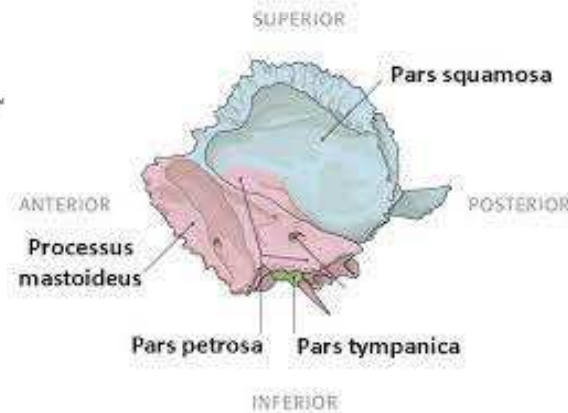
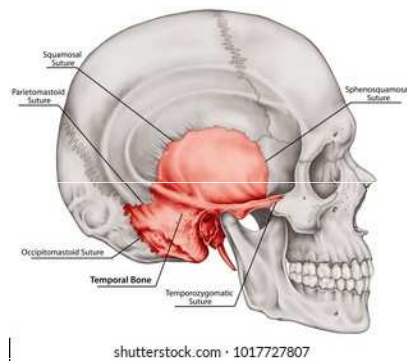


ARCHAICKÁ DNA

Paleogenetika = studium vymřelých organismů prostřednictvím archaické DNA (aDNA, *ancient DNA*)

zdroje aDNA: zuby, kosti (kost skalní). muzejní exponáty atd.
ale také např. vaječné skořápky, koprolity, rozžvýkaná březová kůra,
půda, proteinové obaly hnid vši šatních



další zdroje: keratin, chitin, osteokalcin, kolagen, hemoglobin, pigmenty,
rostlinné polysacharidy → proteomické metody

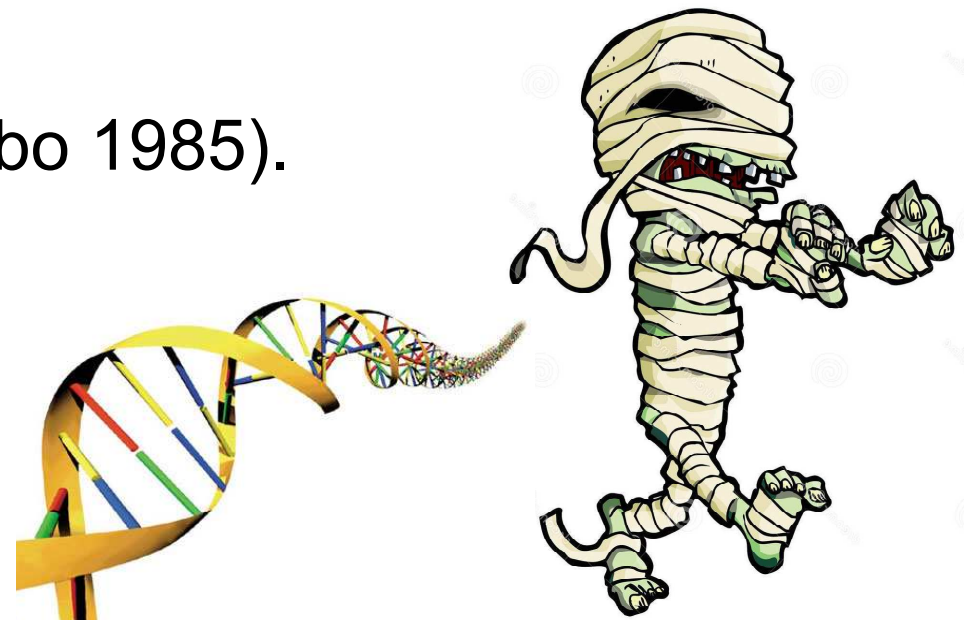
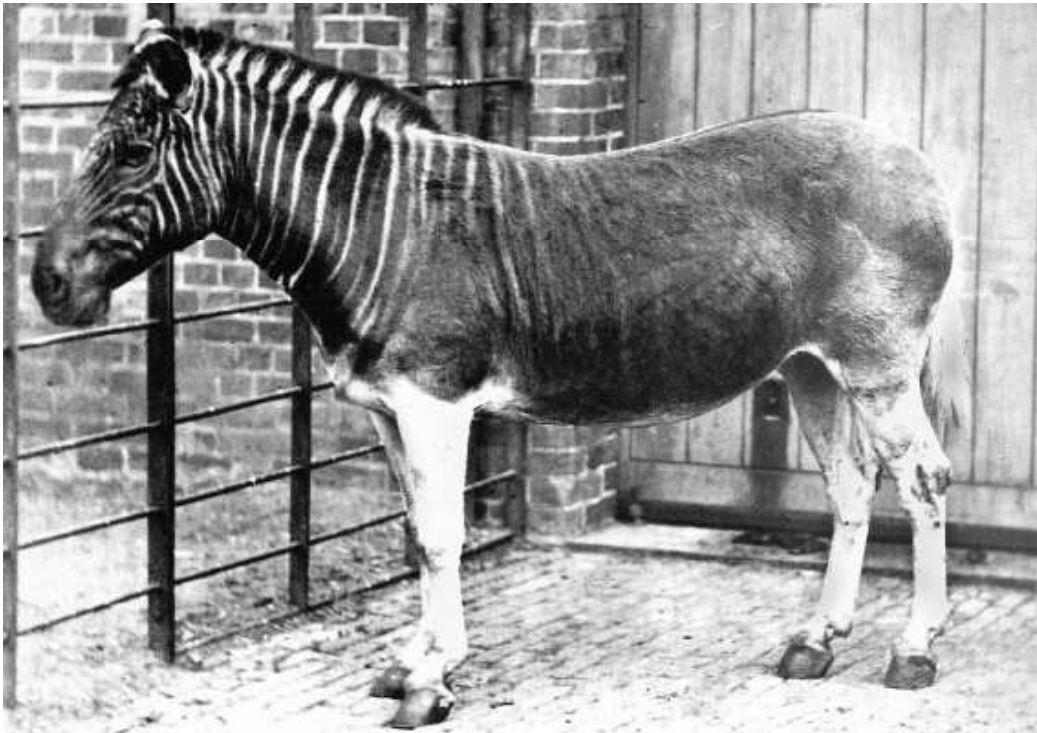
= paleoproteomika

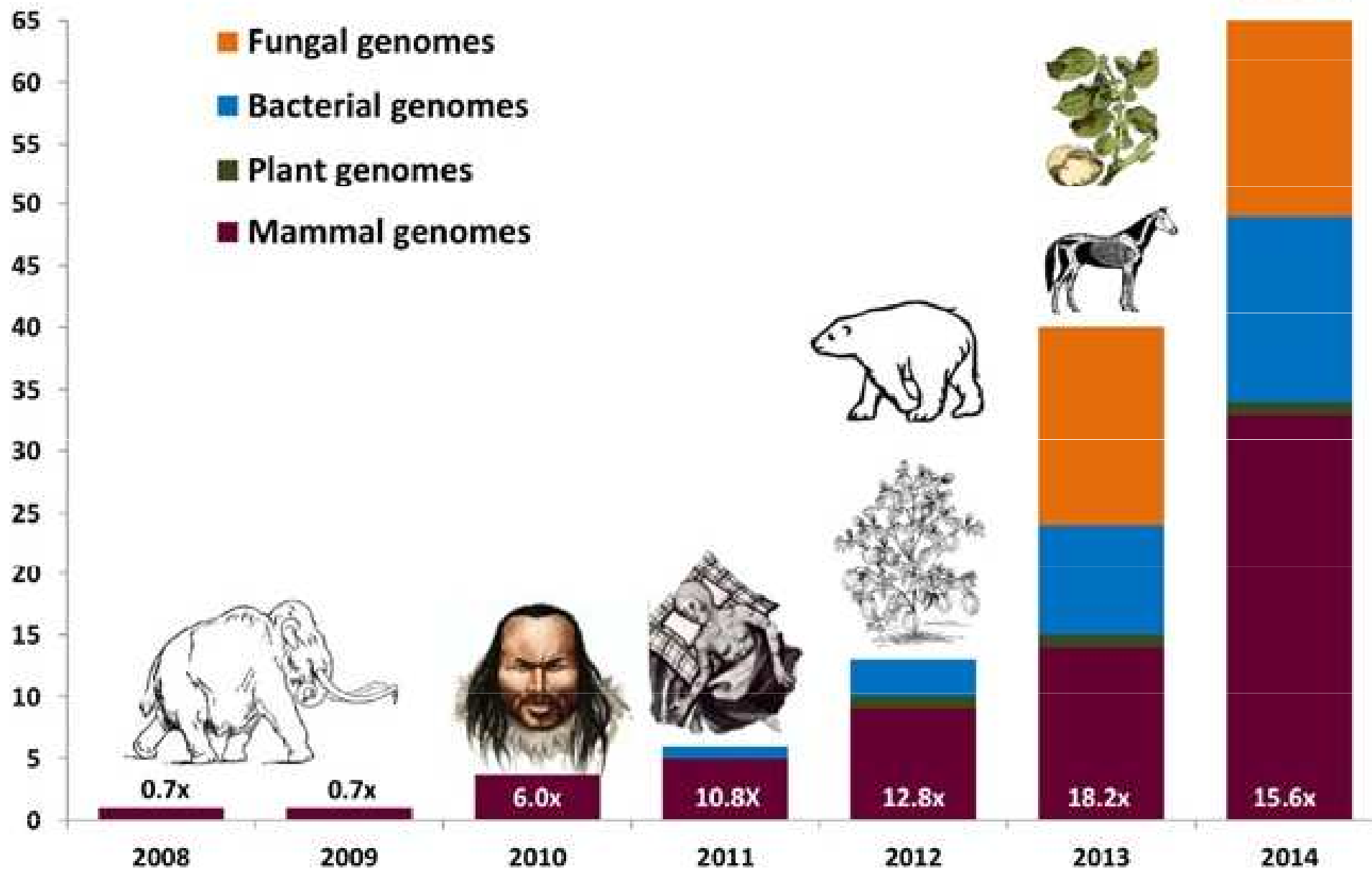
Výhoda: nedochází k fragmentaci proteinů ⇒ větší časové škály

Nevýhoda: proteiny neposkytují tolik informací jako DNA

První sekvence aDNA: zebra kvaga (*Equus quagga*)
150 let (Higuchi et al. 1984)

egyptská mumie: 2400 let (Pääbo 1985).

