Multiple Whole Genome Alignments and Novel Biomedical Applications at the VISTA Portal

Michael Brudno, Alexander Poliakov¹, Simon Minovitsky¹, Igor Ratnere¹, Inna Dubchak¹,²,*

Department of Computer Science, University of Toronto, 6 King's College Rd, Toronto, ON M5S 3G4, Canada; ¹Genomics Division, MS 84-171, Lawrence Berkeley National Laboratory, Berkeley, CA 94720 USA; ²US Department of Energy Joint Genome Institute, Walnut Creek, California 94598, USA.

To whom correspondence should be addressed Tel: +1 510 495 2419; Fax: +1 510 486 5614; Email: ildubchak@lbl.gov

ABSTRACT
The VISTA portal for comparative genomics is designed to give biomedical scientists a unified set of tools to lead them from the raw DNA sequences through the alignment and annotation to the visualization of the results. The VISTA portal also hosts alignments of a number of genomes computed by our group, allowing users to study regions of their interest without having to manually download the individual sequences. Here we describe various algorithmic and functional improvements implemented in the VISTA portal over the last two years. The VISTA Portal is accessible at http://genome.lbl.gov/vista.

INTRODUCTION
Comparing genomic sequences across related species is a fruitful source of biological insight. Functional elements such as exons tend to exhibit significant sequence similarity due to purifying selection, whereas regions that are not functional tend to be neutrally evolved and thus less conserved. The first step in comparing genomic sequences is to align them — to map the letters of one sequence to those of the others. After an alignment is computed, visualization frameworks become essential to enable users to interact with the sequence and conservation data, especially in the context of longer DNA sequences or whole genomes. Visualization frameworks should be easy to understand by a biologist and provide insight into the mutations that a particular region has undergone.

The VISTA portal is a comprehensive comparative genomics resource that provides biomedical scientists with a single unified framework to generate and download multiple sequence alignments, visualize the results in the context of existing annotations and analyze comparative results in search for important sequence signals in alignments.

The VISTA suite of programs has been in development and continued use since 2000 [1-4]. It was originally developed for the alignment and comparative analysis of long genomic sequences and later was expanded to pair-wise and three-way alignment of vertebrate genomes. VISTA has popularized the visualization of the level of conservation in the format of a continuous curve based on the conservation in a sliding window. These concepts proved to be extremely successful due to the easy interpretation of the resulting plots.
VISTA was built through a close collaboration between computational and biological scientists, resulting in a product that is robust, efficient and powerful, yet simple to use for a person without extensive computer experience, as is illustrated by more than 1000 publications citing the various VISTA-associated tools (http://scholar.google.com).

In the last two years the VISTA portal has seen many significant improvements. In addition to updating the whole genome alignments, computed using recent assemblies of vertebrate, insect, plant and microbial genomes, we have added significant new functionality and resources to the Genome Browser and other tools, including:

- A novel multiple whole genome alignment algorithm
- A new server for whole-genome alignment of bacterial genomes
- Base-pair level visualization ability within the VISTA browser
- Visual access to the results of the prediction of potential deleteriousness of non-synonymous Single Nucleotide Polymorphisms (SNPs) by the algorithm PolyPhen [5].
- A novel conservation track, Rank-VISTA, to shows the statistical significance of conserved regions computed by the Gumby algorithm [6].
- Whole-genome rVISTA, that allows for evaluation of which conserved transcription factor binding sites (TFBS) are over-represented in a group of genes.

**VISTA PORTAL**

The suite of VISTA tools is accessible through the website http://genome.lbl.gov/vista. Currently it includes five servers for the analysis of user-submitted sequences: mVISTA that computes alignments of user-submitted sequences; wgVISTA to align whole genomes (up to 10 megabases in length); GenomeVISTA that aligns a user-submitted sequence to a selected genome assembly; rVISTA that searches for conserved transcription factor binding sites; and Phylo-VISTA to analyze multiple DNA sequence alignments of sequences from different species while considering their phylogenetic relationships. In addition, multiple whole-genome alignments of vertebrate, insect and plant species have been built using in-house algorithms and are publicly available for browsing and analysis. The portal provides access to the VISTA Genome Browser, the main visual interface for both the pre-computed whole genome alignments and alignments of user-submitted sequences. In the sections below we will discuss various algorithmic and functional improvements to the VISTA portal in the last two years.

