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Multiple-sequence functional annotation and the generalized hidden Markov phylogeny

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ABSTRACT

Motivation: Phylogenetic shadowing is a comparative genomics principle that allows for the discovery of conserved regions in sequences from multiple closely related organisms. We develop a formal probabilistic framework for combining phylogenetic shadowing with feature-based functional annotation methods. The resulting model, a generalized hidden Markov phylogeny (GHMP), applies to a variety of situations where functional regions are to be inferred from evolutionary constraints.

Results: We show how GHMPs can be used to predict complete shared gene structures in multiple primate sequences. We also describe SHADOWER, our implementation of such a prediction system. We find that SHADOWER outperforms previously reported *ab initio* gene finders, including comparative human–mouse approaches, on a small sample of diverse exonic regions. Finally, we report on an empirical analysis of SHADOWER's performance which reveals that as few as five wellchosen species may suffice to attain maximal sensitivity and specificity in exon demarcation.

Availability: A Web server is available at http://bonaire.lbl.gov/ shadower

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INTRODUCTION

The prediction of functional regions in genomic sequences has traditionally been based on the identification of features associated with genes or regulatory regions (Zhang, 2002). Comparison of homologous genomic sequences facilitates such identification, because functional regions tend to be conserved in sequences that have evolved from a common ancestor, whereas non-functional regions are more likely to mutate. Information about the degree of conservation between pairs of sequences is known to help in the identification of genes (Alexandersson *et al.*, 2003; Korf *et al.*, 2001; Parra *et al.*, 2003; Meyer and Durbin, 2002). Indeed, homology

plays a role in a number of classical sequence analysis methods, such as the BLAST tool suite.

One drawback of pairwise comparative approaches to gene prediction is that non-functional regions are required to have diverged to a degree that enables statistical procedures to distinguish them from biologically active regions; typically, organisms such as human and mouse are used. These methods are therefore not applicable to discovering features present only at close evolutionary proximity, such as primate-specific genes. The phylogenetic shadowing principle of Boffelli et al. (2003) circumvents this problem by seeking to identify conserved regions among multiple closely related organisms. This has numerous advantages: the alignment of the sequences is straightforward, the phylogenetic tree relating the sequences is easy to infer and the identification of conserved regions among the sequences is possible using standard evolutionary models. The principle has been illustrated by Boffelli et al. (2003) in the identification of transcription factor binding sites in the primate-specific apo(a) gene.

To provide a systematic computational methodology for annotating genomic sequences based on the principle of phylogenetic shadowing, we have developed the generalized hidden Markov phylogeny (GHMP). The GHMP is a formal probabilistic model that combines conservation-based constraints deriving from multiple genomic sequences with algorithmic ideas that have proven useful in single-organism gene annotation systems. Our approach synthesizes generalized hidden Markov model (HMM) gene finders, evolutionary models of nucleotide substitution and phylogenetic trees. Similar ideas have been presented by Pedersen and Hein (2003) and Siepel and Haussler (2003); our extensions and contributions are described in the Methods section. We have also implemented SHADOWER, a gene prediction system based on these ideas. We show that SHADOWER outperforms existing ab initio methods, including those taking comparativegenomics approaches, on a multiple-primate dataset of single exons from five separate gene regions. Furthermore, we present an empirical analysis of SHADOWER's performance

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on various subsets of our primates which reveals that just five species, selected according to a formal optimality criterion, suffice to deliver the best results SHADOWER can obtain for these data.

The remainder of the paper proceeds as follows. The Methods section presents theoretical and computational details of the GHMP, placing the GHMP within the general formalism of probabilistic graphical models. We report and discuss the data, parameter estimation procedure and subset-selection optimization underlying our full empirical analysis in the Results section. Finally, the Discussion section offers concluding remarks and outlook.

METHODS

The generalized hidden Markov phylogeny

Graphical models combine ideas from probability theory and graph theory to facilitate the use of sophisticated joint dependency structures in data analysis (Cowell *et al.*, 1999; Jordan, 1999). The nodes of a graphical model correspond to random variables that relate to the problem and data at hand. The edges in the model encode marginal and conditional independencies among these random variables, according to a well-defined formal semantics. The GHMP is a directed graphical model—a model in which the underlying graph is directed and acyclic. In such models, sometimes referred to as Bayesian networks, there is a local conditional probability distribution associated with each node in the graph, conditional on its parents. The joint distribution over all random variables is defined to be the product of these local conditional distributions.

This section details the variables, independence structure and local distributions peculiar to the GHMP. The graphical model perspective allows us to focus our attention on capturing, in the model definition, the essential ingredients of the multi-sequence functional annotation problem. Then, parameter estimation and probabilistic inference are handled using general-purpose graphical modeling algorithms (Jordan, 1999).