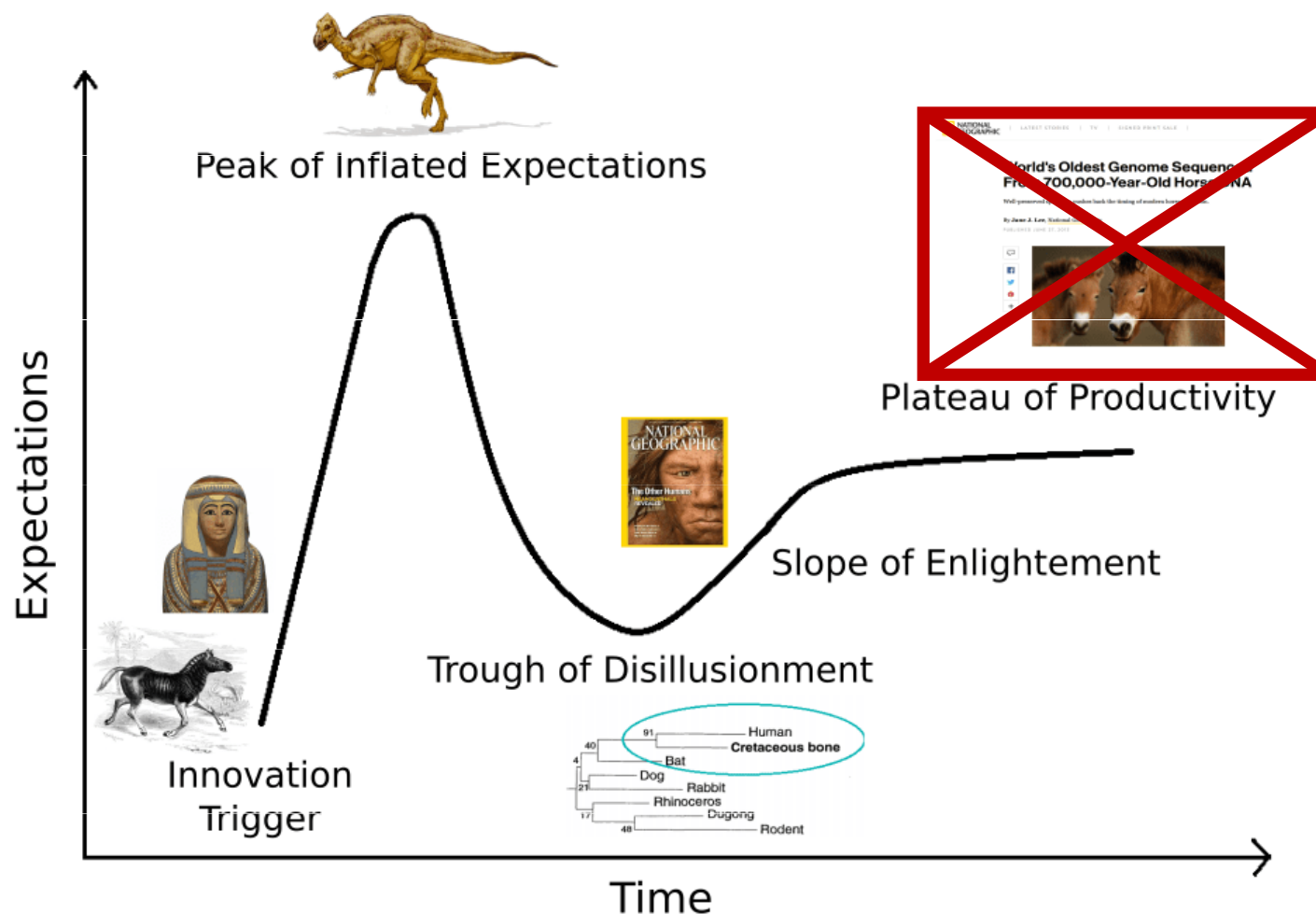




Stáří aDNA:

většinou max. tisíce až desítky tisíc let

The Hype Cycle of Ancient DNA



mamut z permafrostu: 1,10 – 1,65 mil. let

proteinové sekvence:

2019: nosorožec, 1,77 mil., Dmanisi

2016: pštros, 3,8 mil., Laetoli

analýza proteinů vzorků až z rané jury



DNA from multiple mammoth species is illuminating a complex evolutionary picture.

MAMMOTH GENOMES SHATTER RECORD FOR OLDEST ANCIENT DNA

Permafrost-preserved teeth, up to 1.65 million years old, identify a new kind of mammoth in Siberia.

Article


Million-year-old DNA sheds light on the genomic history of mammoths

<https://doi.org/10.1038/s41586-021-03224-9>

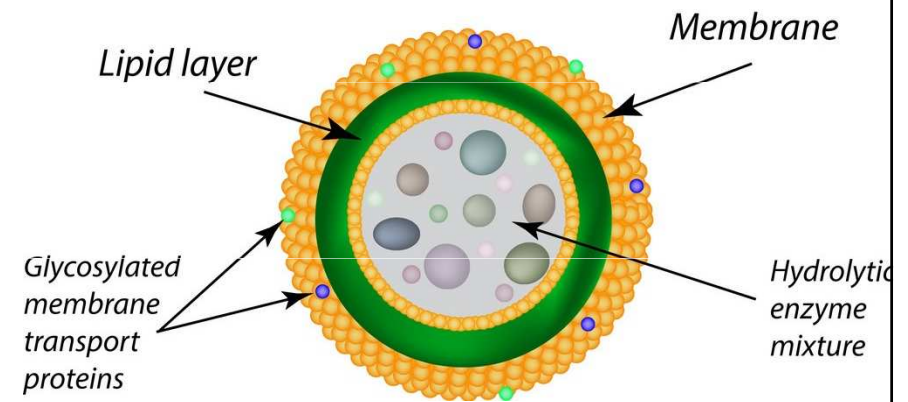
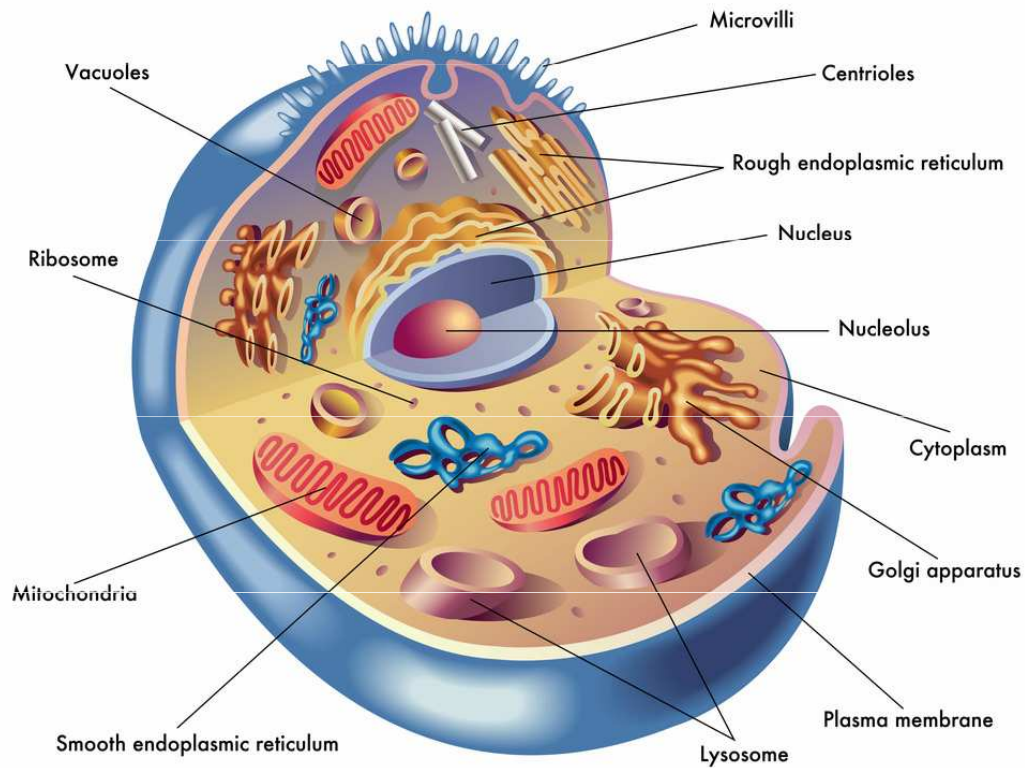
Received: 3 July 2020

Accepted: 11 January 2021

Published online: 17 February 2021

 Check for updates

Tom van der Valk^{1,2,3,17}✉, Patrícia Pečnerová^{2,4,5,17}, David Díez-del-Molino^{1,2,4,17}, Anders Bergström⁶, Jonas Oppenheimer⁷, Stefanie Hartmann⁸, Georgios Xenikoudakis⁹, Jessica A. Thomas⁹, Marianne Dehasque^{1,2,4}, Ekin Sağlıcan⁹, Fatma Rabia Fidan⁹, Ian Barnes¹⁰, Shanlin Liu¹¹, Mehmet Somel⁹, Peter D. Heintzman¹², Pavel Nikolskiy¹³, Beth Shapiro^{14,15}, Pontus Skoglund⁶, Michael Hofreiter⁸, Adrian M. Lister¹⁰, Anders Götherström^{1,16,18} & Love Dalén^{1,2,4,18}✉



Degradace DNA ihned po zániku buňky:

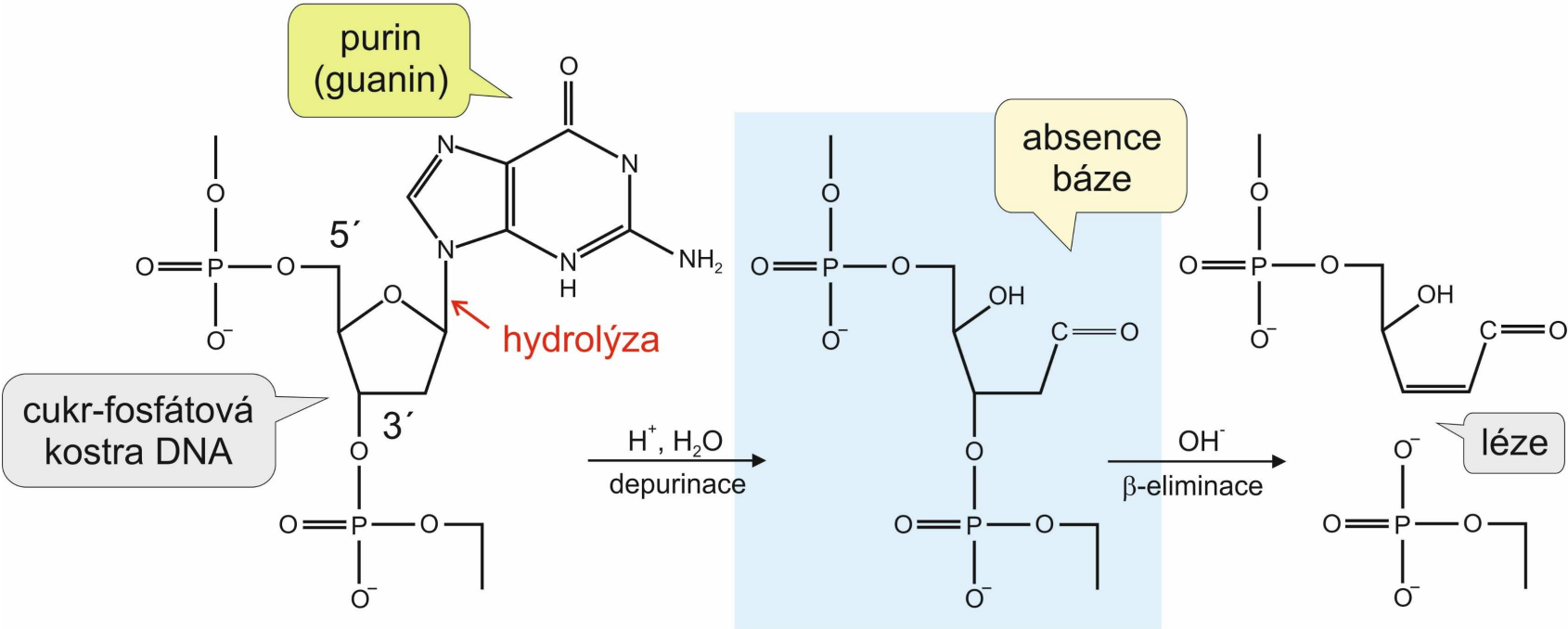
→ fragmentace

→ poškození blokující replikaci DNA

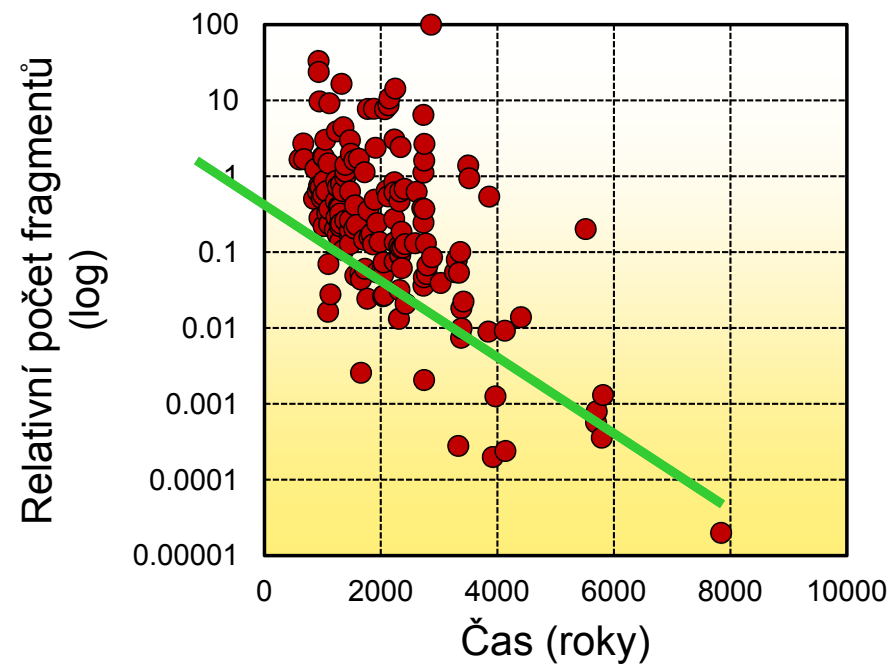
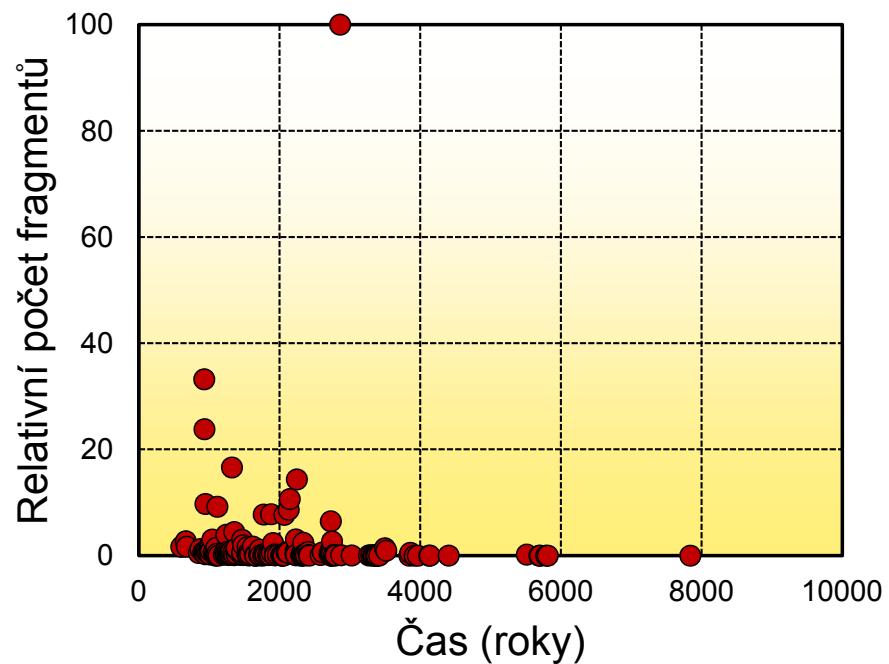
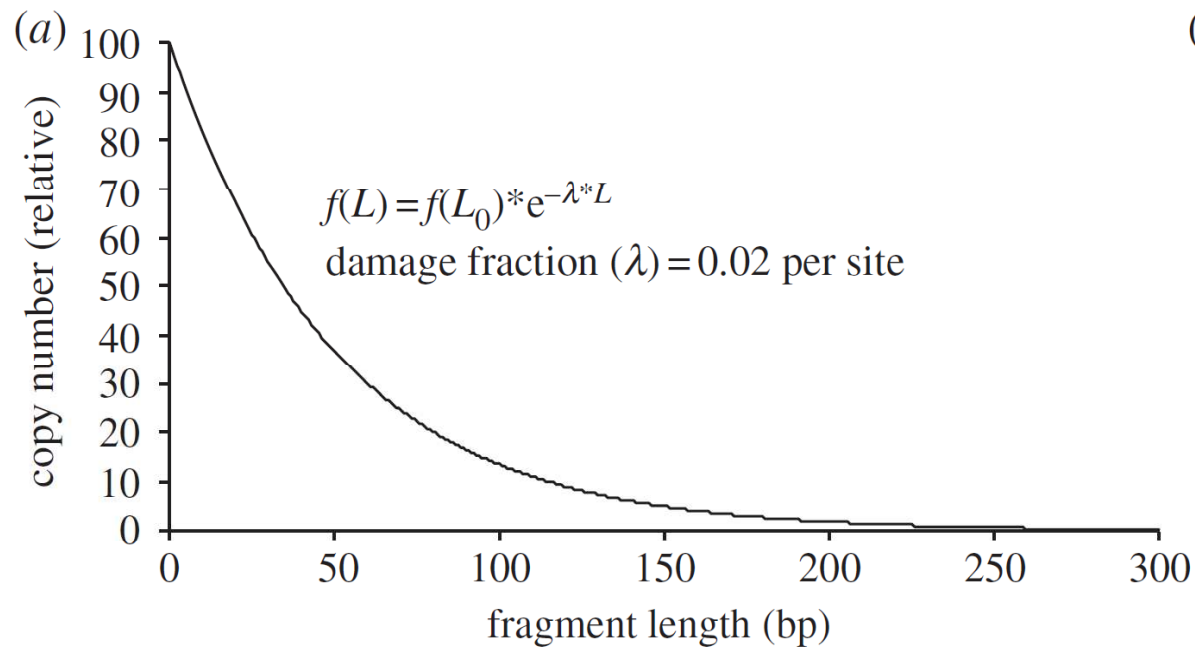
→ poškození způsobující inkorporaci nesprávných bází

Fragmentace:

většinou < 100 bp



Délky fragmentů se zmenšují zhruba exponenciálně



Pruvost et al. (2007):

2 cca. 3200 let staré vzorky téhož jedince pratura (*Bos primigenius*):

- 1) 1947, muzejní sbírky, 54 let → žádná použitelná aDNA
- 2) 2004 → autentická aDNA

To znamená, že 99 % DNA degradováno během pár desítek let

⇒ rychlost rozkladných procesů byla v muzejním prostředí více než 70× vyšší než v průběhu předchozích ~3200 let v půdě!

