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GANESH: Software for Customized Annotation of Genome Regions

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GANESH is a software package designed to support the genetic analysis of regions of human and other genomes. It provides a set of components that may be assembled to construct a self-updating database of DNA sequence, mapping data, and annotations of possible genome features. Once one or more remote sources of data for the target region have been identified, all sequences for that region are downloaded, assimilated, and subjected to a (configurable) set of standard database-searching and genome-analysis packages. The results are stored in compressed form in a relational database, and are updated automatically on a regular schedule so that they are always immediately available in their most up-to-date versions. A Java front-end, executed as a stand alone application or web applet, provides a graphical interface for navigating the database and for viewing the annotations. There are facilities for importing and exporting data in the format of the Distributed Annotation System (DAS), enabling a GANESH database to be used as a component of a DAS configuration. The system has been used to construct databases for about a dozen regions of human chromosomes and for three regions of mouse chromosomes.

One of the underpinning aims of the Human Genome Project is to provide the resources to support genetic analysis of human conditions and disorders. This aim is incomplete in part because neither the finished DNA sequence of the whole of the human genome has yet been established, nor has the task of identifying all of the genes within even the available DNA sequence been completed (Lander et al. 2001). This is an inevitable consequence of the continuing nature of the HGP. But even if the genome were complete, software tools in the public domain are not optimally configured for gene discovery efforts in circumscribed genomic regions.

Presently, there are several partially independent sources of annotated human genomic sequences that include Ensembl (Hubbard et al. 2002), UCSC (Kent et al. 2002), and Celera (Venter et al. 2001). Each of these aims at producing an accurate assembly of the genomic sequence and using available databases of expressed sequences, combined with *in silico* gene prediction, to identify the location and structure of human genes with the greatest possible accuracy. Necessarily, these databases increase their coverage incrementally as both the finished DNA sequence and further computational resources become available, but, in turn, this means that the annotation processes lag behind relevant data. Explicitly, these databases are databases of record, and as a consequence, they tend to sacrifice completeness for the sake of accuracy and certainty. Evidence for the existence or location of a gene may be rejected if the appropriate level of confidence is lacking. Furthermore, we note that Hogenesch et al. (2001) have argued that the concordance between Ensembl and Celera's data is partial, suggesting that neither database can be considered as definitive.

In contrast to the requirements of databases of record, genetic research places rather different constraints upon the anno-

tation and analysis of genomic data. Modern genetic research generally takes a positional cloning approach, frequently using whole-genome scans for linkage in appropriately constructed pedigree collections to identify regions likely to contain a variant gene(s) that predisposes to the condition or disorder under study. Such approaches generally identify regions ranging from a few megabases (in the case of monogenic disorders) to tens of megabases (in the case of multifactorial traits). The challenge for the genetic researcher is then to identify the disease-susceptibility variant or variants within this region.

Genetic analysis software and gene identification software as represented by databases of record might, in time, become congruent. At present, this is not the case. There are three main distinguishing features. First, although gene identification is a key element of both types of software, the ultimate goal of genetic analysis software is to derive an exhaustive list of genes and gene-like objects within a specified region, so that these can be subjected to experimental analysis to identify sequence variants that might be correlated with disorder or condition state. Accuracy of gene prediction, especially the elimination of false positives, is of lesser importance because experimental analyses can be deployed readily to validate the *in silico* predictions (Shoemaker et al. 2001). Secondly, genetic analysis requires as exhaustive an identification of potential genes in the target region as possible, and so must be able to exploit any additional data sources that may be relevant to the target region. In contrast, for databases of record, the sheer scope of the target region necessarily limits the sources of data that can be accommodated. Thirdly, genetic analysis requires comparative analysis of all of the genes within the target region, to allow candidate prioritization. This is not a necessary feature of databases of record, which require classification tools that have a global scope. It has been said that the annotations served by the reference databases of record are broad, but shallow, whereas those required by genetic researchers tend to be narrow, but deep (Stein 2001).