Many biologists are already acquainted with special cases of graphical model methods that preceded the recognition and elaboration of the general framework. Phylogenetic trees can be treated as graphical models, and the likelihood computation of Felsenstein (1981) is an instance of the generalpurpose junction tree algorithm for graphical models (Cowell *et al.*, 1999). Similarly, the forward, backward and Viterbi algorithms for inference in HMMs are also special cases of the junction tree algorithm. The GHMP synthesizes the ideas of HMMs and phylogenetic trees. The corresponding algorithms can be seen, on the one hand, as a synthesis of the HMM and tree inference algorithms, or, on the other hand, as just another instantiation of the universal graphical model procedures.

Combinations of HMMs and evolutionary models have been described previously by Pedersen and Hein (2003), Siepel

and Haussler (2003), Felsenstein and Churchill (1996), Yang (1995) and Goldman *et al.* (1996), and our methods build on these earlier works. The GHMP and SHADOWER introduce:

- Generalized hidden Markov dynamics (non-geometric exon length distributions).
- A frame- and phase-consistent dual-strand hidden state space, supporting single-exon and multi-exon gene prediction (including incomplete multi-exon genes).
- GC isochore-specific parameters.
- Deterministic constraints on repeats, gaps and in-frame stop codons inside aligned exons.
- More complete splice site modeling.
- An automated iterative procedure for alignment and tree building.
- An analysis methodology for optimal species subset selection.

The principle behind our treatment of gaps also differs, as described below. The reader will recognize several ideas from the currently best-performing gene finders (Alexandersson *et al.*, 2003; Korf *et al.*, 2001; Burge and Karlin, 1997; Parra *et al.*, 2003); indeed, our work represents an attempt to import these ideas into a phylogenetic framework. Most important of all, we adhere to the phylogenetic shadowing principle in our exclusive use of closely related species. This motivates and justifies the seemingly stringent requirements we impose; e.g. we rely on exactness of the multiple alignment and so consider only perfectly aligned, gapless splice signals and start/stop codons. This approach should be contrasted with comparison-based gene finders using distantly related organisms, which generally must search in the enormous space of possible alignments.¹

We begin by recasting familiar phylogenetic tree representations within the graphical model framework. Consider the rooted five-primate phylogenetic tree presented in Figure 1A, and the corresponding graphical model shown in Figure 1B. Both diagrams indicate the presence of a specific set of nucleotides at homologous sites in the five primates, and also indicate putative ancestral nucleotides. The distinction between observed nucleotides and ancestral nucleotides is captured in the graphical model by shading; in general, observed random variables are shaded, whereas unobserved (hidden) random variables are left unshaded. In Figure 1A, edge lengths are proportional to evolutionary distance. In the graphical model, on the other hand, edge lengths are uninformative. Instead, the pattern of edges formally encodes the following probabilistic assumption: given the nucleotide of an organism's immediate ancestor, that organism's nucleotide is conditionally independent of all other ancestral nucleotides.

¹For example, there are approximately 10¹⁸ distinct alignments of five sequences, when each sequence is only five bases long.



Fig. 1. Two alternative representations of the same rooted phylogenetic tree. (A) A diagram familiar to biologists, with annotated edge lengths reporting evolutionary distances. (B) The corresponding directed graphical model, in which nodes are random variables (nucleotides) and the pattern of edges encodes Markovian conditional independence assumptions. Shaded nodes (the leaves of the tree) are observed (the entries of an aligned column); unshaded nodes are latent (the unknown orthologous bases of ancestor species).

To complete the specification of the phylogenetic tree graphical model, we require a local conditional distribution for each node v. At nodes other than the root, this distribution is given by an evolutionary nucleotide substitution model: for each possible initial parent nucleotide, the distribution specifies the probability that v evolved to a terminal nucleotide A, C, G or T in time b at some evolutionary substitution rate. In the GHMP, the branch-length parameter b is specific to each node, while one substitution rate is shared by all non-root nodes. Finally, the distribution π of the root can be any probability distribution over the four bases, e.g. equilibrium base frequencies in a region.

In our implementation, we use the Felsenstein substitution model for the conditional distributions (Felsenstein and Churchill, 1996). This model requires as parameters both a transition-transversion ratio and an equilibrium distribution over bases; we take the latter to be the same as the root distribution π .

The GHMP uses this phylogenetic model to define a probability distribution on a single column of a multiple alignment. To define a probability distribution on a full alignment, the GHMP includes additional nodes that represent functional states. In the implementation of the GHMP that we consider in this paper, the functional states include intergenic regions, introns, coding exons and coding exon boundaries (the last includes splice sites, start codons and stop codons). State variables are unobserved (hidden) variables and thus are left unshaded. The nodes representing these variables are arranged as a chain in the graphical model, with the edges between these nodes representing the probability of transitioning between specific values of functional states; this is the graphical model representation of a Markov chain.

The space of allowed functional states is structured to enforce frame and phase consistency across exons, with genes on either strand, as in Kulp *et al.* (1996) and Burge and Karlin (1997). Figure 2 gives a diagrammatic depiction of this state space (see caption). We foresee a major application of the GHMP to be the annotation of complete primary transcript structures, and our choice of functional state space reflects this. Note, however, that by choosing to start and/or finish in an intronic state, the GHMP can predict incomplete multi-exon genes as well. We return to this point in the Results section.