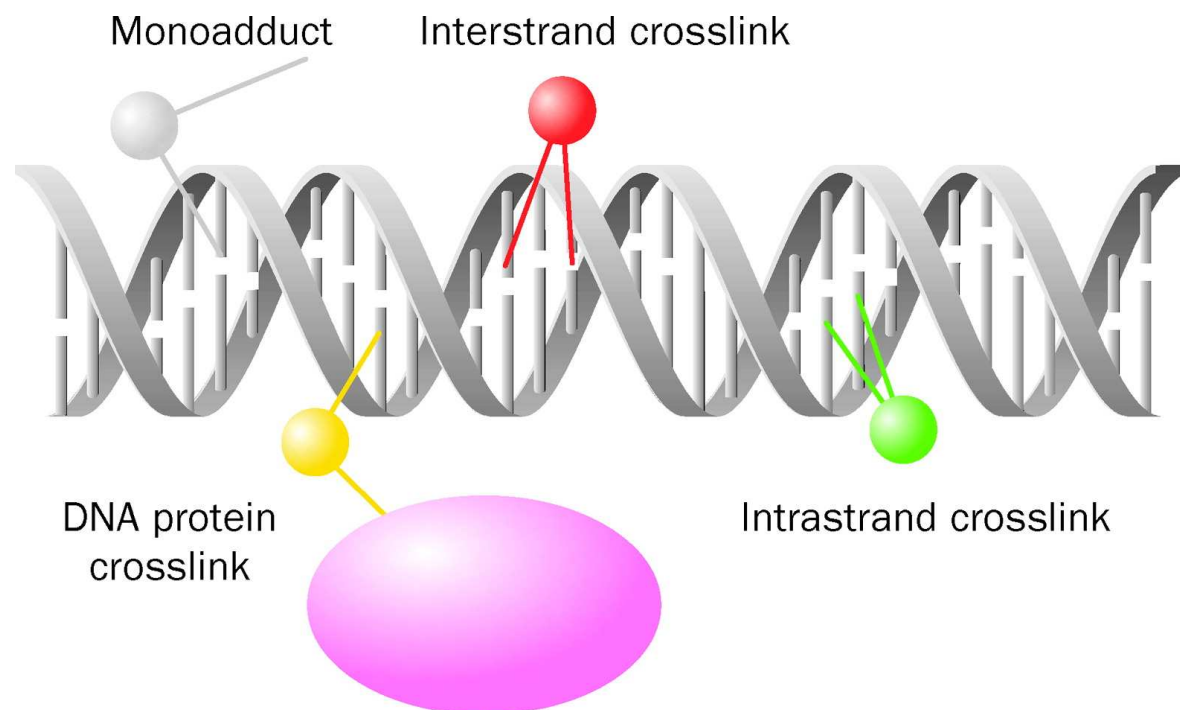
Blokující poškození:

Blokování postupu DNA polymerázy při PCR

modifikace nukleotidů

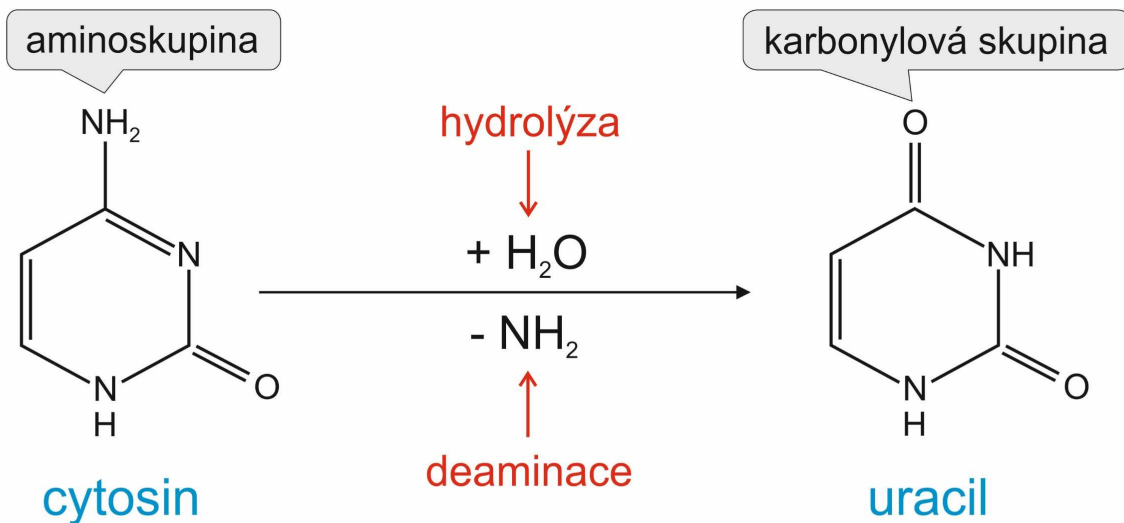
příčné vazby (*crosslinks*):

v rámci jednoho řetězce
mezi řetězci téže molekuly
mezi různými molekulami
mezi DNA a proteinem

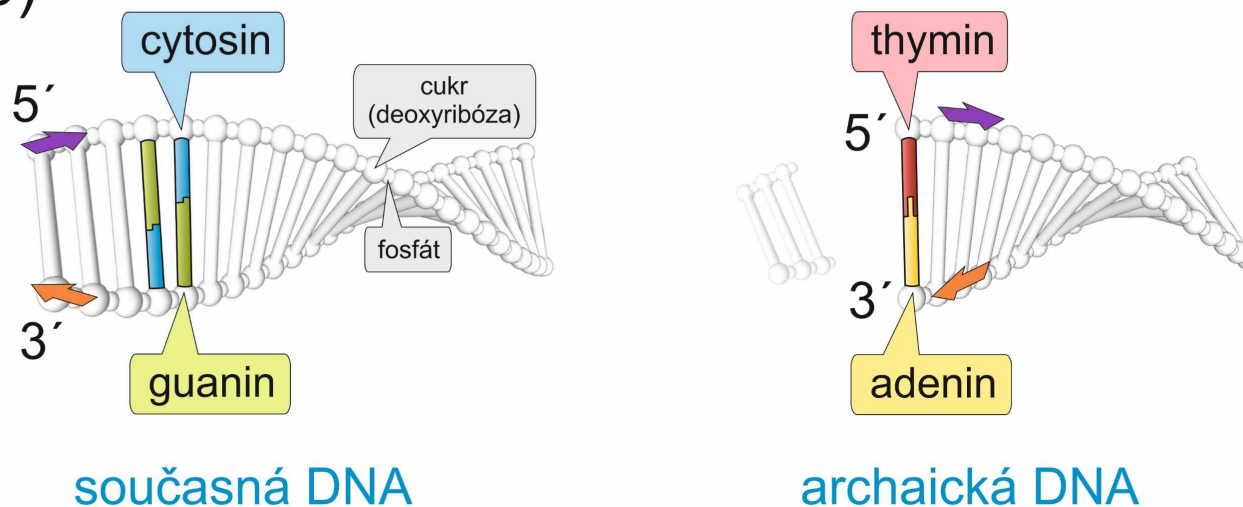


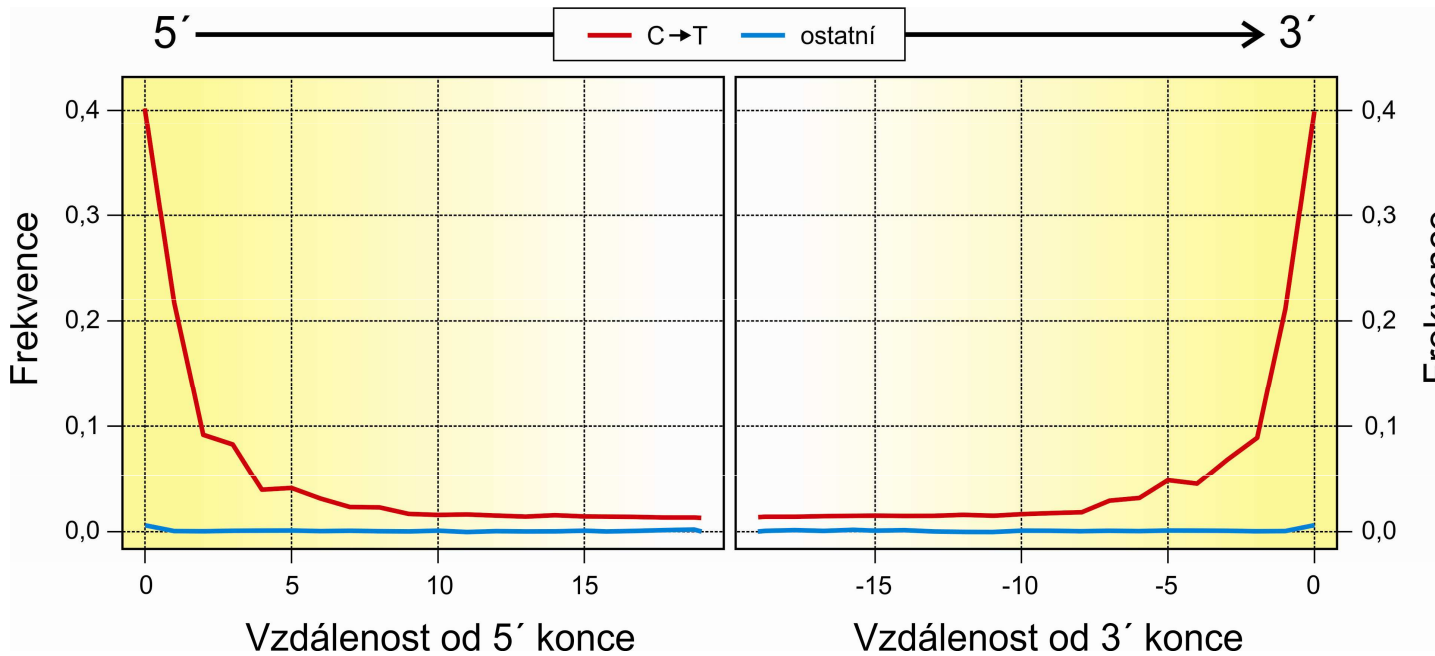
Inkorporace chybného nukleotidu:

a)

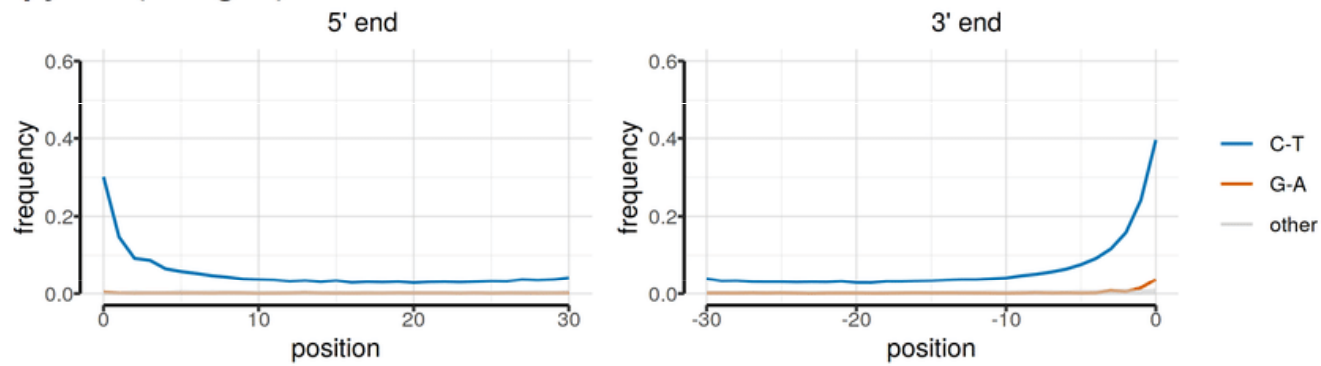


b)