These three design differences make the databases of record cumbersome and restricted in their utility for specifically genetic analysis. To overcome these limitations, we have developed a

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specialist software package (named GANESH), designed in explicit recognition of the differing goals and requirements of geneticist and genomicist. GANESH is a set of software components that may be assembled to construct a new self-updating database providing annotation for a specified region of human (or other) genomic sequence. Sequence and other relevant data for the target region are gathered from various distributed data sources, assimilated, and subjected to a range of database-searching and genome-analysis programs. The results are stored in the database in compressed form and updated on a regular schedule, so that they are always available immediately in their most up-to-date form. A front-end in Java, executed as a stand-alone application or as a web applet, provides a graphical interface for navigating the database and visualization of the genome features detected. There are utilities for importing and exporting data in a variety of formats, including those of DAS, the Distributed Annotation System (Dowell et al. 2001).

At the time of writing, GANESH has been used to construct databases for about a dozen regions of human chromosomes and for three regions of mouse chromosomes. Example databases and further details can be found at <http://zebrafish.doc.ic.ac.uk>. In this work, we describe the structure of GANESH and its components, and demonstrate its role in the genetic analysis of several regions of the human genome.

RESULTS

GANESH: Overview

A GANESH application has the following main components:

- the assimilation module (including downloading scripts, sequence analysis packages, and sequence database searching tools);
- the database;
- the updating module;
- the graphical front-end;
- visualization software; optional analysis tools; utilities.

Figure 1 provides an overview of the system structure.

For construction of a new application, GANESH is focused upon a region of the genome (human or other) by first identifying DNA markers or genomic positions that flank the region of genetic interest. These markers, in turn, are used to define a set of DNA clones that span the interval and that have been, or are being, sequenced. Several databases can be used to select these clones including the UCSC golden path (Kent and Hauser 2001), Ensembl, or the total human fingerprint database (McPherson et al. 2001). The next step is to specify one or more sources of DNA sequences for these clones. In principle, any site is acceptable, provided the files have a fixed location and there is a well-defined update file structure for sequences still to be collected. We have successfully used the Sanger Institute and EMBL. For regions being sequenced by the Sanger Institute, we download from the relevant ftp archive (<ftp.sanger.ac.uk/pub> and [human/sequences](ftp.sanger.ac.uk/pub/human/sequences)) and

check daily for updates. For other regions, the sequences are retrieved initially from EMBL and updates extracted from the EMBL daily update files available from the European Bioinformatics Institute (EBI) ftp site (<ftp.ebi.ac.uk/pub/databases/embl/new>).

All sequence data from the target region is thereafter downloaded by the assimilation module and processed using a range of (standard) genome analysis tools. The standard configuration presently includes RepeatMasker (<http://ftp.genome.washington.edu/RM/RepeatMasker.html>), Genscan (Burge and Karlin 1997) for exon predication, poly(A) addition site and promoter finding, and Pfam analysis (Bateman et al. 2002) using the Wise/halfwise package available from the Sanger Institute (<http://www.sanger.ac.uk/Software/Wise2>). We have chosen this set of programs to provide maximal flexibility for the analysis of any region of any genome; there are, of course, a number of sources of precomputed data, for example, repeats in human DNAs, that we could have used, but this would necessarily have significant implications on the range of DNAs that could be studied with GANESH.

BLAST searching (Altschul et al. 1990) is performed for all of the sequences against EMBL, dbEST, SWISS-PROT, and TrEMBL for homology, and against dbSNP and dbSTS for identity. These searches are all performed on local copies of the databases. Parameters for the different kinds of BLAST searches, and for the other analysis packages, can all be specified by the user though the standard configuration defaults to settings that we have determined to be most useful. Similarly, the configuration of components can also be adjusted. The system is designed to be modular and user-configurable; new components—new genome analysis tools, other forms of sequence database searches, in-house annotations (e.g., from in-house experiments)—are added easily.

