Evolutionarily conserved elements
How to find functional sequences?
Look for sequences that are conserved across species.

Orthologous sequences that are significantly more similar than expected are likely to have critical functional roles.
Based on analyses of human and rodent genomes:
About 5% or more of bases in mammalian genomes are under purifying selection

Protein-coding genes account for only about 1.5% of bases

3.5% conserved (functional) noncoding sequences
Pairwise alignments and simple percent-identity based methods

VISTA, PipMaker, zPicture
‘-’: do not use phylogeny
   use sliding window of fixed size
PhastCons – to identify conserved elements in multiply aligned sequences.

Based on phylogenetic hidden-Markov model and considers
1) the process by which nucleotide substitutions occur at each site in a genome and
2) how this process changes from one site to next

`+` : do not require a sliding window of fixed size;
allow nearly all parameters to be estimated from the data by maximum likelihood;
efficient on large-scale datasets

Siepel et al. 2005

Figure 1. State-transition diagram for the phylo-HMM used by phastCons, which consists of a state for conserved regions (c) and a state for nonconserved regions (n). Each state is associated with a phylogenetic model (ψc and ψn); these models are identical except for a scaling parameter ρ (0 ≤ ρ ≤ 1), which is applied to the branch lengths of ψc and represents the average rate of substitution in conserved regions as a fraction of the average rate in nonconserved regions (see Methods). Two parameters, μ and ν (0 ≤ μ, ν ≤ 1), define all state-transition probabilities, as illustrated. The probability of visiting each state first (indicated by arcs from the node labeled “begin”) is simply set equal to the probability of that state at equilibrium (stationarity). The model can be thought of as a probabilistic machine that “generates” a multiple alignment, consisting of alternating sequences of conserved (dark gray) and nonconserved (light gray) alignment columns (see example at bottom).
Four separate genome-wide multiple alignments (MULTIZ program):

- 4 vertebrates (human, mouse, rat, chicken) – reference genome human
- 4 insects – reference genome *D.melanogaster*
- 2 worms – reference genome *C.elegans*
- 7 yeasts – reference genome *S.cerevisiae*
Predicted elements covered:
4.3% of the human genome
44.5 of the \textit{D.melanogaster} genome
26.4\% of \textit{C.elegans} genome
55.6\% of \textit{S.cerevisiae} genome
Most conserved bases in vertebrates and insects do not code for proteins → the importance of gene regulation in complex eukaryotes.
The lengths of predicted elements:
100-120 bp for vertebrate, insects and yeast groups
270 bp for worm group
(5bp – thousands of bp)

In vertebrates:
Highly conserved elements (HCE)

- Longer than UCEs (ultraconserved elements) (in vertebrates average 780 bp)
- Less extreme sequence conservation than in UCEs (due to the length dependency of log-odds scores)
- Based on different set of species
- Set of vertebrate HCEs is 10-fold larger than the set of UCEs
- Vertebrate HCEs include 80% of human/rodent UCEs
- More strongly associated with genes as genome sizes become smaller and gene densities increase
- 14.3 % top 100 vertebrate HCE’s overlap 3’UTR’s (in all conserved elements 5.6%)
• Significant enrichment for local secondary structures in 3’UTR, 5’ UTR, introns and intergenic regions HCE’s. *Manuscript in preparation (Pedersen, Bejerano and Haussler)*

• In vertebrates, intergenic HCEs are strongly enriched in stable gene deserts, suggesting that many of them may act as distal *cis*-regulatory elements
Highlights of the study:

• Larger genome and more complex organism – more conserved bases outside of known or suspected exons of protein-coding genes

• Some of the most extreme conservation is in 3’UTRs of vertebrates genes which regulate other genes

• HCEs in vertebrate 3’UTRs, introns and intergenic regions are enriched with local RNA secondary structures

• Intergenic HCEs in vertebrates are strongly enriched in stable gene deserts