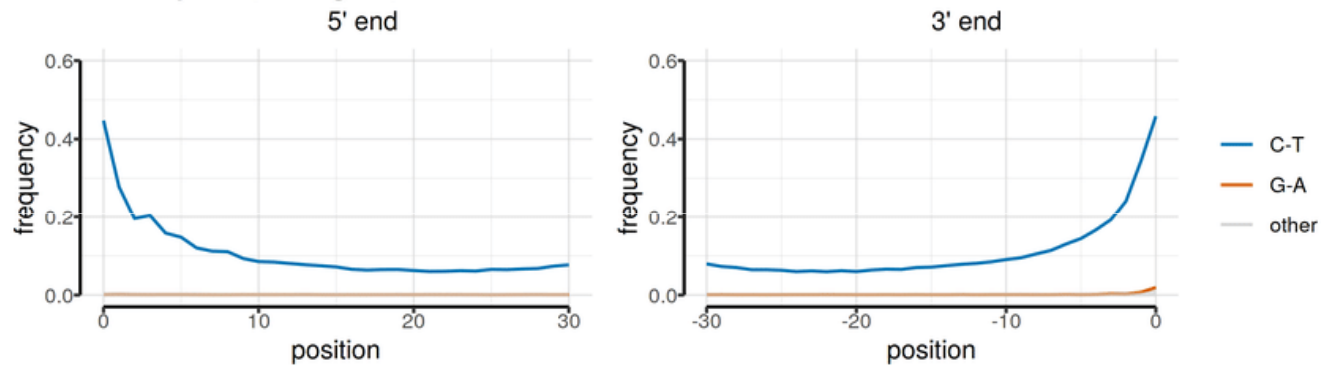




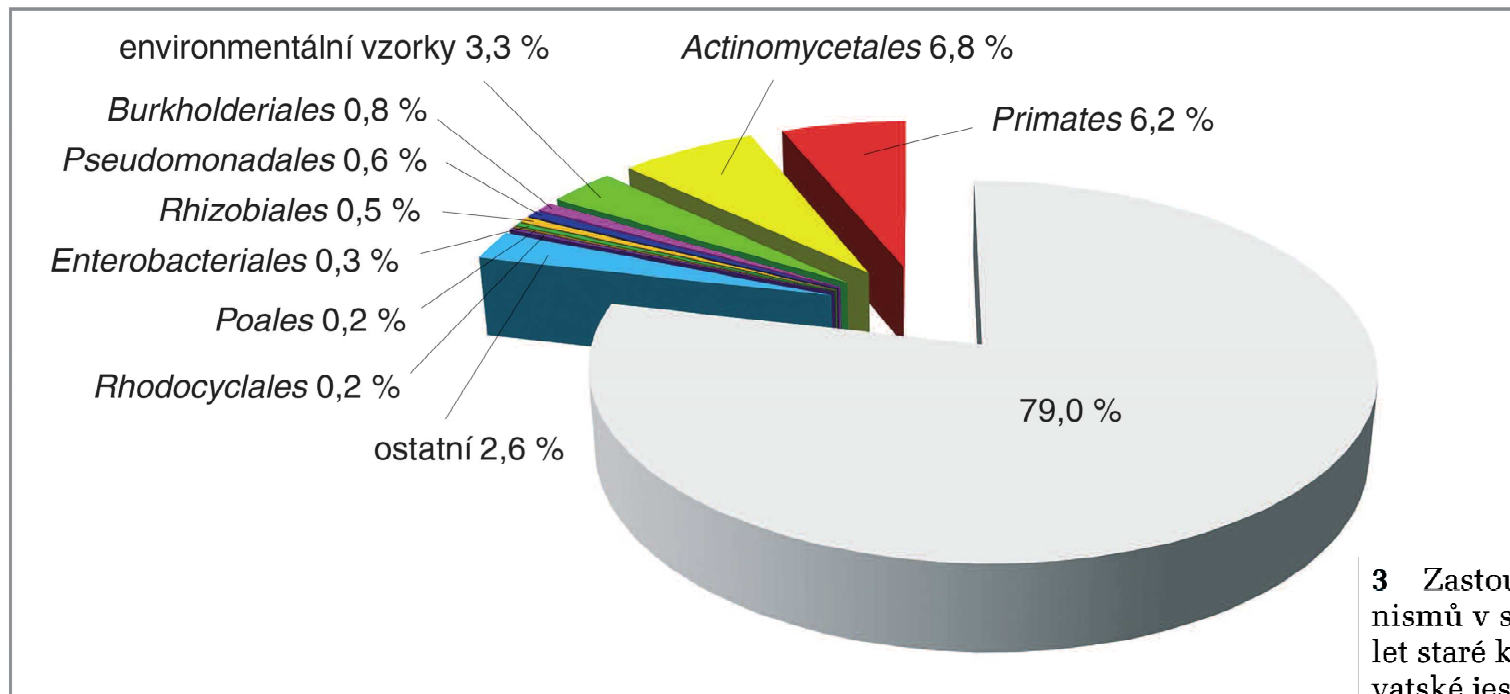
Spy 94a (shotgun)



Mezmaiskaya 2 (shotgun)



Kontaminace



3 Zastoupení nejhojnějších řádů organismů v sekvencích získaných z 38 tisíc let staré kosti neandertálce Vi-80 z chorvatské jeskyně Vindija. Největší podíl v sekvenovaných úsecích DNA měly půdní bakterie řádu *Actinomycetales* (6,8 %) a primáti (6,2 %). Průměrná shoda s databázovými sekvencemi primátů byla 98,8 % (u ostatních řádů 92–98 %), což ukazuje na blízkou taxonomickou příbuznost identifikovaných sekvencí primátů ve srovnání s ostatními řády. Skupina tzv. environmentálních vzorků, což jsou neidentifikované organismy vyskytující se ve vzorcích půdy, mořské vody, obsahu střeva apod., tvořila 3,3 % sekvencí. Za povšimnutí stojí, že naprostou většinu (79 %) sekvencí se nepodařilo přiřadit k žádné databázové sekvenci. Upraveno podle: R. E. Green a kol. (2006)



PRÁCE SE SOUBORY

DNA databáze:

EMBL (European Molecular Biology Laboratory) – European Bioinformatics Institute, Hinxton, UK: <http://www.ebi.ac.uk/embl/>

GenBank – NCBI (National Center for Biotechnology Information), Bethesda, Maryland, USA: <http://www.ncbi.nlm.nih.gov/Genbank/>

DDBJ (DNA Data Bank of Japan) – National Institute of Genetics, Mishima, Japan: <http://www.ddbj.nig.ac.jp/>

Proteinové databáze:

SWISS-PROT – University of Geneva & Swiss Institute of Bioinformatics: <http://www.expasy.ch/sprot/> a <http://www.ebi.ac.uk/swissprot/>

PIR (Protein Information Resource) – NBRF (National Biomedical Research Foundation, Washington, D.C., USA) & Tokyo University & JIPID (Japanese International Protein Information Database, Tokyo) & MIPS (Martinsried Institute for Protein Sequences, Martinsried, Germany): <http://www-nbrf.georgetown.edu/>

PRF/SEQDB (Protein Resource Foundation) – Ósaka, Japan: <http://www.prf.or.jp/en/os.htm>

PDB (Protein Data Bank) – University of New Jersey, San Diego & Super-computer Center, University of California & National Institute of Standards and Technology: <http://www.rcsb.org/pdb/>

Formáty souborů:

FASTA:

>H_sapiens

```
ATGACCCCAATACGCAAAATTAACCCCTAATAAAATTAATTAACCACTCATTTCATCGACCTCCCCACCC
CATCCAACATCTCCGCATGATGAACTTCGGCTCACTCCTTGGCGCCTGCCTGATCCTCCAAATCACCAC
AGGACTATTCCTAGCCATACTACTCACCAGACGCCTCAACCGCCTTTTCATCAATCGCCCACATCACT
CGAGACGTAAATTATGGCTGAATCATCCGCTACCTTCACGCCAATGGCGCCTCAATATTCTTTATCTGCC
TCTTCCTACACATCGGGCGAGGCCTATATTACGGATCATTTCTCTACTCAGAAACCTGAAACATCGGCAT
```

...

>P_troglod

```
ATGACCCCGACACGCAAAATTAACCCACTAATAAAATTAATTAATCACTCATTTTATCGACCTCCCCACCC
CATCCAACATTTCCGCATGATGGAECTTCGGCTCACTTCTCGGCGCCTGCCTAATCCTTCAAATTACCAC
AGGATTATTCCTAGCTATACTACTCACCAGACGCCTCAACCGCCTTCTCGTCGATCGCCCACATCACC
CGAGACGTAAACTATGGTTGGATCATCCGCTACCTCCACGCTAACGGCGCCTCAATATTTTTTTATCTGCC
TCTTCCTACACATCGGCCGAGGTCTATATTACGGCTCATTTCTCTACCTAGAAACCTGAAACATTGGCAT
```

...

>P_paniscus

```
ATGACCCCAACACGCAAAATCAACCCACTAATAAAATTAATTAATCACTCATTTTATCGACCTCCCCACCC
CATCCAATATTTCCACATGATGAACTTCGGCTCACTTCTCGGCGCCTGCCTAATCCTTCAAATCACCAC
AGGACTATTCCTAGCTATACTACTCACCAGACGCCTCAACCGCCTTCTCATCGATCGCCCACATTACC
CGAGACGTAAACTATGGTTGAATCATCCGCTACCTTCACGCTAACGGCGCCTCAATACTTTTCATCTGCC
TCTTCCTACACGTCGGTCGAGGCCTATATTACGGCTCATTTCTCTACCTAGAAACCTGAAACATTGGCAT
```

...