The results of all of these computations are compressed and stored in a standard relational database (specifically, the MySQL system (<http://www.mysql.com>), which we chose because it is

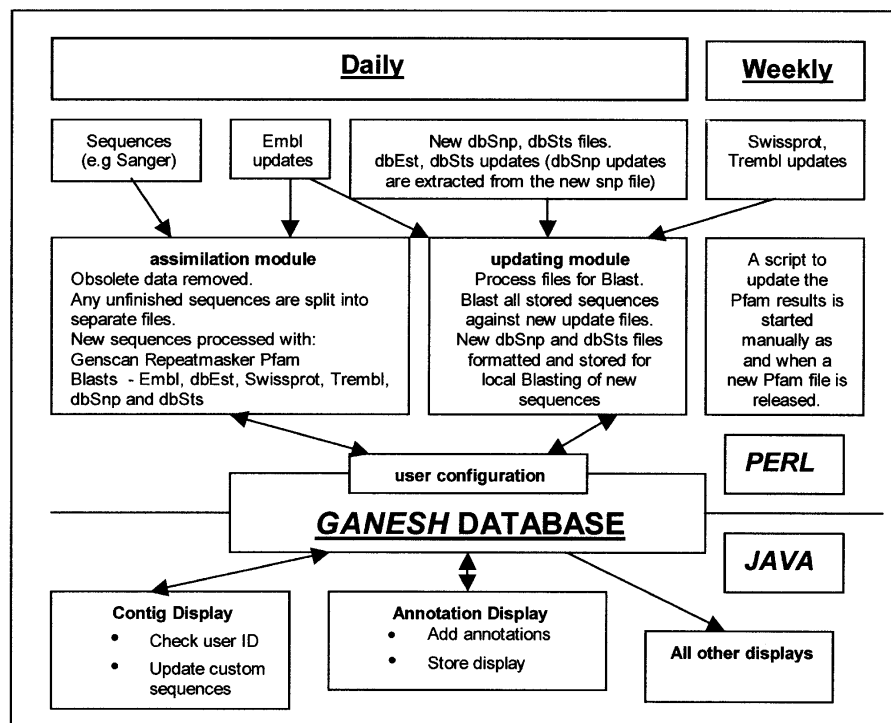


Figure 1 GANESH Overview.

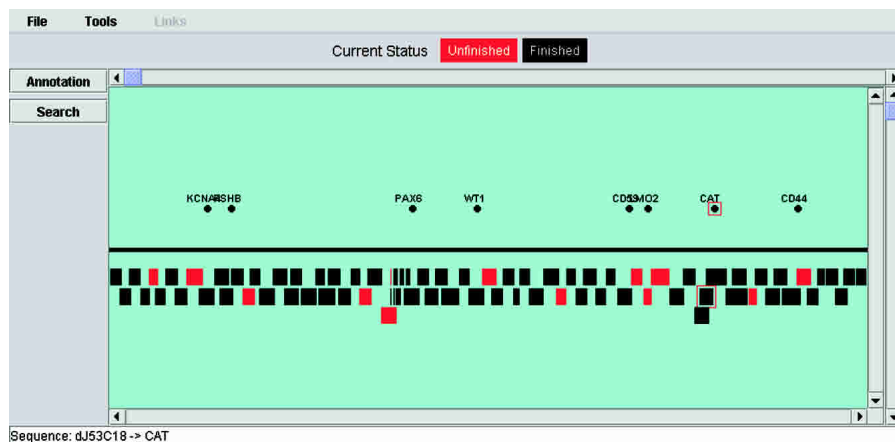


Figure 3 Contig display. The initial display shows the minimal tiling path for the region (in this case the chromosome 11 WAGR region) with markers above to assist in navigation. The markers are linked to the associated sequence clones and vice versa. The display can be zoomed in and out. From it, the Annotation Display can be launched. Window image is clipped for presentation purposes.

Neomorphic Genome Software Development Kit (<http://www.neomorphic.com>), with which it shares some look and feel. Interactive access to GANESH, allowing individual user annotations, is controlled by a user login.

Figure 3 shows the opening screen of the display, in this case a human chromosome region, 11p13, that is deleted in WAGR syndrome patients. The display can be scrolled, and zoomed in and out to focus on areas of interest. Selection of a clone in this screen (by clicking) accesses information about it and (by double-clicking) brings up the annotation display shown in Figure 4. The content of this display is configurable. The user can also add or remove displayed features as required during viewing. As with all displays, it is also possible to zoom in and out; increased detail (names, markers, etc.) appears automatically as display resolution allows.