Let us refer to the collection of adjacent columns generated from a single functional state variable as a *slice*. Suppose the total length of the alignment is N and that the hidden Markov chain comprises $K \leq N$ realizations of the state variables. Since each state variable generates exactly one slice, there are K slices s_1, \ldots, s_K as well, having corresponding lengths d_k , with $\sum_k d_k = N$. Write z_k for the value assumed by the k-th state variable, $p(d_k|z_k)$ for the probability that a slice of length d_k comes out of state z_k and $p(s_k|z_k, d_k)$ for the probability of the observed slice s_k given the postulated functional state and slice length. The joint probability distribution over hidden state variables, slice lengths and observed slices under the GHMP is then

$$p(z_1)p(d_1|z_1)p(s_1|z_1,d_1)$$

$$\times \prod_{k=2}^{K} p(z_k|z_{k-1})p(d_k|z_k)p(s_k|z_k,d_k),$$

where $p(z_1)$ is an initial probability and $p(z_k|z_{k-1})$ is a transition probability for the functional state chain.

We now describe the structure of the emission probability distributions $p(s_k|z_k, d_k)$ and length distributions $p(d_k|z_k)$. Consider first the simplest case, that of the intron state, in which a slice containing only a single column is associated with the state (Fig. 3). In the figure, the chain of functional



Generalized hidden Markov phylogeny



Fig. 2. The state space of biological functions through which the GHMP maneuvers. This is not a graphical model; it is a diagram depicting which functional states (exon, intron, intergenic) have non-zero transition probabilities to and from each other. IG is the intergenic state. $I_0, \ I_1 \mbox{ and } I_2 \mbox{ are introns of phase 0, 1 and 2, i.e.}$ they interrupt the previous exon's final codon after its 0th, 1st or 2nd base. All shaded nodes are exons: the subscript indicates the initial phase (0, 1, 2, 'I' for initial exon) and the trailing digit indicates the terminal phase (0, 1, 2, 'T' for terminal exon). Thus, E₁₂ is an exon that begins in phase 1 (it finishes a codon whose first base occurs in the previous exon) and ends in phase 2 (its last codon is interrupted after two bases). Separating the functional notions of 'exon' and 'intron' into these distinct states allows us to enforce phase and frame consistencies. For example, a phase-1 intron has positive transition probability only to exons of initial phase 1, and a gene is completed only by a terminal ('T') exon, ensuring it contains a whole number of codons. The entire structure above the IG node is duplicated in the GHMP, to allow annotation on both the forward and reverse strands.

state nodes proceeds from left to right; for concreteness, the three nodes of the chain depicted here have all taken on the intron state. Below each hidden state node appears the phylogenetic tree of Figure 1B (for clarity, only the tree associated with the middle node is depicted). The same tree topology and parameters are used in every instance of the tree.

Observe that every node in the tree has the functional state node as a parent. This allows the nucleotide substitution models to depend on the functional role of the slice being generated. In particular, the evolutionary substitution



Fig. 3. An excerpt of the GHMP graphical model corresponding to an aligned intronic column. The hidden chain of functional states runs along the top. The phylogenetic tree structure culminates in an observed alignment column, which populates its leaves. Every nucleotide distribution in the tree depends on the rate parameter given by the functional state (here, 'intron'); thus, there is an edge from the functional state node to every tree node. There are similar tree structures beneath the other two state nodes, omitted here for visual clarity.

rate from ancestor to descendant varies with function. The version of the GHMP we implemented uses two substitution rates, functional (exons, exon boundaries) and non-functional (introns, intergenic regions). Since the substitution model we consider allows us to establish evolutionary rates only up to a positive scaling factor, we take the functional rate r_{slow} to be 1.0 with no loss of generality, leaving the non-functional rate as a free parameter $r_{fast} > 1.0$.

Formally, let \mathcal{I} be the set of intronic and intergenic states (a subset of the overall hidden state space, namely the unshaded states in Fig. 2). When $z_k \in \mathcal{I}$, the slice length d_k is deterministically one, and $p(s_k|z_k, d_k)$ is the marginal probability of the leaves in a phylogenetic tree with leaf configuration given by the single-column slice s_k . We denote the nucleotides in this column by s_k^1, \ldots, s_k^M and the unobserved ancestor nucleotides by a_k^1, \ldots, a_k^{M-1} , where M is the number of aligned species. The rate parameter used for the tree is r_{fast} . Here, 'marginal' means the ancestor nodes are integrated out of the phylogenetic distribution:

$$p(s_k | z_k \in \mathcal{I}, d_k = 1) = \sum_{a_k^1} \cdots \sum_{a_k^{M-1}} p_{\text{tree}}(a_k^1, \dots, a_k^{M-1}, s_k^1, \dots, s_k^M | r_{\text{fast}}),$$