Formáty souborů:

GenBank:

ORIGIN

```
1  tgaaatgaag atattctctt ctcaagacat caagaagaag gaactactcc ccaccaccag
61  cacccaaagc tggcattcta attaaactac ttcttgtgta cataaattta catagtacaa
121 tagtacatth atgtatatcg tacattaaac tattttcccc aagcatataa gcaagtacat
181 ttaatcaatg atataggcca taaaacaatt atcaacataa actgatacaa accatgaata
241 ttataactaat acatcaaatt aatgctttaa agacatatct gtgttatctg acatacacca
301 tacagtcata aactcttctc ttccatatga ctatcccctt ccccatthtg tctattaatc
361 taccatcctc cgtgaaacca acaacccgcc caccaatgcc cctcttctcg ctccggggcc
421 attaaacttg ggggtagcta aactgaaact ttatcagaca tctgggttct acttcagggc
481 catcaaatgc gttatcgccc atacgttccc cttaaataag acatctcgat ggtatcgggt
541 ctaatcagcc catgaccaac ataactgtgg tgtcatgcat ttggtattht tttatthtgg
601 cctactthca tcaacatagc cgtcaaggca tgaaaggaca gcacacagtc tagacgcacc
661 tacgggtgaag aatcattagt ccgcaaaacc caatcaccta aggctaatta ttcatgcttg
721 ttagacataa atgctactca ataccaaatt ttaactctcc aaacccccca acccctcct
781 cttaatgcca aacccccaaa aactaagaa cttgaaagac atatattht aactatcaaa
841 ccctatgtcc tgatcgattc tagtagttcc caaaatatga ctcatattht agtacttgta
901 aaaatthtac aaaatcatgc tccgtgaacc aaaactctaa tcacactcta ttacgcaata
961 aatattaaca agttaatgta gcttaataac aaagcaaagc actgaaaatg cttagatgga
1021 taatthtatc cca
```

//

Formáty souborů:

PHYLIP (“interleaved” format):

6 1120

```
H_sapiens      ATGACCCCAA TACGCAAAAT TAACCCCCTA ATAAAATTAA TTAACCACTC
P_troglod      ATGACCCCGA CACGCAAAAT TAACCCACTA ATAAAATTAA TTAATCACTC
P_paniscus     ATGACCCCAA CACGCAAAAT CAACCCACTA ATAAAATTAA TTAATCACTC
G_gorilla      ATGACCCCTA TACGCAAAAC TAACCCACTA GCAAAACTAA TTAACCACTC
P_pygmaeus     ATGACCCCAA TACGCAAAAC CAACCCACTA ATAAAATTAA TTAACCACTC
H_lar          ATGACCCCCC TGCGCAAAAC TAACCCACTA ATAAAACTAA TCAACCACTC

                ATTCATCGAC CTCCCCACCC CATCCAACAT CTCCGCATGA TGAAACTTCG
                ATTTATCGAC CTCCCCACCC CATCCAACAT TTCCGCATGA TGGAACTTCG
                ATTTATCGAC CTCCCCACCC CATCCAATAT TTCCACATGA TGAAACTTCG
                ATTCATTGAC CTCCCTACCC CGTCCAACAT CTCCACATGA TGAAACTTCG
                ACTCATCGAC CTCCCCACCC CATCAAACAT CTCTGCATGA TGGAACTTCG
                ACTTATCGAC CTTCCAGCCC CATCCAACAT TTCTATATGA TGAAACTTTG
```

Formáty souborů:

NEXUS (PAUP*, “interleaved”):

```
#NEXUS
begin data;
dimensions ntax=6 nchar=1120;
format datatype=DNA interleave datatype=DNA missing=? gap=-;
matrix
P_troglod   ATGACCCCGACACGCAAAATTAACCCACTAATAAAATTAATTAATCACTC
P_paniscus  ATGACCCCAACACGCAAAATCAACCCACTAATAAAATTAATTAATCACTC
H_sapiens   ATGACCCCAATACGCAAAATTAACCCCTAATAAAATTAATTAACCACTC
G_gorilla   ATGACCCCTATACGCAAAACTAACCCACTAGCAAACTAATTAACCACTC
P_pygmaeus  ATGACCCCAATACGCAAAACCAACCCACTAATAAAATTAATTAACCACTC
H_lar       ATGACCCCTGCGCAAAACTAACCCACTAATAAACTAATCAACCACTC

P_troglod   ATTTATCGACCTCCCCACCCCATCCAACATTTCCGCATGATGGAACTTCG
P_paniscus  ATTTATCGACCTCCCCACCCCATCCAATATTTCCACATGATGAAACTTCG
H_sapiens   ATTCATCGACCTCCCCACCCCATCCAACATCTCCGCATGATGAAACTTCG
G_gorilla   ATTCATTGACCTCCCTACCCCGTCCAACATCTCCACATGATGAAACTTCG
P_pygmaeus  ACTCATCGACCTCCCCACCCCATCAAACATCTCTGCATGATGGAACTTCG
H_lar       ACTTATCGACCTTCCAGCCCATCCAACATTTCTATATGATGAAACTTTG

end;
```

Formáty souborů:

Clustal:

```
P_troglod ATGACCCCGACACGCAAAATTAACCCACTAATAAAATTAATTAATCACTCATTATCGAC
P_paniscus ATGACCCCAACACGCAAAATCAACCCACTAATAAAATTAATTAATCACTCATTATCGAC
H_sapiens ATGACCCCAATACGCAAAATTAACCCCTAATAAAATTAATTAACCACTCATTATCGAC
G_gorilla ATGACCCCTATACGCAAAACTAACCCACTAGCAAAACTAATTAACCACTCATTATCGAC
P_pygmaeus ATGACCCCAATACGCAAAACCAACCCACTAATAAAATTAATTAACCACTCACTCATCGAC
H_lar ATGACCCCCCTGCGCAAAACTAACCCACTAATAAAACTAATCAACCACTCACTTATCGAC
*****          *****          *****  ***          *****  *****  ** *****  * **  ***
```

```
P_troglod CTCCCCACCCCATCCAACATTTCCGCATGATGGAACTTCGGCTCACTTCTCGGCGCCTGC
P_paniscus CTCCCCACCCCATCCAATATTTCCACATGATGAAACTTCGGCTCACTTCTCGGCGCCTGC
H_sapiens CTCCCCACCCCATCCAACATCTCCGCATGATGAAACTTCGGCTCACTCCTTGGGCGCCTGC
G_gorilla CTCCCTACCCCGTCCAACATCTCCACATGATGAAACTTCGGCTCACTCCTTGGTGCCTGC
P_pygmaeus CTCCCCACCCCATCAAACATCTCTGCATGATGGAACTTCGGCTCACTTCTAGGCGCCTGC
H_lar CTTCCAGCCCCATCCAACATTTCTATATGATGAAACTTTGGTTCACTCCTAGGCGCCTGC
** ** ***** ** ** ** ** ***** ***** ** ***** ** ** *****
```

Formáty souborů:

FASTQ:

Line 1 begins with a '@' character and is followed by a sequence identifier and an *optional* description (like a [FASTA](#) title line).

Line 2 is the raw sequence letters.

Line 3 begins with a '+' character and is *optionally* followed by the same sequence identifier (and any description) again.

Line 4 encodes the quality values for the sequence in Line 2, and must contain the same number of symbols as letters in the sequence.

Illumina sequence identifiers:

@HWUSI-EAS100R:6:73:941:1973#0/1

HWUSI-EAS100R	the unique instrument name
6	flowcell lane
73	tile number within the flowcell lane
941	'x'-coordinate of the cluster within the tile
1973	'y'-coordinate of the cluster within the tile
#0	index number for a multiplexed sample (0 for no indexing)
/1	the member of a pair, /1 or /2 (<i>paired-end or mate-pair reads only</i>)

BLAST (*Basic Local Alignment Search Tool*):

Nucleotide BLAST: nucleotide → nucleotide

blastx: nucleotide → protein

tblastn: protein → nucleotide

Protein BLAST: protein → protein

megablast

blastn

discontiguous megablast

BLAST

Basic Local Alignment Search Tool

BLAST finds regions of similarity between biological sequences. The program compares nucleotide or protein sequences to sequence databases and calculates the statistical significance. [Learn more](#)

Using BLAST+ in Docker and on the cloud: [Webinar](#) on December 9, 2020.

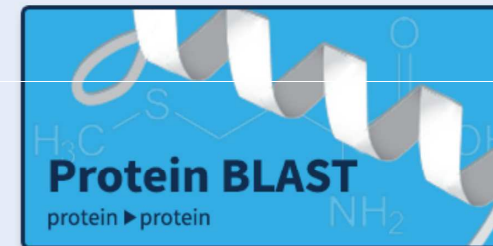
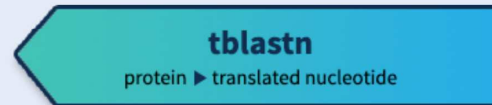
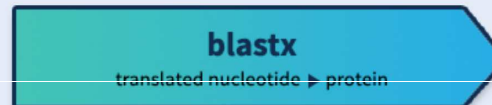
NEWS

In this webinar, the NCBI BLAST team will demonstrate containerized BLAST+ in Docker that is ready to use locally and in the cloud.

Wed, 02 Dec 2020 12:00:00 EST

[More BLAST news...](#)

Web BLAST



BLAST Genomes

Enter organism common name, scientific name, or tax id

Search

Human

Mouse

Rat

Microbes

Standard Nucleotide BLAST

blastn blastp blastx tblastn tblastx

BLASTN programs search nucleotide databases using a nucleotide query. [more...](#)


[Reset page](#) [Bookmark](#)

Enter Query Sequence

Enter accession number(s), gi(s), or FASTA sequence(s)

```
CAAAAACACTAAGAAGCTTGAAGACATATACTATTAACATCTAACCCCTATGTCCTGATCAATTCTAGTAGTT
CAAAAAATATGACTTATATTTTAGTCTTGTAAAAATTTTGCAAAAAATGCCCCATAAGCCAAAACCTCTAAT
TATACCCCTATACGCAATAAACAAATAGTAAGTAAATGTAGCTTAAATAAAAAGCAAAGCCTGAAAAATGCTTAG
ATGGATAATTTTATCCATAAACACAAAGGTTTGGTC
```

Clear Query subrange
From
To

New columns added to the Description Table
Click 'Select Columns' or 'Manage Columns'. 