Clicking on any of the individual features brings up details of the annotation, depending on the nature of the probe. There is a separate graphical display for viewing BLAST hits. Clicking on a BLAST feature brings up the BLAST window shown in Figure 5. This can be scrolled and zoomed as usual. The BLAST significance level for display is controlled with a slider (shown to the left of the screen in Fig. 5), which is used to specify the stringency threshold for displayed BLAST hits. Clicking on a BLAST hit brings up a detailed view of the BLAST alignment, as shown in Figure 6. Links are created automatically to access source data, via the WWW, from GenBank, Entrez, dbSNP, UniGene, and SRS. Because of security restrictions, it is not possible to copy and paste from an applet. Some output is therefore displayed in a separate browser window via a CGI Perl script to enable the user to cut and paste information into other software. Additional windows have also been created to display the complete FPC clone map of the region (Marra et al. 1997), YAC-based maps, and several radiation hybrid maps (James et al. 1994; Deloukas et al. 1998). As with all displays, these can be scrolled and zoomed to access display details at varying levels of resolution.

A key feature superimposed on these displays is the automatic notification of updated information from the daily/weekly BLAST analyses or resulting from updates to the sequence itself. The last viewed version can be retrieved for comparison to the latest information. There is a further option to view just the information added since the user last accessed the system. This feature is particularly useful when combined with the tools provided for users to add their own annotations. These are keyed to

individual operators and are intended to be the primary tool for recording acceptance or rejection of features within the displays—genes, gene-like objects, SNPs, regulatory regions, etc. It is important to stress that these annotations do not necessarily have to represent definitive genome features as in the databases of record, and can take the form of a simple statement of the form ‘gene X is located at position Y’ in the sequence. The purpose of GANESH is primarily to facilitate the identification of as complete a list as possible of genes and other genome features of interest within the target region, and only secondarily to provide definitive descriptions of their structures.

Unfinished Sequences

A complicating factor in both the processing and display of a sequence is the presence of unfinished DNA sequences. These are treated by GANESH as a series of smaller sequences contained within a large BAC clone; in order not to mislead, the display has a prominent warning that these fragments are being located in the display in arbitrary order. As the appropriate genome center further finishes these sequences, GANESH updates its displays so that the number of subsequences for the BAC decreases and their length increases. Eventually, they are displayed as a continuous finished DNA sequence.

All annotations, and user annotations in particular, have to be updated as each new release of the sequence is assimilated. We deal with this by first performing a Smith-Waterman comparison of the new sequence with the previously stored version to identify the parts that have changed. We use crossmatch (<http://www.phrap.org/phrap.docs/general.html>) for this purpose. Typically, large fragments of the sequence will have remained the same. Only those parts that are new are reprocessed (subject to the sequence length requirements discussed below), thus reducing the computational effort, and more importantly, preserving user annotations as far as possible. As already mentioned in the description of the display components, users are able to compare updated sequences with the previously stored and analyzed versions.

Clearly, the problems of dealing with unfinished sequence for the human genome will reduce as more and more genome sequence is finished, but the need to deal with unfinished sequence will not disappear. Although GANESH was developed originally to support the genetic analysis of human genomic regions, it is being used for other model organisms as well (at present, for mouse), and it is expected that it will be used increasingly for other organisms, including organisms whose genomes will remain in largely unfinished draft form.

Long Sequences

In some cases, the regions that are displayed within a GANESH application are not organized around (BAC) clones, but are contained within long, continuous, merged DNA sequences. This is particularly true of those chromosomes and regions that have been the subject of concentrated finishing analyses, such as human chromosomes 21 and 22, and now increasingly more and more regions of the other chromosomes. Most of the analysis packages deployed within GANESH work optimally on sequences up to ~250 Kb, and thus, initially the sequences are cut into 300-Kb lengths with a 50-Kb overlap. Any feature