Fig. 4. An excerpt of the GHMP graphical model corresponding to an aligned internal exon on the forward strand. The hidden chain of functional states runs along the top. The bounding box (plate) around the phylogenetic tree denotes duplication, D_k times. Each copy of the tree corresponds to an alignment column, which populates the tree's leaves. D_k too is random, allowing the length of aligned exons to follow an arbitrary distribution (thus GHMP). The ovals labeled as splice sites are not part of the language of graphical models. They appear here to reduce visual clutter (Fig. 5).

with p_{tree} obtained from the conditional substitution probabilities at each branch or leaf node v^j given its parents $par(v^j)$:

$$p_{\text{tree}}(v^1, \dots, v^{2M-1}|r) = \prod_j p[v^j | \text{par}(v^j), r].$$

While intronic states generate only a single-column slice, implying a geometric distribution for the lengths of aligned introns (Kulp et al., 1996; Burge and Karlin, 1997), exonic states are associated with multiple-column slices. Consider the GHMP fragment shown in Figure 4, where the middle node has taken on the state of a shared exon. This hidden exon state is associated with a left exon boundary slice (here, a 3' splice site), an internal exonic slice and a right exon boundary slice (here, a 5' splice site). The square containing the phylogenetic tree is a piece of graphical model notation called a plate. The plate indicates that the entire tree structure inside the plate is repeated D_k times, corresponding to an aligned exon spanning a D_k -column slice. Of course, different exons must be allowed different lengths, so D_k itself is a random variable, making the overall structure a generalized plate (i.e. a GHMP). The conditional distribution $p(d_k|z_k)$ of D_k given an exon type is arbitrary, so that aligned exon length distributions appropriate to the species at hand may be modeled (Kulp *et al.*, 1996; Burge and Karlin, 1997). (Note that some authors refer to generalized HMMs as hidden semi-Markov models.) Each tree in the figure, including those of the boundary slices we now describe, evolves at the functional rate $r_{slow} = 1.0$.

Formally, for $z_k \in \mathcal{E}$, the exonic hidden states (shaded in Fig. 2), let $\mathbf{s}_k^j = (s_k^{j,1}, \ldots, s_k^{j,M})$ be the observed nucleotides in column j of the slice, $j = 1, \ldots, d_k$. Similarly, write $\mathbf{a}_k^j = \begin{bmatrix} a_k^{j,1}, \ldots, a_k^{j,(M-1)} \end{bmatrix}$ for the ancestor nucleotides in column j's tree. Then,

$$p(s_k | z_k \in \mathcal{E}, d_k) = q_{\text{left}}(s_k, z_k)$$

$$\times \left\{ \prod_{j=1}^{d_k} \sum_{a_k^{j,1}} \cdots \sum_{a_k^{j,(M-1)}} p_{\text{tree}} \left(\mathbf{a}_k^j, \mathbf{s}_k^j | r_{\text{slow}} \right) \right\}$$

$$\times q_{\text{right}}(s_k, z_k)$$

Here, q_{left} and q_{right} assign probabilities for boundaries appropriate to the type of exon given by z_k .

Note that Figure 4 depicts a shared forward-strand internal exon. Other possibilities are an initial or terminal exon in



Fig. 5. Detail of the 5' donor splice site submodel omitted from Figure 4. The bounding box (plate) denotes repetition of the tree structure it contains; in this case there are nine copies ($r = -3, -2, -1, 1, \dots, 6$). The *r*-th tree is matched to the *r*-th column in a window surrounding a shared 5' splice signal (with shared GT in columns 1, 2). The equilibrium nucleotide distribution depends on window position, as indicated by the node labeled π_r . The dot inside the node identifies it as a parameter rather than a random variable.

a multi-exon gene, a single-exon gene or any of these on the reverse strand. The splice-site ovals of Figure 4 are not part of the graphical model nomenclature; we utilize them only to simplify the diagram. We now consider the exon boundary models, i.e. splice site, start codon and stop codon slices. which are substituted in the place of these ovals. In particular, Figure 5 shows the donor splice site model the reader should envision in place of the right oval in Figure 4. It is a plate denoting nine independent copies of the phylogenetic tree, numbered r = -3 to 6, with no zero index. Each copy generates one column of the full donor slice, i.e. the window in the alignment surrounding the shared GT signal at columns 1 and 2. The columns are not identically distributed, but rather have position-dependent equilibrium base distributions π_r . This is explicitly depicted in the plate via the node containing an internal dot. (The dot indicates that the π_r are to be construed as fixed parameters, rather than in a Bayesian fashion as random variables.) The position-specific distributions allow us to exploit varying nucleotide usages in the splice signal's flanking region. This phenomenon has been studied in human genomic sequence by Zhang (1998) and others.