Or, upload file

Soubor nevybrán

Job Title

Enter a descriptive title for your BLAST search

Align two or more sequences

Choose Search Set

Database Standard databases (nr etc.): rRNA/ITS databases Genomic + transcript databases Betacoronavirus

Nucleotide collection (nr/nt)

Organism

Optional

Enter organism name or id--completions will be suggested exclude

Enter organism common name, binomial, or tax id. Only 20 top taxa will be shown

Exclude

Optional

Models (XM/XP) Uncultured/environmental sample sequences

Limit to

Optional

Sequences from type material

Entrez Query

Optional

[YouTube](#) [Create custom database](#)

Enter an Entrez query to limit search

Program Selection

Optimize for

- Highly similar sequences (megablast)
- More dissimilar sequences (discontiguous megablast)
- Somewhat similar sequences (blastn)

Choose a BLAST algorithm

BLAST

Search database **Nucleotide collection (nr/nt)** using **Megablast (Optimize for highly similar sequences)**

Show results in a new window

[+ Algorithm parameters](#)

BLAST is a registered trademark of the National Library of Medicine

[Support center](#) [Mailing list](#) [YouTube](#)

NCBI

National Center for Biotechnology Information, U.S. National Library of Medicine
8600 Rockville Pike, Bethesda MD, 20894 USA



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[Save Search](#)

[Search Summary](#) ▼

[? How to read this report?](#)

[▶ BLAST Help Videos](#)

[↶ Back to Traditional Results Page](#)

Job Title	Nucleotide Sequence
RID	WX41A1JA013 <small>Search expires on 12-09 03:27 am</small> Download All ▼
Program	BLASTN ? Citation ▼
Database	nt See details ▼
Query ID	lcl Query_50051
Description	None
Molecule type	dna
Query Length	950
Other reports	Distance tree of results MSA viewer ?

Filter Results

Organism only top 20 will appear exclude

Type common name, binomial, taxid or group name

[+ Add organism](#)

Percent identity to **E value** to **Query Coverage** to

[Filter](#) [Reset](#)

Descriptions

[Graphic Summary](#)

[Alignments](#)

[Taxonomy](#)

Sequences producing significant alignments

[Download](#) ▼

New [Select columns](#) ▼

Show

100 ▼

[?](#)

select all 100 sequences selected

[GenBank](#)

[Graphics](#)

[Distance tree of results](#)

	Description	Common Name	Max Score	Total Score	Query Cover	E value	Per. Ident	Acc. Len	Accession
<input checked="" type="checkbox"/>	Mus cypriacus isolate MM1375 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gen...	Cypriot mouse	878	1756	100%	0.0	100.00%	1062	EU106210.1
<input checked="" type="checkbox"/>	Mus cypriacus isolate MM1373 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gen...	Cypriot mouse	878	1751	100%	0.0	100.00%	1062	EU106208.1
<input checked="" type="checkbox"/>	Mus cypriacus isolate MM1372 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gen...	Cypriot mouse	874	1747	100%	0.0	99.79%	1062	EU106207.1
<input checked="" type="checkbox"/>	Mus cypriacus isolate MM1391 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gen...	Cypriot mouse	872	1653	95%	0.0	99.79%	1015	EU106216.1
<input checked="" type="checkbox"/>	Mus cypriacus isolate MM1388 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gen...	Cypriot mouse	872	1739	100%	0.0	99.79%	1062	EU106214.1
<input checked="" type="checkbox"/>	Mus cypriacus isolate MM1387 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gen...	Cypriot mouse	872	1712	100%	0.0	99.79%	1062	EU106213.1
<input checked="" type="checkbox"/>	Mus cypriacus isolate MM1374 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gen...	Cypriot mouse	872	1717	100%	0.0	99.79%	1062	EU106209.1
<input checked="" type="checkbox"/>	Mus cypriacus isolate MM1364 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gen...	Cypriot mouse	872	1734	100%	0.0	99.79%	1062	EU106204.1
<input checked="" type="checkbox"/>	Mus cypriacus isolate MM1360 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gen...	Cypriot mouse	872	1706	100%	0.0	99.79%	1062	EU106200.1
<input checked="" type="checkbox"/>	Mus cypriacus isolate MM1371 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gen...	Cypriot mouse	872	1739	100%	0.0	99.79%	1062	EU106199.1
<input checked="" type="checkbox"/>	Mus cypriacus isolate MM1383 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gen...	Cypriot mouse	872	1712	100%	0.0	99.79%	1062	EU106198.1
<input checked="" type="checkbox"/>	Mus cypriacus isolate MM1358 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gen...	Cypriot mouse	872	1706	100%	0.0	99.79%	1062	EU106195.1

100 sequences selected

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Mus cypriacus isolate MM1375 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gene, complete sequence; mitochondrial

Sequence ID: [EU106210.1](#) Length: 1062 Number of Matches: 2

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Range 1: 1 to 475 [GenBank](#) [Graphics](#) Next Match Previous Match

Score	Expect	Identities	Gaps	Strand
878 bits(475)	0.0	475/475(100%)	0/475(0%)	Plus/Plus
Query 1	TGTA AACCTGAAATGAAGATATTTCTTTCTCAAGACATCAAGAAGAAGGAAC TTATTCCC			60
Sbjct 1	TGTA AACCTGAAATGAAGATATTTCTTTCTCAAGACATCAAGAAGAAGGAAC TTATTCCC			60
Query 61	CACCACCAACACCCAAAGCTGGTATTCTAGTTAAACTACTTCTTGAGTACATAAATTTAC			120
Sbjct 61	CACCACCAACACCCAAAGCTGGTATTCTAGTTAAACTACTTCTTGAGTACATAAATTTAC			120
Query 121	ATAGTACATTAGTACATTTATGTATATCGTACATTA AATTATATTTCCCAAGCATATAAG			180
Sbjct 121	ATAGTACATTAGTACATTTATGTATATCGTACATTA AATTATATTTCCCAAGCATATAAG			180
Query 181	CACGTAAATTA AATTAATGACATAGCACATAAAACGATATTTAACATAAAA TACTACACA			240
Sbjct 181	CACGTAAATTA AATTAATGACATAGCACATAAAACGATATTTAACATAAAA TACTACACA			240
Query 241	ACATGAATATTATATTA AATACATTAAGTTAATGCTTTAAAGACATATCTGTGTTATCTG			300
Sbjct 241	ACATGAATATTATATTA AATACATTAAGTTAATGCTTTAAAGACATATCTGTGTTATCTG			300
Query 301	ACATACACCATAAAGTCATAAACCCTTCTCTTCCATATGACTATCCCCTTCCCATTGG			360
Sbjct 301	ACATACACCATAAAGTCATAAACCCTTCTCTTCCATATGACTATCCCCTTCCCATTGG			360
Query 361	TCTATTAATCTACCATCCTCCGTGAAACCAACAACCCGCCACCTATGCCCTCTTCTCG			420
Sbjct 361	TCTATTAATCTACCATCCTCCGTGAAACCAACAACCCGCCACCTATGCCCTCTTCTCG			420
Query 421	CTCCGGGCCCATTA AACTTGGGGGTAGCTAAACTGAACTTTATCAGACATCTGG			475
Sbjct 421	CTCCGGGCCCATTA AACTTGGGGGTAGCTAAACTGAACTTTATCAGACATCTGG			475

Range 2: 588 to 1062 [GenBank](#) [Graphics](#) Next Match Previous Match First Match

Sekvence conversion:

Complementary Sequence Conversion Tool

GenBank:

původně Los Alamos National Laboratory

90's → NCBI (*National Center for Biotechnology Information*)

20 000 přímých podání, 200 000 hromadných podání denně

každých 10 měsíců zdvojnásobení počtu

GenBank

ncbi.nlm.nih.gov/nuccore/?term=mus+cypriacus

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Species Summary 20 per page Sort by Default order Send to: Filters: [Manage Filters](#)

Animals (29) Customize ...

Molecule types genomic DNA/RNA (29) Customize ...

Source databases INSDC (GenBank) (29) Customize ...

Sequence Type Nucleotide (29)

Genetic compartments Mitochondrion (28)

Sequence length Custom range...

Release date Custom range...

Revision date Custom range...

Items: 1 to 20 of 29

<< First < Prev Page 1 of 2 Next > Last >>

- [Mus cypriacus mitochondrial partial cytb gene for cytochrome b](#)
1. 1,140 bp linear DNA
Accession: FR751074.1 GI: 323713991
[Protein](#) [PubMed](#) [Taxonomy](#)
[GenBank](#) [FASTA](#) [Graphics](#)
- [Mus cypriacus isolate MM1381 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gene, complete sequence; mitochondrial](#)
2. 1,062 bp linear DNA
Accession: EU106281.1 GI: 157266050
[PubMed](#) [Taxonomy](#)
[GenBank](#) [FASTA](#) [Graphics](#) [PopSet](#)
- [Mus cypriacus isolate MM1377 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gene, complete sequence; mitochondrial](#)
3. 1,062 bp linear DNA
Accession: EU106280.1 GI: 157266049
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- mus cypriacus control region (0)

Sem zadejte hledaný výraz

CES 20:09
CSQ 07.12.2020

Mus cypricus isolate MM1381 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gene, complete sequence; mitochondrial

GenBank: EU106281.1

[FASTA](#) [Graphics](#) [PopSet](#)

Go to:

LOCUS EU106281 1062 bp DNA linear ROD 16-NOV-2007
DEFINITION Mus cypricus isolate MM1381 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gene, complete sequence; mitochondrial.

ACCESSION EU106281
VERSION EU106281.1
KEYWORDS .
SOURCE mitochondrion Mus cypricus (Cypriot mouse)

ORGANISM [Mus cypricus](#)
Eukaryota; Metazoa; Chordata; Craniata; Vertebrata; Euteleostomi; Mammalia; Eutheria; Euarchontoglires; Glires; Rodentia; Myomorpha; Muroidea; Muridae; Murinae; Mus; Mus.

REFERENCE 1 (bases 1 to 1062)
AUTHORS Macholan,M., Vyskocilova,M., Bonhomme,F., Krystufek,B., Orth,A. and Vohralik,V.

TITLE genetic variation and phylogeography of free-living mouse species (genus Mus) in the Balkans and the Middle East

JOURNAL Mol Ecol 16 (22), 4774-4788 (2007)
PUBMED [17908218](#)

REFERENCE 2 (bases 1 to 1062)
AUTHORS Macholan,M., Vyskocilova,M., Bonhomme,F., Krystufek,B., Orth,A. and Vohralik,V.

