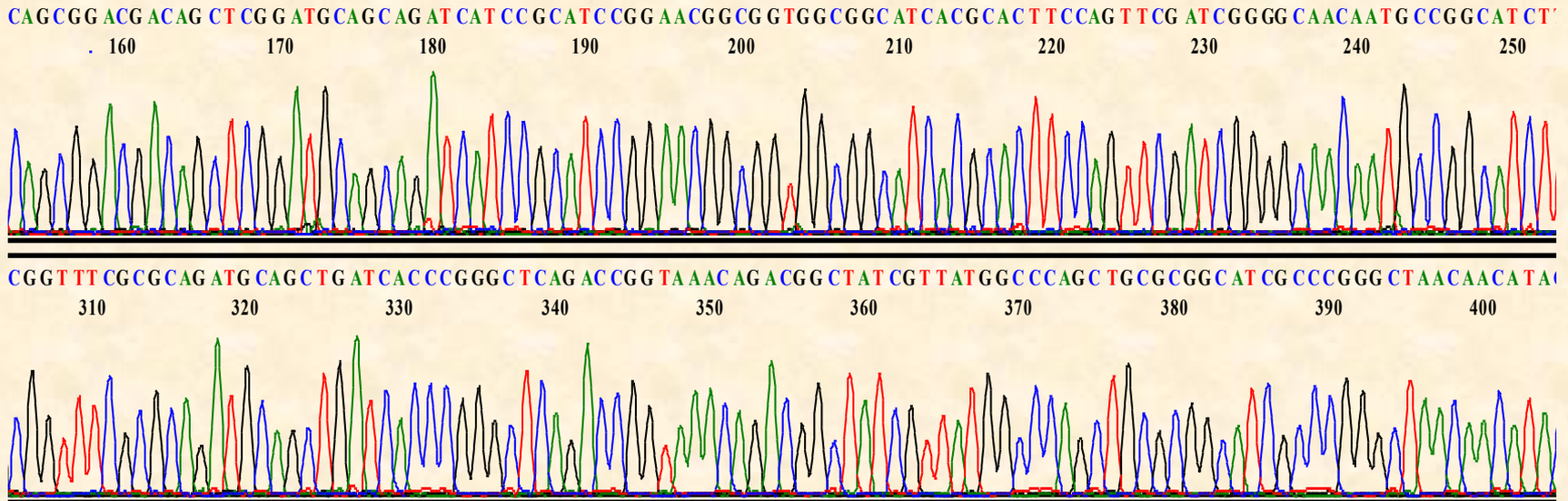


Predikce genů



GATAGCGTAATGATCGGCTGGCTGCCGATTTTCATGCTGGTTTCCCAACGAAAATAACCGCTCACGGTGCCATCACGATCGCACACCGCAAATCGGCGG
TACAGGTGGTCGCGCCCGCCGACACATCGCTGCGCCAATAATGATCTTTCAGCGGACGACAGCTCGGATGCAGCAGATCATCCGCATCCGGAACGGC
GGTGGCGGCATCACGCACCTCCAGTTCGATCGGGGCAACAATGCCGGCATCTTTCAGGGCAAAGCGAATAAACAGCACGCTCACCTCCGCGGCAGCGCC
AGCGCGGTTTCGCGCAGATGCAGCTGATCACCCGGGCTCAGACCGGTAAACAGACGGCTATCGTTATGGCCAGCTGCGCGGCATCGCCCGGGCTAAACA
CATACAGGTGGCGACCATCAATCACGGTCGGGGCGCCGGATCACGGCTGGCTTCCGGATAGGCGCTCAGCAGGGTAACGGCATCCACAATCACCAGCAT

GATAGCGTAATGATCGGCTGGCTGCCGATTTTCATGCTGGTTTCCCAACGAAAATAACCGCTCACGGTGCCATCACGATCGCACACCCGAAAAATCGGCGG
TACAGGTGGTCGCGCCCGCCGCCAGCACATCGCTGCGCCAATAATGATCTTTTCAGCGGACGACAGCTCGGATGCAGCAGATCATCCGCATCCGGAACGGC
GGTGGCGGCATCACGCAC TTCCAGTTCGATCGGGGCAACAATGCCGGCATCTTTTCAGGGCAAAGCGAATAAACAGCACGCTCACTTCGCGCGCAGCGCC
AGCGCGGTTTCGCGCAGATGCAGCTGATCACCCGGGCTCAGACC GGTAACAGACGGCTATCGTTATGGCCCAGCTGCGCGGCATCGCCCGGGCTAACAA
CATACAGGTGGCGACCATCAATCACGGTCGGGGCGGCCGGATCACGGCTGGCTTCCGGATAGGCGCTCAGCAGGGTAACGGCATCCACAA TCACCAGCAT