Treatment of the 3' acceptor site's window is analogous to the case of the donor slice. Since the start codon ATG is nonstochastic, it requires no model. Finally, a stop codon slice is generated using a phylogenetic tree of stop codons (not pictured), as follows. A progenitor stop codon TAA, TAG or TGA is chosen at the root of the tree according to a stop codon equilibrium distribution. This codon is evolved towards the leaves; they then constitute a three-column slice of the multiple alignment (each row of which is some valid stop codon). The stop-codon substitution model is defined by first independently evolving each of an ancestor's 3 nt using a nucleotide substitution model. This evolution is then normalized by removing outcomes that are not stop codons and scaling the remaining outcomes by their total probability mass. In this manner, with probability one, a valid stop codon is produced at every node in the tree.

Note the simplicity of the exon model employed. Conditional on being in an exonic hidden state, the columns of the aligned exon interior are independent and identically distributed. The concepts of codon and peptide are not incorporated, nor is sequential dependence along the exon a part of the setup. This is in some sense the most naïve possible model of a shared exon: only a non-geometric length distribution and lower substitution rate, together with the boundary structures, distinguish exons from intronic and intergenic regions of the multiple alignment. As a reflection of what is known biologically about the exon structure, this exon representation is subordinate to the more sophisticated methods of Kulp et al. (1996); Burge and Karlin (1997); Korf et al. (2001); Alexandersson et al. (2003) and others. On the other hand, our approach has scientific virtue: by comparing the performance of a simple model for multiple closely related species with a complex model for a single organism, or distant paired organisms, we learn something about the relative advantages of these gene-finding strategies. In this regard, consult the Results section.

Our discussion of the GHMP model closes with the issue of gaps in the multiple alignment. We have not attempted to include nucleotide insertion and deletion events in our model. Other authors (Pedersen and Hein, 2003; Siepel and Haussler, 2003) treat gaps as missing data, marginalizing gapped leaves out of aligned columns. This approach can be accommodated readily within the probabilistic inference mechanism of the GHMP, but it has practical drawbacks. A gap is not a nucleotide we failed to observe; instead, it is more like a nucleotide that evolved out of the phylogenetic tree at a given homologous position. As such, for purposes of functionally annotating the alignment, it evidences lack of conservation and should not just be integrated away during the probability computations. To incorporate this consideration into the model, we replace all gaps in an aligned column with that column's least-occurring base,² as a heuristic penalization. However, before this is done, deterministic constraints involving gaps are enforced (see below). Note that, due to our use of closely related species, the importance of any particular gap heuristic is greatly diminished: e.g. the aligned exons in our dataset were entirely gapless, so any approach that preferentially assigns gaps outside exons is likely to perform comparably.

²Ties are broken according to the equilibrium base distribution.

Estimation and inference

We now discuss the parameter estimation methods used in our implementation. Starting with raw homologous sequence data from multiple organisms, we first obtain a multiple alignment and phylogenetic tree by repeated alternation between tree-based alignment and maximum-likelihood tree estimation over the aligned sequences. We use MAVID (Bray and Pachter, 2003) for the former and FASTDNAML (Olsen et al., 1994; Felsenstein, 1981) for the latter. The corresponding nucleotide substitution model is described in Felsenstein and Churchill (1996), with equilibrium base frequencies estimated by maximum likelihood from the raw sequence data. The transition-transversion ratio is fixed at 2.0, except where this is incompatible with the estimated equilibrium base distribution, in which case the smallest admissible value is utilized. Once the alignment and tree have been estimated, they are fixed during all subsequent inference on the GHMP, and the same tree topology and branch lengths are used for every column of the alignment.

The hidden Markov chain of functional states requires an initial probability distribution $p(z_1)$ over the functional state space, as well as a matrix of transition probabilities $p(z_k|z_{k-1})$. While these parameters can be estimated using expectation-maximization or other likelihood-based approaches, given appropriate data (Pedersen and Hein, 2003; Siepel and Haussler, 2003), the phylogenetic shadowing principle lets us finesse the issue. Since we work only with immediate primate neighbors of the human, a satisfactory approximation to the model's Markov chain parameters is obtained simply by using widely available maximum-likelihood estimates from annotated human genomic sequences. Indeed, we transferred the reported GC isochore-specific *Homo sapiens* parameters of Alexandersson *et al.* (2003) directly to the GHMP.

The same rationale applies to GC isochore-specific aligned length distributions for exons (by type), introns and intergenic regions, as well as the equilibrium stop codon distribution: previously reported maximum-likelihood estimates on the human genomic sequence are employed. However, since observed intron and intergene lengths in the human sequence do not reflect the increased length in a multiple alignment due to gaps, the geometric distribution mean parameters are scaled up by a factor involving the fraction of gapped columns in the given alignment. This is not necessary for exonic lengths, because of the extreme rarity of gaps, as described previously. Furthermore, the position-dependent equilibrium nucleotide distributions of our donor and acceptor models are fixed at the human occurrence frequencies reported by Zhang (1998). This leaves only one parameter, the non-functional evolutionary substitution rate r. Its treatment as a model selection parameter is discussed in the Results section.