1. Algorithmic improvements

Most of the algorithms for multiple alignment either rely on a reference genome, against which all of the other sequences are laid out, or require a one-to-one mapping, where each nucleotide of one genome is constrained to align to at most one place on the other genome [4, 7-9] Both approaches have drawbacks for whole-genome comparisons. The first approach requires computation, storage, and analysis of several alignments, one for each base species. The second approach fails to align any gene that has undergone duplications since the divergence of the species being compared. Additionally, “referenced” alignments commonly fail to include elements conserved among some
genomes, but missing in the base genome. We have developed and implemented a novel alignment algorithm that treats all genomes symmetrically.

**Improved alignment pipeline.** Initially our whole genome alignment pipeline used an alignment strategy where one genome was split up into contigs of about 250 Kilobases (Kb) [3, 4]. The potential orthologs for each contig were found in the second genome with the BLAT local aligner (Kent 2002). This step was followed by a global alignment of two orthologous sequences. Although this approach produces a map that is more accurate within large syntenic blocks than all-by-all local alignment, it has two main weaknesses: (1) small syntenic blocks, resulting from rearrangements within a larger region, may be missed; (2) the initial arbitrary division of one genome into segments can split a syntenic region, making it difficult to map the region to its true orthologue.

To address these issues we have developed a “glocal” alignment method which treats the rearrangement events explicitly. The more recent algorithms for whole genome alignments attempt to incorporate the likely evolutionary events as “operations” into their scoring schemas. There have been several algorithms that decide whether to accept or reject a local alignment based on other alignments near it, and thus allow for the direct treatment of the various rearrangement events, including Shuffle-LAGAN (S-LAGAN, [10]) which currently serves as the underlying engine for our whole genome alignments. The pair-wise algorithm described below is based on a novel chaining tool, called SuperMap. The multiple alignment algorithm is a progressive extension of the pair-wise one, where at every internal node we pick an ordering of the alignments that simplifies the next alignment that we will conduct. The algorithms for multiple alignment will be discussed in detail in a separate publication (Brudno et al. in preparation).

**A. Pair-wise Alignment.** To align two genomes this algorithm uses a novel approach based on a reimplementation of the original S-LAGAN chaining algorithm [10, 11] combined with a novel post-processing stage called SuperMap. The S-LAGAN chaining takes as input a set of local alignments between the two sequences generated by BLAT [12] or any other local aligner and returns the maximal scoring subset of these under certain gap criteria. This subset is called a 1-monotonic conservation map. In order to allow S-LAGAN to catch rearrangements, the map is allowed to be non-decreasing (monotonic) in only one sequence, without putting any restrictions on the second sequence. The 1-monotonic chain can capture all rearrangement events besides duplications in the second genome. In order to allow our alignments to incorporate these events we have introduced the novel SuperMap algorithm that takes two S-LAGAN outputs to make our algorithm symmetric. We run S-LAGAN twice, using each sequence as the base. This gives us three pieces of data: the original local alignments, which were common to the two runs of S-LAGAN, and two chains of these alignments, each corresponding to the S-LAGAN 1-monotonic maps. We then classify all local alignments as belonging to both chains, and consequently orthologous (best bi-directional hits) or being in only one chain, and hence a duplication (see Figure 2 for a graphical overview of the algorithm).
SuperMap has several advantages over regular S-LAGAN: 1. It is able to locate duplications in both sequences, a major weakness of the original algorithm; 2. In case of translocations, two of the pieces are no longer arbitrarily joined together; 3. This approach locates both regions of one-to-one similarity (those that were in both 1-monotonic chains) and likely duplications (those in only one chain).