„Syrové“ sekvence DNA

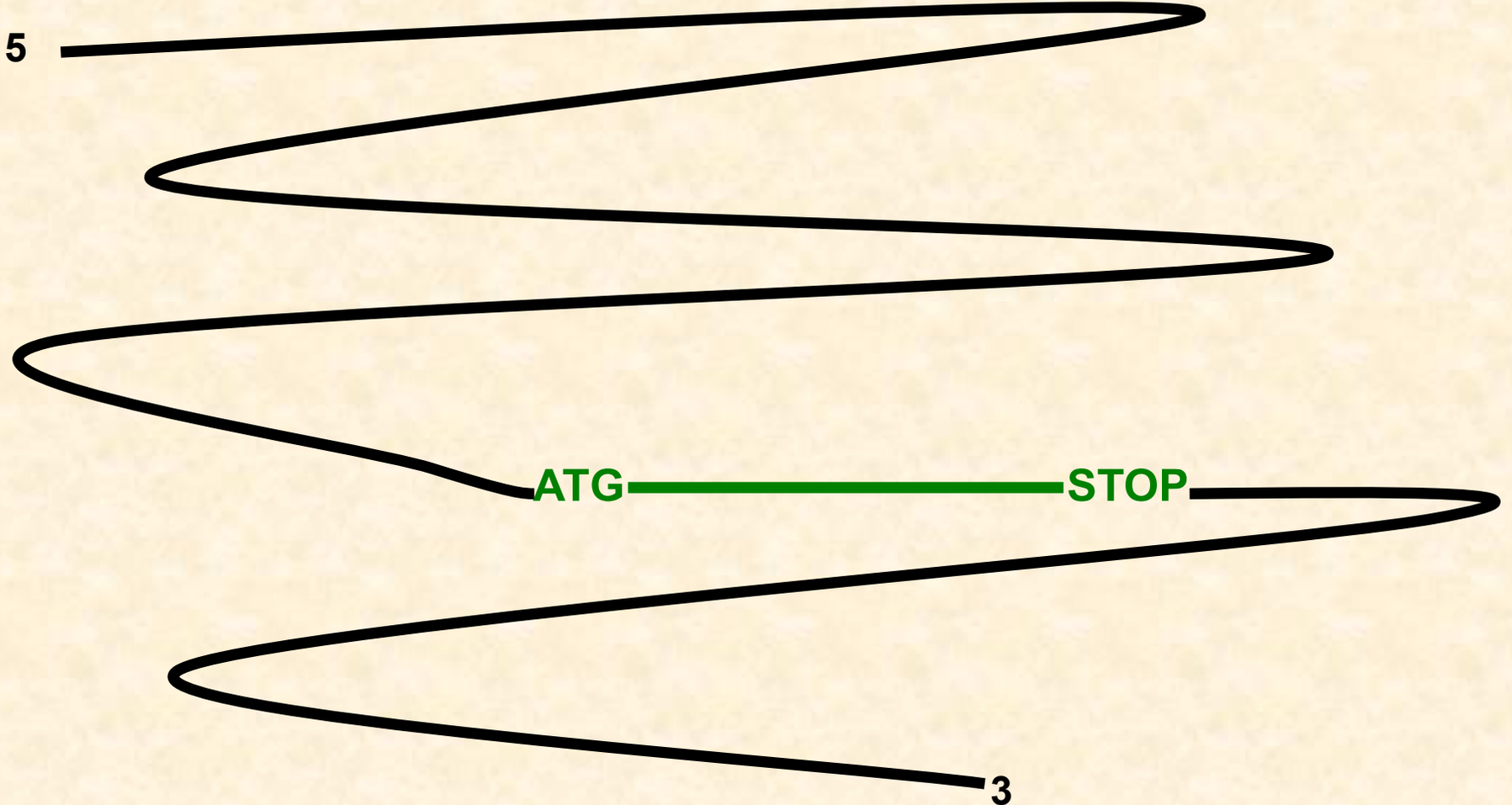


Identifikace a anotace genů a proteinů

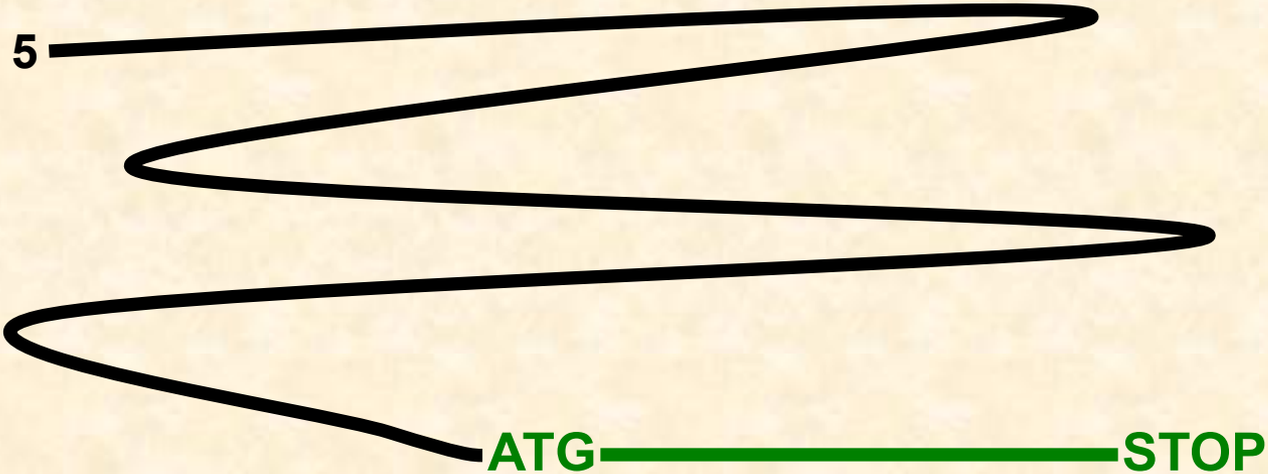
Table 1
Software commonly used for bacterial genome annotation and comparison

<i>DNA level annotation</i>		
GeneMark	http://exon.gatech.edu/genemark/	Protein gene prediction
Glimmer	http://www.genomics.jhu.edu/Glimmer/	Protein gene prediction
SHOW	http://genome.jouy.inra.fr/ssb/SHOW/	Protein gene prediction
tRNAscan-SE	http://lowelab.ucsc.edu/tRNAscan-SE/	tRNA gene prediction
RNAmmer	http://www.cbs.dtu.dk/services/RNAmmer/	rRNA gene prediction
RepSeek	http://www.abi.snv.jussieu.fr/%98public/RepSeek/	Search for approximate repeats in complete DNA sequences
IslandPath	http://www.pathogenomics.sfu.ca/islandpath/	Identification of genomic islands
<i>Protein level annotation</i>		
BLAST	http://www.ncbi.nlm.nih.gov/BLAST/	Compare a novel sequence with those contained in nucleotide and protein databases
InterProScan	http://www.ebi.ac.uk/InterProScan/	Search for domains/motifs in the InterPro database
COGNITOR	http://www.ncbi.nlm.nih.gov/COG/old/xognitor.html	Compare a query sequence to the COG (Cluster of Orthologous Groups of proteins) database
PRIAM	http://bioinfo.genopole-toulouse.prd.fr/priam/	Detection of enzymatic function in a fully sequenced genome, based on all sequences available in the ENZYME database
GOAnno	http://bips.u-strasbg.fr/GOAnno/	BLAST search on the Gene Ontology database