Having described the estimation of all free parameters in the GHMP, we turn now to the inference procedure. First, we enforce a set of deterministic constraints: start codons, stop codons and splice signals must be exactly aligned and gapless. Gaps are allowed only in codon-sized runs within exon slices. Additionally, in-frame stop codons are disallowed for every species inside an exon slice. Taken together, these constraints lead to the identification of all candidate aligned exons. These then underlie a generalized Viterbi algorithm, which computes the most probable trajectory through the hidden functional state chain, conditional on the observed alignment data. This version of the Viterbi algorithm supports non-geometric durations in exonic states, as well as the computation of phylogenetic-tree emission probabilities.

We emphasize again that this algorithm, which involves conditioning on the alignment data and marginalizing out all ancestor branch nodes in the GHMP, is a special case of the general-purpose machinery for graphical model inference. Nonetheless, since we are dealing with generalized exon durations, a naive implementation would have prohibitive complexity on large alignments. As mentioned, we identify candidate shared exons in a preprocessing step, and the Viterbi algorithm contemplates intron-exon or exon-intron transitions only at these candidates. Furthermore, since each exonic state has a unique in-transition and a unique out-transition, both to intronic states (a consequence of generalized exon output distributions), we can avoid maintaining data structures for exon states in the Viterbi recursion, further reducing computational complexity (Burge and Karlin, 1997; Alexandersson et al., 2003).

RESULTS

All-species analysis

We have implemented SHADOWER, a system for automated functional annotation based on the ideas described in the previous section. Here, we report on a re-examination of five exonic regions across a number of primates varying, by region, from 13 to 18. The datasets are described in Boffelli *et al.* (2003). Each region spans roughly 2 kb and contains a single exon from one of the five genes apolipoprotein(a), apolipoprotein(b), cholesteryl ester transfer protein (cetp), liver x-receptor α (lxr α) and plasminogen (plg). Human sequence was used in every region; beyond that, there is modest overlap among the sets of primates sequenced for each dataset.

In Table 1, we show the accuracy of SHADOWER's exon predictions as the non-functional rate r is varied from 1.0 (the functional evolutionary rate) to 2.5. For these datasets, the predicted exon count increases monotonically with r. We estimate performance using cross-validation, leaving out one dataset at a time. At each step, r is chosen on four of the datasets to maximize sensitivity; in the case of multiple maximizing values, the smallest is used. Performance is then assessed on the remaining fifth dataset, as presented in the top row of Table 2. Total nucleotide-level sensitivity and partial-match exon-level sensitivity (which Table 1. Each row shows SHADOWER prediction results on the named single-exon dataset as the non-functional rate parameter r varies from 1.0 to 2.5

	Non-f	unctional	evolution	ary rate (r)											
	1.0	1.1	1.2 ^a	1.3	1.4	1.5	1.6	1.7	1.8	1.9	2.0	2.1	2.2	2.3	2.4	2.5
apo(a)	×	×	\rightarrow	\rightarrow	\rightarrow	\rightarrow	1—	1—	1—	1—	1—	1—	1—	4—	4—	4—
apo(b)	×	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	2√	2√	4√	4√
cetp	1√	1√	1√	1√	$1\checkmark$	$1\checkmark$	1√	$1\checkmark$	1√	$1\checkmark$	$1\checkmark$	1√	$1\checkmark$	$1\checkmark$	2√	2√
$lxr \alpha$	×	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark
plg	—	—	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	\checkmark	1√	1√	1√	1√	1√	1√	1√	3√

Key: \checkmark means the exon is predicted exactly; \times means the exon is completely missed; — means the exon is predicted but both boundaries are incorrect; \rightarrow means the exon is predicted but only the downstream boundary is correct; and the number *n* means there are additionally *n* false positive exons. ^aThe cross-validated choice of *r* turns out to be 1.2 for every dataset.

 Table 2. Sensitivity and specificity of various gene finders on the primate exon datasets

	Nucl. (%	5)	Exon	partial	Exon exact		
	Sn	Sp	Sn	Sp	Sn	Sp	
GENSCAN	44.7	34.0	2/5	2/3	1/5	1/3	
MZEF	37.4	63.2	3/5	3/4	1/5	1/4	
SHADOWER	100.0	89.6	5/5	5/6	4/5	4/6	
SHADOWER ^b	42.7	42.2	2/5	2/5	1/5	1/5	
SLAM	80.2	100.0	3/5	3/3	3/5	3/3	

Results are shown at the nucleotide, partial exon and exact exon levels (see text). GENSCAN predicts complete or incomplete genes, using only the human sequence data. MZEF predicts individual internal exons (without frame or phase consistency), using only the human sequence data. SHADOWER employs the GHMP to analyze multiple orthologous sequences.

SHADOWER^b indicates an analysis that excludes exon boundary models (see text). SLAM uses human-mouse homology in a generalized pair HMM.

forgives inexact boundary demarcations) are both 100% all coding bases in all five exons are detected. Exact exon sensitivity is 4/5, because of a single boundary failure the upstream start codon boundary of the apo(a) exon is incorrectly localized at a nearby splice signal. This is the unique initial exon in our data; we conjecture that this incorrect localization is partly due to the observation that is lacking of the remaining downstream apo(a) exons, whose presence would interact with the hidden Markov dynamics to create a stronger preference for an initial exon at this location.