**B. Progressive Multiple Alignment.** After the SuperMap algorithm is used to align the two pairs of sister taxa we use a progressive generalization of the pair-wise SuperMap algorithm to align all of the genomes, by following the species’ phylogenetic tree. After aligning two genomes, our algorithm joins together syntenic blocks (regions of genomes without rearrangements) based on their order in the outgroups (those sequences that will be aligned at a later stage: for example if we have aligned mouse with rat, then human, dog, and chicken are all outgroups). We use an algorithm based on finding a maximum weighted matching in a graph, with the weights specified by the outgroup genomes, to order the individual alignment blocks in the order that will create the simplest alignment problem when we align the result to the ancestor, and use the SuperMap based pair-wise alignment algorithm algorithm to align the alignments to each other using the regular LAGAN aligner [13]. This algorithm is summarized in the flowchart in Figure 3.

By picking an order of the syntenic blocks which is closest to the outgroup we facilitate alignment of the more distant genomes. Our approach has several advantages over previous algorithms: 1) it does not assume a base genome, to which all other genomes are aligned, but creates a symmetric alignment equally valid for all genomes; 2) it penalizes various rearrangement events based on an evolutionary tree, creating a set of alignments that mirrors the evolutionary history of the sequences; and 3) it is able to align short, low similarity syntenic blocks based on their adjacency to higher similarity areas even when there has been a rearrangement event between them.

**C. wgVISTA: Whole Genome Alignment for user’s genomes.** In order to allow our users to compare whole genomes using the whole genome alignment algorithms described above we have developed whole genome VISTA (wgVISTA), a tool which accepts sequences up to 10 megabasepairs in length, aligns them using our alignment pipeline and visualizes the results through the VISTA browser.

**2. Visualization improvements**

The VISTA Browser allows for the exploration of alignment and annotations of DNA sequences. It shows any number of alignments on a particular base genome and is scalable to the size of whole mammalian chromosomes. At the larger scale, visual presentation of rearrangements, inversions, and gaps in the alignment are also available through the browser. Because all of our alignments are built in a symmetric fashion (see above section) the user may select any sequence or a genome as the reference or base, and display the level of conservation between this reference and the sequences of other organisms. The browser has a number of options, such as zoom, extraction of a region to be displayed, user-defined parameters for conservation level and selection of sequence elements for study. We have recently introduced two significant improvements to the
browser that allow for a more detailed analysis of the areas of conservation detected in alignments. The VISTA Browser also gives access to the Text Browser that provides a user with all data related to alignments, analysis of conservation, and access to other resources.

The scrollable nucleotide-level alignment window displays not only the details of the underlying pair-wise or multiple alignment, but also additional nucleotide-level annotation such as the SNPs (Figure 1). Unlike the main VISTA window, the base-pair window does not have a selected continuous base sequence, but rather shows a real pair-wise or multiple alignment where a user can analyze gaps and substitutions in any sequence.

Another visual representation of conservation is the RankVISTA plot, that allows the users to judge the statistical significance of any conserved region. It produces a histogram-like plot where block width is proportional to median conserved element length in human, and block area is proportional to the median of $-\log(P\text{-value})$ [6]. Block height thus represents degree of evolutionary constraint at the basepair level.

Finally we have integrated the mVISTA server for user-submitted sequences with the VISTA Browser: when a user submits sequences to mVISTA, instead of just being e-mailed the VISTA curves in a PDF document, we now make the alignment into a track on the VISTA browser, allowing the user to zoom in on a region of interest and view the detailed alignment in the nucleotide window.

3. New applications

One of the main emphases of our development in the last two years has been on better integration of the existing VISTA tools and of novel biological data into the VISTA portal. Several new applications developed in the group and by the collaborators have been integrated in the VISTA portal, allowing biologists to easily access and visualize the results of these analyses.