PSORTb	http://www.psort.org/psortb/	Prediction of bacterial protein subcellular localization
TMHMM	http://www.cbs.dtu.dk/services/TMHMM/	Prediction of transmembrane helices in protein sequences
SignalP	http://www.cbs.dtu.dk/services/SignalP/	Prediction of signal peptide cleavage sites in protein sequences
<i>Comparative genomic tools</i>		
Mauve	http://gel.ahabs.wisc.edu/mauve/	Multiple genome alignments in the presence of large-scale evolutionary events
MOSAIC	http://mig.jouy.inra.fr/mig/mig_eng/presentation/project/mosaic	Define the set of backbones and loops in closely related bacterial genomes
ACT	http://www.sanger.ac.uk/Software/ACT/	Comparative genome analysis and visualization tools for multiple genome alignments
CGAT	http://mbgd.genome.ad.jp/CGAT/	
MaGe	http://www.genoscope.cns.fr/agc/mage/	Computation of gene order conservation (syntenies) between available bacterial genomes
Pathologic	http://biocyc.org/	Metabolic network reconstruction and comparative pathway analysis
PUMA2	http://compbio.mcs.anl.gov/puma2/	Metabolic pathway reconstruction
The SEED	http://theseed.ucchicago.edu/FIG/	Comparative analysis and annotation tools using the subsystem approach
STRING	http://string.embl.de/	Search Tool for the Retrieval of Interacting Proteins
PyPhy	http://www.cbs.dtu.dk/staff/thomas/pyphy/	Reconstruction of phylogenetic relationships of complete microbial genomes
HoSeqI	http://pbil.univ-lyon1.fr/software/HoSeqI/	Automatically assign sequences to homologous gene families from the HOGENOM database