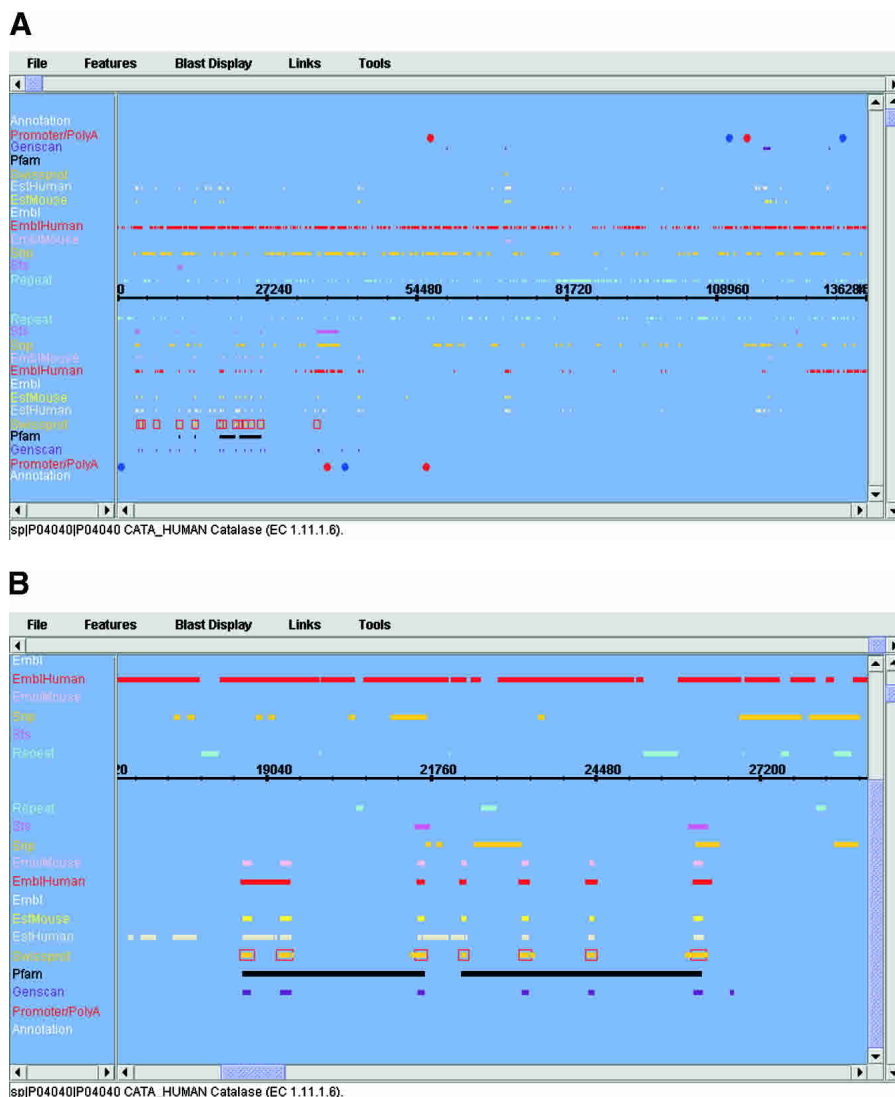


Figure 4 Annotation Display. (A) The Annotation display provides an overview of the features of the sequence. For the application version, a configuration file can set the features to be displayed. The display can be scrolled and zoomed. From this display, the details of an individual feature, including a BLAST hit or all of the BLAST hits, can be viewed. There are also links to several external data sources and options to view the sequence (or partial sequence) in fasta format or a six-frame translation. Window image is clipped for presentation purposes. (B) Zoomed view, at small-medium resolution.

that starts within the first 270 Kb of a fragment is stored, and those that start after this position are ignored, as the next subsequence will record them. For Genscan, sequences are broken into 3-Mb fragments with 300-Kb overlaps, as this software requires a longer sequence context to operate optimally. GANESH automatically detects large sequences and displays a sliding selection box to allow the operator to select a region of the sequence for viewing in the annotation display. The position of the selection box within the sequence is displayed in the text box.

These large genomic sequences also cause some problems of focus, because not all regions are of similar interest to all users. Provision is therefore made within GANESH for the creation of custom sets of sequences. Users can pick out and register subregions and sets of subsequences as being of special interest. For the purposes of the annotation display and update notification, these custom sets are treated in the same way as normal (BAC-

indexed) sequences. The user is notified of any new annotation data within their range as the parent sequences are updated, as usual. The difference is that displays of custom sets of sequence are only available to their registered users. Different users may create their own custom sets, and these regions may overlap, but they are only displayed when the registered user logs in, and then only their own custom sets are displayed.