**Whole-genome rVISTA.** Gene expression studies generate extensive lists of co-expressed genes which can share regulatory factors that control their synchronous expression. We have developed a computational tool, called Whole-Genome rVISTA, designed to discover which conserved between pairs of species transcription factor binding sites (TFBS) are over-represented in upstream regions in a group of genes. The tool uses whole-genome alignments computed in the group, and TRANSFAC Professional from Biobase [14] with the MATCH program [15] to predict TFBS. The effectiveness of Whole Genome rVISTA was recently illustrated in a study of responsiveness to cAMP regulation [16]. That expression study indicated that several circadian rhythm clock genes are induced by cAMP. We used Whole Genome rVISTA to scan 5Kb upstream of the transcription start site of the cAMP-regulated genes, and found that up-regulated genes contained more cAMP Response Elements (CRE) than all other genes on the array.
**Gumby in RankVISTA.** With more genomes available it has become essential to introduce new statistically motivated methods for conservation analysis that take into consideration neutral rates and phylogeny of the species. Gumby [6] makes no prior assumptions about evolutionary rates and requires no adjustment of parameters as the phylogenetic scope is varied from primates to vertebrates. Gumby uses a dynamically generated phylogenetic log-odds scoring scheme to identify local segments of any length that evolve slower than the background neutral rate, and ranks these conserved segments by P-value using the Karlin-Altschul statistic. This scoring technique demonstrated its efficiency in analyzing conservation both in evolutionary distant [17], and very close [6] [18] [19] species. Rank-VISTA plots of Gumby analysis allow the users to judge the statistical significance of any conserved regions and are available through VISTA Browser for genome-wide alignment of a number of genomes as well as for user-submitted mVISTA queries (Fig 1).

**PolyPhen on the nucleotide alignment track.** (Poly/morphism /Phen/otyping) [5] is a tool which predicts possible impact of an amino acid substitution on the structure and function of a human protein using straightforward physical and comparative considerations. For each non-synonymous SNP in the dbSNP database [20] the VISTA Browser provides access to the results of the PolyPhen analysis of its deleteriousness.

**FUTURE DIRECTIONS**

The main emphasis of our future work within the VISTA portal will be on integration of additional data that is necessary for biological and medical researchers to carry out their analyses. We plan on integrate into our portal information about human variation, especially where it is known that some variation has a correlation with medical disorders. We will also continue to work on providing the users with a simple-to-use interfaces for browsing genomic data – we have been developing methodologies to display various evolutionary events in the context of the underlying phylogenetic trees [21, 22] and expect to make similar improvements for visualizing rearrangements between the various genomes.

Finally the new alignment pipeline implemented within the VISTA portal should be both flexible and powerful enough to analyze many of the genomes that are currently being sequenced. Consequently the majority of our alignment-related work in the near future will be on maintaining up-to-date versions of novel genomes, including low coverage genomes that are currently being sequenced.

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Figure 1. VISTA Browser display of 7.8 Kb fragment of LEP gene on Chr. 7 of the human genome (hg17). VISTA plots for the 4-way Human-Mouse-Dog-Rat alignment are shown. Conserved sequences in VISTA (70%/100 bp cutoff) are colored according to the annotation (exons - dark blue, UTRs – turquoise, non-coding - pink). Rank-VISTA peaks identified by Gumby ($P$-value < 0.5) are shown as vertical bars following the same coloring convention. At the bottom of the window one can see the base-pair browser and SNP data (dbSNP annotation and PolyPhen [5] prediction of functionality for coding SNPs).
Figure 2: SuperMap Algorithm
a) Local alignment hits: regions A & B correspond to duplications in Organism 1; regions C & D correspond to duplications in Organism 2; b) S-Lagan chain for Organism 1 as a base. Chain increases in direction of X axis, but can jump up and down in Y direction (Organism 2), region D is left out; c) S-Lagan chain for Organism 2 as a base - chain increases in direction of Y axis, region B is left out; d) SuperMap output - combines regions of Figures b & c - regions that are in both maps are colored green.

Figure 3: Multiple alignment with LAGAN in the VISTA Genome Pipeline (VGP). After running a local alignment program, SuperMap Chaining is used to identify all rearrangements. The resulting regions are aligned with LAGAN, and finally a maximum matching algorithm is used to predict ancestral contigs. These ancestral contigs are then used to align to outgroup genomes in the higher levels of the phylogenetic tree.