Regulatory Elements
part of “Genomik der Genregulation”

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TFs are organized into *cis*-regulatory elements

- promoter regions: directly upstream and/or downstream of the transcription start site (TSS)
- proximal promoter: within 500nt upstream of the TSS
- enhancer elements: “anywhere” in the genome
<table>
<thead>
<tr>
<th>RNA polymerase</th>
<th>Promoter Description</th>
<th>Location relative to start site</th>
<th>Transcript</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pol I</td>
<td>core element UCE (upstream control element)</td>
<td>-45 to +20, -180 to -107</td>
<td>pre-rRNA (28S, 18S, 5.8S)</td>
<td>components of the ribosome; translation</td>
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<tr>
<td></td>
<td>TATA-Box Initiator CpG islands, no</td>
<td>-25 to -35, -100</td>
<td>mRNA, snRNA (U1-4), LINEs</td>
<td>protein coding genes; components of the spliceosome; mRNA splicing Retrotransposon</td>
</tr>
<tr>
<td>Pol III</td>
<td>type 1: A-box, C-box, type 2: A-box, B-box, type 3: TATA-Box</td>
<td>+50 to +80, +10 to +60, -30 to -70</td>
<td>5S rRNA, tRNA, snRNA (U6)</td>
<td>component of large ribosomal subunit translation; components of the spliceosome; mRNA splicing component of the SRP (signal recognition particle); protein transport to ER (endoplasmatic reticulum)</td>
</tr>
</tbody>
</table>
Core motifs of the different promoter types. Motifs in dark gray are less dispensable than motifs in light gray. Any specific promoter may contain just a subset or, in the worst case, none of these motifs. UCE = upstream control element, BRE = TFIIB recognition element, Inr = initiator element, DPE = downstream core promoter element, Oct = octamer binding site, PSE = proximal sequence element. The arrow indicates the transcription start site at +1.
Regulatory Elements
cis-regulatory sequences - enhancer elements

- carry out a regulatory function that is a subfunction of a complex regulatory pattern
- execute this subfunction when linked to a reporter gene
- 100-300bp long
- may be separated from the target gene by several 100kb
- contain multiple sequence-specific binding motifs for transcription factors
  - short
  - gapless
  - more or less conserved
computational approaches

- local (window of 100-200nt) overrepresentation of a particular TFBS (more observed BS than expected given a background model)
- mapping of genome-wide ChIP-seq data for the TF
- phylogenetic footprinting (for distant species)
- phylogenetic shadowing (for closely related species)
Phylogenetic Footprinting

- functional DNA evolves much slower than non-functional DNA
- sequences of divergent species show some conservation
- conserved regions are thought to be functional elements
- density and length of conserved regions decrease as evolutionary distances increase
Phylogenetic Footprinting

divergent sites are indicated by red arrows

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Phylogenetic Shadowing

divergent sites are indicated by red arrows

- sequences of closely related species show some differences
- differences from many sequences taken together reveal variable regions
- variability is assumed to be detrimental
- functional regions are said to lie in less variable regions
sequences of closely related species show some differences

differences from many sequences taken together reveal variable regions

variability is assumed to be detrimental

functional regions are said to lie in less variable regions
nascent transcripts can recruit specific RNA-binding proteins or RNAs to an actively transcribed genomic locus (e.g. TF binds RNA instead of DNA)

non-coding sequences on mRNAs (UTRs) can sequence/structural elements regulating mRNA stability, and translation (e.g. upstream ORFs, IRES, miRNA target sites)

small non-coding RNAs, like miRNAs (microRNAs), can a.o. regulate mRNA degradation or inhibition of translation

long non-coding RNAs can be involved in various regulatory mechanisms
Factor requirement of different viral IRESes. (A) Most viral IRESes including Picornaviruses and Lentiviruses. All canonical initiation factors are required with the exception of eIF4E and the Nt of eIF4G. (B) Flaviviral/pestiviruses IRES. Ribosome entry does not require eIFs 4F/4A/4B/1/1A, but need eIF5, eIF2 and eIF3. (C) Dicistrovirus IGR IRESes. This 200 nt long IRES is able to contact directly a 40S subunit and to assemble an 80S elongation competent ribosome without any of the initiation factors.
upstream ORFs
micro RNAs in left and right asymmetric neural fate decision in *C. elegans*
the ncRNA is necessary to hold proteins of a protein complex together
- the ncRNA specifies the set of proteins forming the protein complex
the ncRNA links the protein complex to the genomic locus
ncRNAs are not important only the act of transcription is
ncRNA hotair lies between hoxC and hoxC on the opposite strand

- it regulates expression of hoxD genes in *trans*
- it bind polycomb repressive complex 2 (PRC2) with its 5’-end which tri-methylates H3K27 (effect: repressive)
- it binds LSD1 with its 3’-end which demethylate mono- and di-methylated H3K4 and H3K9 (effect: repressive)