Turning to false positive exon predictions using the crossvalidated choice of r, we find a specificity of 89.6% at the nucleotide level, 5/6 at the partial-match exon level and 4/6 at the exact exon level. The main failure here is a single false positive terminal exon in the cetp region. It is interesting that this false positive appears at every value of r shown in Table 1, including even the functional rate 1.0. A look at the alignment reveals a highly probable acceptor site slice and stop codon slice flanking an exon of typical length; taken together, these are the likely determinants of this prediction. It is less clear that additional upstream exons would ameliorate this problem, as they would in the case of apo(a). Instead, the situation calls for the enhancement of our site-independent exon slice model. In the Discussion section we expand on this point.

To contrast SHADOWER with state-of-the-art gene-finding methods, we ran GENSCAN, at default settings (Burge and Karlin, 1997), on the human sequence data from each of the five regions (Table 2). Its nucleotide sensitivity came out at 44.7%, versus 100% using SHADOWER, and its nucleotide specificity was 34.0%, versus 89.6% with SHADOWER. GENSCAN entirely missed three of the five exons, partially matched one (lxr α) and demarcated one exactly (cetp), while producing one false positive exon in the apo(b) region.

Given that SHADOWER and GENSCAN both use a functional state space designed primarily to detect complete multi-exon genes, it is unlikely that GENSCAN's poor relative performance is due to an inherent bias toward predicting multiple-exon gene structures. Nonetheless, in order to study the issue of multiple-exon prediction bias, we analyzed the human sequence data using MZEF (Zhang, 1997), a method designed specifically to predict individual internal exons. [Note that for this purpose we construed the one initial exon in our dataset, apo(a), as internal.] Table 2 summarizes the results. As one can see, MZEF is less sensitive (37.4%) but more specific (63.2%) than GENSCAN at the nucleotide level; the same holds at the exact and partial exon levels. It is clear that SHADOWER improves on both MZEF and GENSCAN for these data by exploiting additional constraints from conservation.

We also compared SHADOWER with SLAM (Alexandersson *et al.*, 2003) using human-mouse homology (Table 2). It is known that no homologous mouse sequence exists for the primate-specific apo(a) exon; in addition, we found no cetp homolog. The remaining three exons were demarcated exactly. This gives SLAM a nucleotide sensitivity of 80.2% with 100% specificity. Thus, the results for these three exons are similar to those of SHADOWER, but an important point of this comparison is that the evolutionary distance of human and mouse prevents SLAM from competing on functional annotation of some genomic regions under study.

Finally, it is of interest to determine how SHADOWER performs relative to less sophisticated methods that exploit

multiple-sequence homology. We examined this by analyzing a reduced version of the GHMP with no boundary model components (cf. Fig. 5). As Table 2 shows, the results are poor: nucleotide sensitivity and specificity are both \sim 42%. Only one exon is demarcated exactly. In the absence of boundary probabilities, SHADOWER labels a region as exonic essentially on the basis of a higher likelihood under the conserved phylogenetic model. It would be quite surprising if an algorithm that uses homology alone, ignoring the informative cues of exon boundary structure, did well on test data such as these.

Species-subset analysis and maximal Steiner subtrees

In the context of functional annotation using multi-species sequence datasets, the relationship between the set of species chosen and the consequent annotation quality has enormous practical significance. Given the expense and resources required for large-scale sequencing of an organism, we particularly need to determine how few species suffice to deliver adequate prediction of biological function. Of course, there are many available species sets of a given size, so one arrives naturally at a sequence of optimization problems: for every size, which collection of that size yields the highest-quality functional annotation?

To fix ideas, focus now on the 18 primates sequenced in the apo(a) region. There are 261 972 subsets of these primates having size at least two; we want to avoid producing a SHADOWER analysis for all of them. So, instead, we take the following approach. The results in the previous section show that the total evolutionary divergence among these primates is large enough to distinguish conserved from non-conserved regions but still small enough to enable exact alignments of exon boundaries. Now, we can measure divergence using the maximum-likelihood phylogenetic tree grown on the full apo(a) dataset, construing branch lengths as expected nucleotide substitution counts. In this tree, each available primate belongs to one leaf. From this viewpoint, the total divergence of all the apo(a) primates corresponds to the total weight of the phylogenetic tree, i.e. the sum of all branch lengths.

Similarly, the total divergence of any subset of the apo(a) primates corresponds to the weight of that subset's 'family tree', i.e. the lowest-weight subtree covering all the leaves in the subset (also known as the Steiner subtree for those leaves). This is the tree that a SHADOWER analysis restricted to the given subset would utilize. We take the Steiner subtree weight of an apo(a) primate subset as our surrogate for the annotation 'quality' that subset would provide. Finding the subset of size *k* having the maximal-weight Steiner subtree—the *k*-mss problem—is a well-defined optimization problem that admits a dynamic programming solution linear in tree size. A complete discussion on this topic is in preparation.