Gene Identification Tools

An optional GANESH module is provided, which can be used to predict the presence of genes/exons. Predictions can be made by comparing evidence from the following three sources: (1) similarity to known expressed sequences, (2) in silico prediction programs such as Genscan, and (3) similarity to genomic regions of closely related organisms. The strategy adopted is to consider all lines of evidence in parallel and retain all gene/exon predictions, however low the likelihood. This is in contrast to the strategies adopted by databases of record, in which genes are only predicted if several lines of evidence are available. Our reason for adopting this approach is that in any positional cloning strategy, particularly once localization is down to the submegabase level, researchers will wish to be aware of every sequence within that region with evidence supporting transcriptional or regulatory function, less the disease-susceptibility gene be overlooked. We therefore categorize the gene predictions on the basis of the strength of evidence available, enabling the researcher to focus on the stronger candidates initially, but without eliminating other weaker candidates.

The gene predictions are subdivided into:

- category 1 predictions (Ganesh-1): match a known Ensembl gene;
- category 2 predictions (Ganesh-2): evidence from all three main sources of evidence: (1) similarity to known expressed sequences, (2) in silico prediction programs such as Genscan, and (3) similarity to genomic regions of closely related organisms;
- category 3 predictions (Ganesh-3): evidence from two of the three lines of evidence;
- category 4 predictions (Ganesh-4): evidence from a single line of evidence.

We make our gene predictions available to our collaborators by utilizing the Distributed Annotation System (DAS; Dowell et al. 2001; <http://www.biodas.org>). A DAS server (zebrafish.doc.ic.ac.uk:8080) was set up using the Dazzle server (<http://www.biojava.org/dazzle>), enabling our collaborators to view the predictions using the standard Ensembl Web-based browser with the relevant GANESH database selected as an additional DAS source.

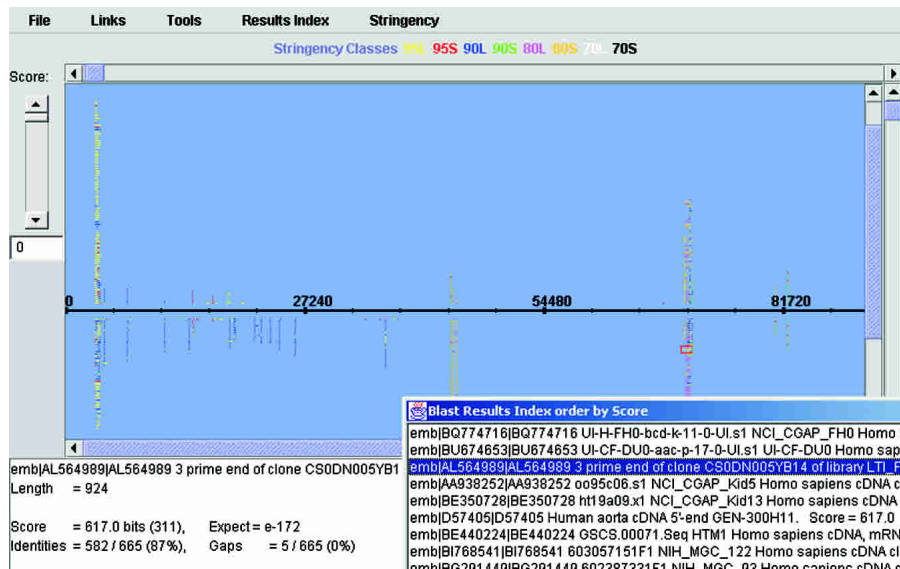


Figure 5 BLAST Display. The BLAST Display shows a summary of all the BLAST hits. The display can be scrolled and zoomed. An index of all of the hits can be displayed optionally—as shown—which is linked to the main display. The slider on the left of the display is used to screen the BLAST hits on minimum score value. Window images are clipped for presentation purposes.

Figure 7 shows the Ensembl display of chromosome 17: 12–12.3 Mbp, highlighting the Ensembl-predicted transcripts and GANESH category 1, 2, 3, and 4 gene predictions. They are summarized in Table 1 for the first 100 Kbp.