Gen



Gen



DNA



RNA

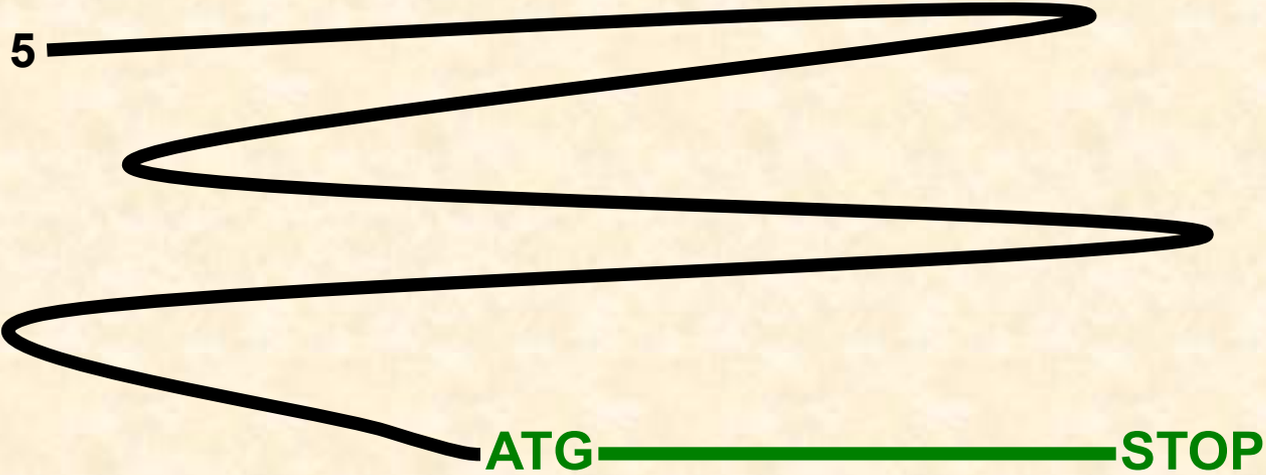


Protein



Met

Gen



DNA



RNA



Protein



Met



Predikce genů kódujících proteiny

- **Prokaryotické geny**

Nepřerušované úseky DNA mezi **startovním kodonem** (ATG, GTG, TTG, CTG) a **stop kodonem** (TAA, TGA, TAG).

- **Úsek kóduje protein, který je podobný již dříve popsanému proteinu (prohledávání DATABÁZÍ pomocí ALIGNMENTU).**

Překlad DNA sekvence

- **ExPASy**

<http://www.expasy.org/tools/dna.html>

- **ORF Finder (NCBI)**

<http://www.ncbi.nlm.nih.gov/gorf/gorf.html>

nonpolar polar basic acidic (stop codon)

Překlad DNA sekvence

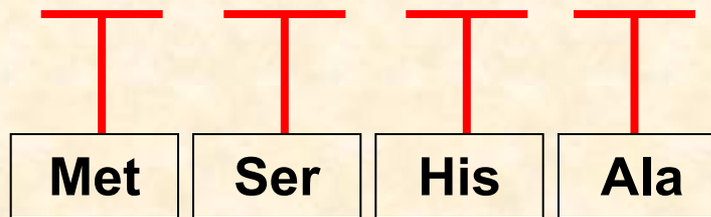
The table shows the 64 codons and the amino acid for each. The **direction** of the mRNA is 5' to 3'.