In Boffelli *et al.* (2003), it was shown that the percentage of total divergence attained by the k-mss primate subsets in these five regions increases rapidly for k up to five or six, then



Fig. 6. The performance of SHADOWER on various-sized primate collections corresponding to maximal (solid) and minimal (dashed) Steiner subtrees. The *k*-maximal Steiner subtree corresponds to that subset of *k* primates whose total evolutionary divergence is largest among all size-*k* primate subsets at the leaves of the phylogenetic tree. The *k*-minimal Steiner subtree is understood analogously. The minimal Steiner subtree primates yield comparable sensitivity with, but far worse specificity than, the maximal Steiner subtree primates. Also, the performance available using all 13 primates is attained by a subset of just five, provided they are the maximal Steiner subtree primates.

gradually for larger values of k. If our postulated connection between total divergence and annotation quality holds, we expect to see a similar relationship to exon sensitivity and specificity. To study this, we first solved the k-mss problem in all five regions, for each k from 2 to 13. Then, for each k, we ran SHADOWER on the k-mss primate subsets, region by region. The non-functional rate parameter r was chosen as described in the previous subsection.

The resulting nucleotide-level exon sensitivity and specificity are shown as a function of k in Figure 6 (solid lines). As the figure shows, using just the five primates of each region's 5-mss allows SHADOWER to recover the same level of performance it obtained on the full primate collections. As expected, annotation quality improves rapidly at values of k up to five. The results are the same for exact and partial-match exon detection (data not shown). Figure 1A shows the five primates in the apo(a) 5-mss, situated in the phylogenetic tree grown on just their sequence data for the SHADOWER analysis.

In contrast, we repeated the analysis using k-minimal Steiner subtrees, e.g. the k primates that are nearest to one another in the sense of family tree weight. The resulting nucleotide sensitivity and specificity appear in Figure 6 (dashed lines). Note that, although sensitivity is comparable for these

data once $k \ge 3$, specificity is drastically reduced when the *k*-minimal Steiner subtree species are chosen. This accords with intuition, since increased evolutionary distance offers more opportunities to accrue the mutations that reveal false positives. One concludes that proper selection of target primates for sequencing will have a major impact on the future success of comparative functional annotation projects.

DISCUSSION

We have developed the generalized hidden Markov phylogeny, a graphical model architecture that provides a rigorous probabilistic underpinning for the phylogenetic shadowing principle. Empirical results on a small dataset of five orthologous primate exon regions support the premise of phylogenetic shadowing, showing that a straightforward implementation of a GHMP can yield competitive performance in the identification of primate-specific elements in the human genome.

We have described a relatively simple implementation of the GHMP, in which the elementary component models (splice sites, exons, introns, intergenic regions) do not attempt to capture probabilistic dependence among the aligned columns. Our success on the available datasets makes a strong case for the viability of such a simplified model. This success is, of course, predicated on the strength of the signal in the data. For basic *ab initio* annotation of exons, introns and splice sites, our analysis has suggested this signal is sufficiently strong for accurate annotation when the data consist of sequences from as few as five primates.

We showed that making the GHMP simpler by eliminating its splice site models has a strong detrimental effect on its performance. A similar question relates to our use of non-geometric exon length distributions: how important are they? Since human exon lengths are known to be nothing like exponentially distributed (Kulp et al., 1996; Burge and Karlin, 1997), we anticipate that non-generalized hidden state durations would also cause the GHMP's performance to deteriorate dramatically in the setting of larger-scale analyses. Unfortunately, little can be learned from an analysis on five distinct exons, precisely because the effects of the length distribution cannot manifest themselves on a sample of size five. This is in contrast to, say, nucleotide-level sensitivity and specificity analyses: even our limited dataset comprises many thousands of aligned columns in total, which suffices to illuminate the strengths and shortcomings of the approach.

The graphical model framework underlying the GHMP readily accommodates architectural variations and extensions, and several are of immediate interest. First, the GHMP can be extended to allow for the identification of regulatory elements and binding sites. The known regulatory similarities of closely related organisms suggest that such sites may be conserved in position and number; we already have empirical evidence for this from the apo(a) gene (Boffelli *et al.*, 2003).

Second, the GHMP model described here does not explicitly assign probabilities to insertion or deletion events. A model incorporating gapped slices would be of general interest, and in particular would be useful in the context of the regulatory element modeling problem, where, for instance, varying-sized boxes of short repetitive elements are known to be homologous across species. In addition, we anticipate that a small percentage of orthologous genes among closely related primates will exhibit more complicated evolutionary structure, so that their *ab initio* annotation will be possible only with a more realistic probabilistic treatment of insertions and deletions.

Finally, a more powerful codon-based exon model would not only help in the reduction of false positives (as, e.g. in the cetp gene), but could also be used to incorporate functional annotation methods for proteins (e.g. Simon *et al.*, 2002) into genomic sequence annotation.

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