At the time of writing, we have not completed experimental work on the predicted genes in this table beyond some preliminary microarray analysis to confirm/discard some of the gene predictions. It seems likely that a proportion of the low-category predictions represent genuine transcripts. According to a recent microarray-based validation of predicted genes on chromosome 22q (Shoemaker et al. 2001), as many as 57% of sequences predicted to be transcribed on the basis of *in silico* evidence alone had their expression confirmed experimentally.

Portability and Requirements

We maintain a GANESH system at Imperial College for our own users and collaborators annotating the regions detailed. The GANESH system itself is freely available for other researchers to install to annotate their own regions, and GANESH can be deployed to any region of the human genome, or to any region of another species once the parameters for the various genome analysis packages have been adjusted accordingly. The default version of GANESH requires the installation of the analysis programs and other software detailed, all of which are either open source or free to academic users. If any of the default analysis programs are not required, the system requires minimal modification to accommodate the changes. GANESH requires access to the EMBL, SWISS-PROT, and TrEMBL databases, and if these are not available locally to the installer, a Perl script is provided to build the required databases, and GANESH auto-

matically maintains the updates. Also required are the installation of Perl, including DBD, DBI, and FTP modules, and Java 1.3. It uses the MySQL database system (although it does not rely on anything specific to that system and can be adapted easily for use with any relational database system). GANESH can be installed and run satisfactorily on a single processor linux machine, either as a standalone or a client-server system. The Java viewer can be run on any operating system that has access to the machine hosting the GANESH system.

The applet requires Netscape 6(+), or for other browsers, the installation of the Java Runtime Environment 1.3.1_02(+) and can be installed on any web server that can access the GANESH installation database.

GANESH is designed to be as straightforward to install and maintain as possible, but some knowledge of the unix/linux operating system is required. Furthermore, some knowledge of Perl would be beneficial in order to modify

the scripts if required, particularly if new analysis programs are to be added.

We have also developed a portable version of the system that allows databases to be stored and queried off-line. A laptop or notebook is sufficient to run this version. These stand-alone databases can be synchronized with the central database over the internet.

The GANESH software is available under an Open Source license. Details may be found at <http://zebrafish.doc.ic.ac.uk>.

DISCUSSION

Our main objective in developing GANESH was to provide a local tool for genetic analysis, but clearly, such studies can contribute to whole-genome annotation efforts. GANESH shares many fea-

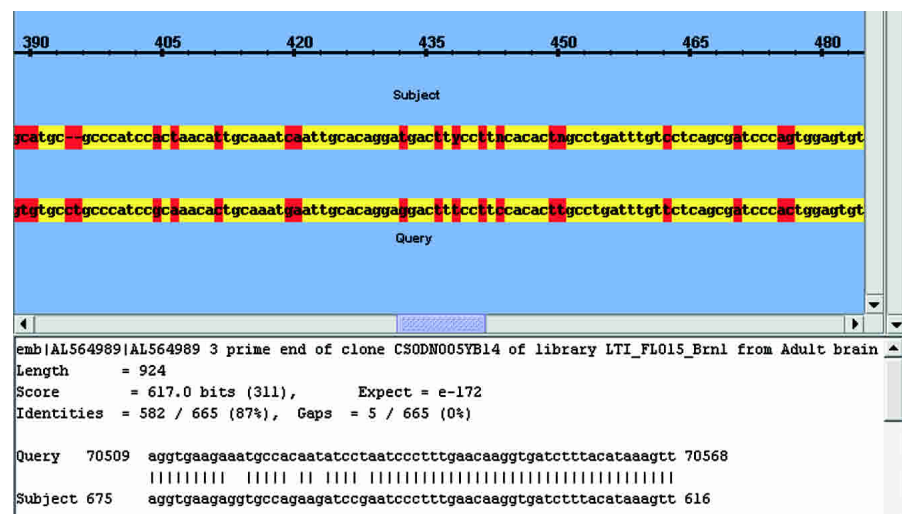


Figure 6 BLAST Alignment Display. The BLAST Alignment Display shows the detail of an individual BLAST hit with the gaps/mismatches highlighted. The display can be scrolled and zoomed. The standard BLAST alignment and details are also provided in a text window from which they can be copied. Window image is clipped for presentation purposes.