		2nd base			
		U	C	A	G
1st base	U	UUU (Phe/F) Phenylalanine	UCU (Ser/S) Serine	UAU (Tyr/Y) Tyrosine	UGU (Cys/C) Cysteine
		UUC (Phe/F) Phenylalanine	UCC (Ser/S) Serine	UAC (Tyr/Y) Tyrosine	UGC (Cys/C) Cysteine
		UUA (Leu/L) Leucine	UCA (Ser/S) Serine	UAA Ochre (Stop)	UGA Opal (Stop)
		UUG (Leu/L) Leucine	UCG (Ser/S) Serine	UAG Amber (Stop)	UGG (Trp/W) Tryptophan
	C	CUU (Leu/L) Leucine	CCU (Pro/P) Proline	CAU (His/H) Histidine	CGU (Arg/R) Arginine
		CUC (Leu/L) Leucine	CCC (Pro/P) Proline	CAC (His/H) Histidine	CGC (Arg/R) Arginine
		CUA (Leu/L) Leucine	CCA (Pro/P) Proline	CAA (Gln/Q) Glutamine	CGA (Arg/R) Arginine
		CUG (Leu/L) Leucine	CCG (Pro/P) Proline	CAG (Gln/Q) Glutamine	CGG (Arg/R) Arginine
	A	AUU (Ile/I) Isoleucine	ACU (Thr/T) Threonine	AAU (Asn/N) Asparagine	AGU (Ser/S) Serine
		AUC (Ile/I) Isoleucine	ACC (Thr/T) Threonine	AAC (Asn/N) Asparagine	AGC (Ser/S) Serine
		AUA (Ile/I) Isoleucine	ACA (Thr/T) Threonine	AAA (Lys/K) Lysine	AGA (Arg/R) Arginine
		AUG (Met/M) Methionine, Start ^[A]	ACG (Thr/T) Threonine	AAG (Lys/K) Lysine	AGG (Arg/R) Arginine
	G	GUU (Val/V) Valine	GCU (Ala/A) Alanine	GAU (Asp/D) Aspartic acid	GGU (Gly/G) Glycine
		GUC (Val/V) Valine	GCC (Ala/A) Alanine	GAC (Asp/D) Aspartic acid	GGC (Gly/G) Glycine
		GUA (Val/V) Valine	GCA (Ala/A) Alanine	GAA (Glu/E) Glutamic acid	GGA (Gly/G) Glycine
		GUG (Val/V) Valine	GCG (Ala/A) Alanine	GAG (Glu/E) Glutamic acid	GGG (Gly/G) Glycine

Překlad DNA sekvence

ATG ————— STOP

ATG TCG CAT GCC STOP



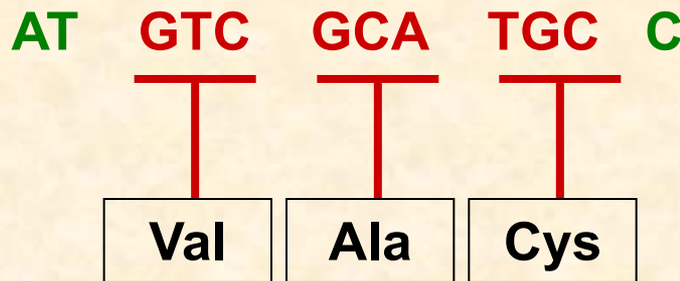
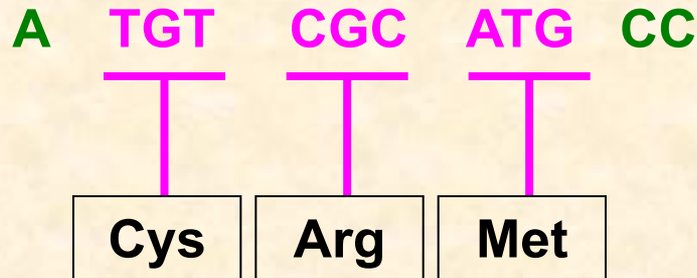
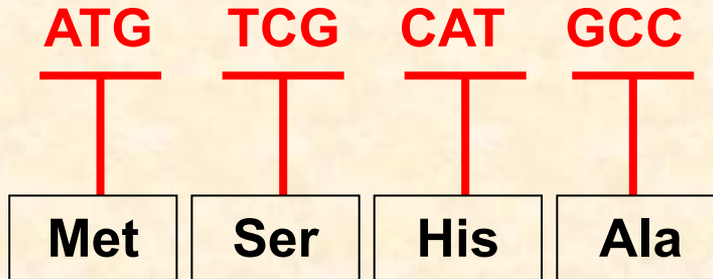
N-konec

Met —————

C-konec

NH₂ ————— COOH

↓ ↓ ↓
ATGTCGCATGCC



Čtení tripletů závisí na tom, u kterého nukleotidu stanovíme počátek čtení.

ATGCGCAGGAATGCATAG sekvence DNA

Met	His	?
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Protein 2

ATGCGCAGGAATGCATAG

Met	Arg	Arg	Asn	Ala	Stop
-----	-----	-----	-----	-----	------

Protein 1

Opravdu ORF kóduje protein?

- ORF kóduje protein, který je podobný již dříve popsanému proteinu (prohledávání DATABÁZÍ pomocí ALIGNMENTU) = **nejspolehlivější ověření.**
- **Nástroje pro překlad DNA jsou propojeny s prohledáváním databází.**