2016) GenBank. Nucleic Acids Res., 44, D67–D72.
10. Benson, G. (1999) Tandem repeats finder: a program to analyze DNA sequences. Nucleic Acids Res., 27, 573–580.
11. Stanke, M., Diekhans, M., Baertsch, R. and Haussler, D. (2008) Using native and syntenically mapped cDNA alignments to improve de novo gene finding. Bioinformatics, 24, 637–644.
12. Stanke, M. (2004) Gene prediction with a hidden Markov model.