Figure 7 Ensembl DAS display. Ensembl display of the 12–12.3-Mbp region of human chromosome 17 highlighting the Ensembl predicted transcripts and GANESH category 1, 2, 3, and 4 gene predictions.

tures with other systems for the automated annotation of genomic sequence. The main distinguishing features of GANESH as we see them are as follows:

1. GANESH provides a means to set up and tailor a sequence database and annotation system for small groups with limited computational resources, or groups dealing with unusual model organisms. We see the latter point as being a significant contribution, as the major annotation systems such as Ensembl focus on a few organisms; GANESH, in contrast, provides the opportunity to annotate any organism of choice.
2. GANESH is designed to support the detailed analysis of relatively small, often <10–20 cM (~10–20 Mb), regions of the genome, and to incorporate varied, and sometimes speculative, tools, external data sources, and in-house annotations and experimental data, that could not be included in the archival databases of record.
3. GANESH provides update mechanisms to ensure that data and annotations for the target region are as up-to-date as possible.

We have recently added facilities for importing and exporting data in the format of the Distributed Annotation System (DAS), enabling a GANESH database to be used as a component of a DAS configuration. A GANESH database of annotated human, and now also mouse, sequence can thus be accessed and viewed in two different ways as follows: (1) via the Ensembl site, using the Ensembl Web-based browser with the GANESH database selected

as an additional DAS source; or (2) using the GANESH front end, either as a stand-alone application or as a web applet, with relevant data extracted from Ensembl and stored in GANESH as desired. The advantage of the first method is that the Ensembl browser is used widely and is increasingly familiar to researchers, and is likely to be the subject of sustained further development. It also provides an easy mechanism for disseminating results to our collaborators (as there is less for us to maintain). The advantages of the second method are that the GANESH front end is (comparatively) easy to adapt and customize, and can also be used as a stand-alone application, which some users seem to prefer. Only the second method is applicable, of course, to the analysis of genomes not presently stored in Ensembl.

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Table 1. GANESH Gene Predictions for the First 100 Kbp of Human Chromosome 17:12-12.3

Ensembl transcripts	Ganesh category 1	Ganesh category 2	Ganesh category 3	Ganesh category 4
ENST00000226207	SQ:chr17 - :8844		SQ:chr17 + :43979	SQ:chr17 + :41289
ENST00000245503	SQ:chr17 - :37683		SQ:chr17 + :27635	SQ:chr17 + :41756
			SQ:chr17 + :8826	SQ:chr17 + :55977
			SQ:chr17 + :32398	SQ:chr17 + :50033
			SQ:chr17 + :24862	SQ:chr17 + :57127
			SQ:chr17 + :58939	SQ:chr17 + :43131
			SQ:chr17 + :44749	SQ:chr17 - :42224
			SQ:chr17 + :17368	SQ:chr17 + :54196
			SQ:chr17 + :15128	SQ:chr17 + :51808
			SQ:chr17 + :61779	SQ:chr17 + :48052
			SQ:chr17 + :20406	SQ:chr17 + :65951
			SQ:chr17 + :63852	SQ:chr17 + :56461
				SQ:chr17 + :49818
				SQ:chr17 - :32446
				SQ:chr17 - :20850
				SQ:chr17 + :31403
				SQ:chr17 + :51606
				SQ:chr17 - :32444
				SQ:chr17 + :48088
				SQ:chr17 - :9840
				SQ:chr17 - :9411
				SQ:chr17 + :55738
				SQ:chr17 + :2773
				SQ:chr17 + :68464
				SQ:chr17 + :65763
				SQ:chr17 + :62328
				SQ:chr17 + :66994
				SQ:chr17 + :23916
				SQ:chr17 + :38183
				SQ:chr17 + :34809
				SQ:chr17 + :72196
				SQ:chr17 + :71100
				SQ:chr17 + :35753
				SQ:chr17 + :99468

ing, Imperial College. We thank Win Hide for many useful discussions and for his continuing support of this project.

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