TITLE Direct Submission
JOURNAL Submitted (20-AUG-2007) Laboratory of Mammalian Evolutionary Genetics, Institute of Animal Physiology and Genetics, Acad. Sci. Czech Rep., Veveri 97, Brno CZ-60200, Czech Republic

FEATURES
Location/Qualifiers
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/mol_type="genomic DNA"
/isolate="MM1381"
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[tRNA](#) 38..105
/product="tRNA-Pro"
[D-loop](#) 106..982
gap 476..587
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[tRNA](#) 983..1049
/product="tRNA-Phe"

ORIGIN
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61 caccaccaac acccaagct ggtattctag ttaactact tcttgagtac ataaatttac
121 atagtacatt agtacattta tgtatatcgt acattaaatt atattcccca agcatataag
181 cacgtaaat aaattaatga catagcacat aaaacgata ttaacataaa atactacaca
241 acatgaatat tatattaat acattaagtt aatgctttaa agacatact gtgttatctg
301 acatacaca taaagtcata aacccttct tcccatatga ctatcccctt ccccatgttg
361 tctattaatc taccatctc cgtgaaacca acaaccgcc cacctatgcc cctcttctcg
421 ctccgggcc attaaactg gggtagcta aactgaaact ttatcagaca tctgg
[gap 112 bp] [Expand Ns](#)
588 cat ttggtattt
601 tttatttgg tctactttca tcaacatagc cgtcaaggca tgaaggaca gcacacagtc
661 tagacgacc tacggtgaag aatcattagt ctcataacc caatcaccca aggctaatta
721 ttcagcttg ttagacataa aattattcaa taccagatt taactctcca aaccccccc
781 accccatcc tcttaatgcc aaacccaaa aacactaaga acttgaaga catatactat
841 taactatcta accctatgct ctgatcaatt ctgtagttc aaaaaatag acttatattt
901 tagttctgt aaaaatttg caaataatg ccccataagc caaactcta attataccct
961 attacgcaat aaacaatag agttaatgt agcttaataa aaagcaagc actgaaaatg
1021 cttagatgga taattttatc cataaacac aaagtttgg tc

//

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Mus cypricus isolate MM1381 tRNA-Thr gene, partial sequence; and tRNA-P Nucleotide

mus cypricus (29) Nucleotide

mus cypricus control region (0) Nucleotide

On the Probability of Fixation of Mutant Genes in a Population

The frequency of multiple paternity suggests that sperm competition is common in...

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Mus cypriacus isolate MM1381 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gene, complete sequence; mitochondrial

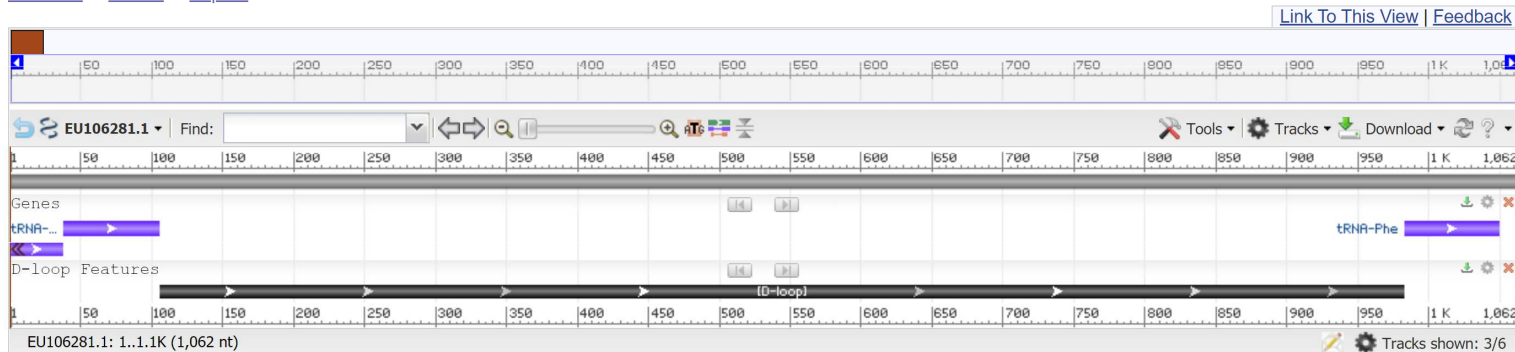
GenBank: EU106281.1

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gene, D-loop, and tRNA-Phe gene, complete sequence; mitochondrial
TGTAACCTGAAATGAAGATATTCTTTCTCAAGACATCAAGAAGAAGGAAC TTATCCCCACCACCAAC
ACCCAAAGCTGGTATTCTAGTTAACTACTTCTTGAGTACATAAATTTACATAGTACATTAGTACATTTA
TGTATATCGTACATTAATTATATTCCCCAAGCATATAAGCACGTAATAATAATTAATGACATAGCACAT
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NNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNN
NNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNN
NNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNNN
CGTCAAGGCATGAAAGGACAGCACAGTCTAGACGCACCTACGGTGAAGAATCATTAGTCTCATAACC
CAATCACCCAAGGCTAATTATTCATGCTTGTAGACATAAAATATTCAATACCAGATTTAACTCTCCA
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TAATCTAACCCTATGCTCTGATCAATTCTAGTAGTTCAAAAAATAGACTTATATTTAGTCTTGT
AAAAATTTGCAAAATGATGCCATAAGCCAAACTCTAATTATACCCTATTACGCAATAAACAATAGT
AAGTTAATGTAGCTTAATAAAAAGCAAAGCACTGAAAATGCTTAGATGGATAATTTATCCATAAACAC
AAAGTTTGGTC
```

Mus cypriacus isolate MM1381 tRNA-Thr gene, partial sequence; and tRNA-Pro gene, D-loop, and tRNA-Phe gene, complete sequence; mitochondrial

GenBank: EU106281.1

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Mus cypriacus isolate MM1381 tRNA-Thr

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 Mus cypriacus isolate MM1381 tRNA-Thr gene, partial
sequence; and tRNA-Pro gene,... Nucleotide

 mus cypriacus (29) Nucleotide

mus cypriacus control region (0)

Seřazení sekvencí:

Sekvence 1 TTGTACGACGG
 Sekvence 2 TTGTACGACG

```

TTGTACGACGG   TTGT---ACGACGG
|||||         |||   |||
TTGTACGACG    TTGTACGACG
    
```

$$GP = g + hl$$

g - gap penalty
h - gap extension penalty
l - gap length

Gap penalty:

Sekvence 1 ACTTGTGCTTC
 Sekvence 2 ACGTGCTGCTC

Path 1
 ACTTG-TGCTTC
 ||| ||| |||
 ACGTGCTGCTC

Path 2
 ACTTGTGCTTC
 || | ||| ||| |
 AC--GTGCTGCTC

g = penalizace za výskyt mezery (1×)

h = extenze za každou „pomlčku“

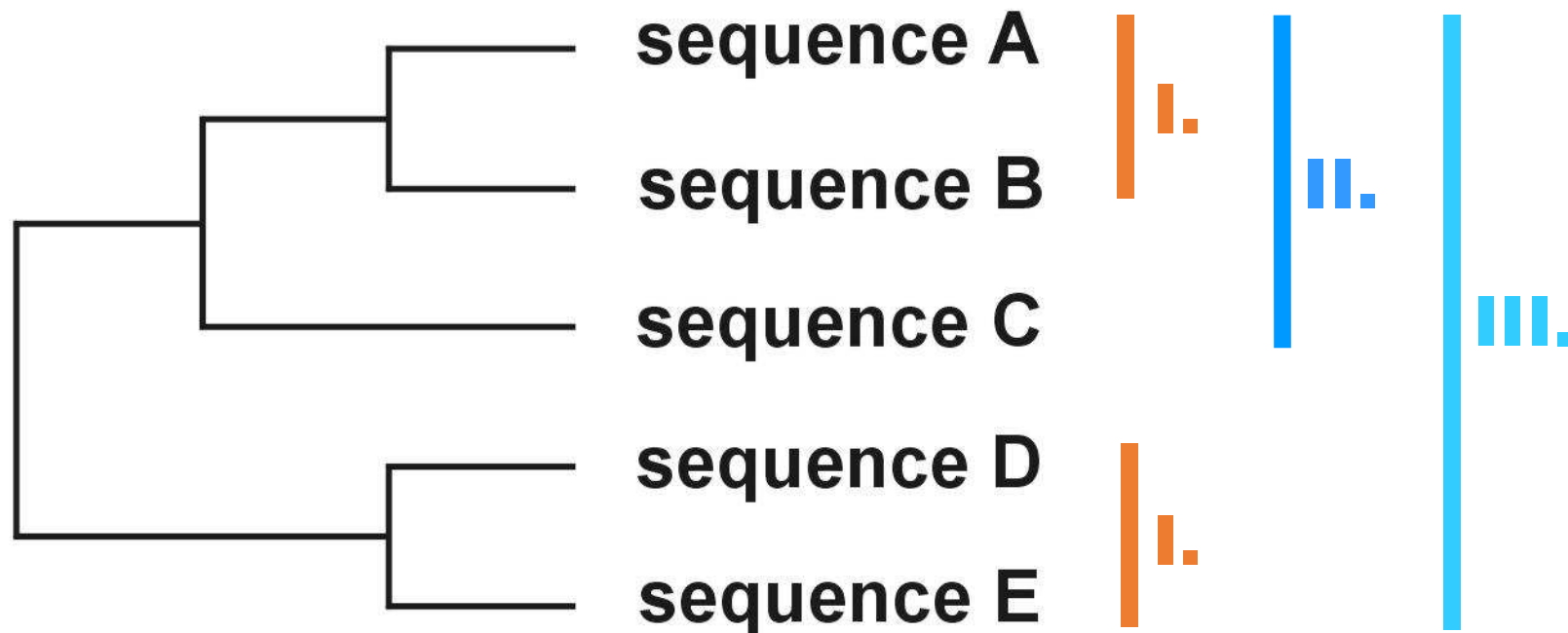
l = délka mezery (= počet „pomlček“)

Př.: GC-----TTAA
l = 5, *g* = *x*, *h* = 5*x*

Progresivní seřazení - ClustalX

3 phases:

1. Alignment of sequence pairs → pairwise distances
2. Construction of „guide tree“ (NJ)
3. Alignment of all sequences according to the guide tree



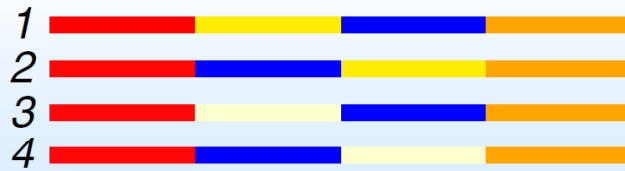
Další programy:

Clustal Omega – online

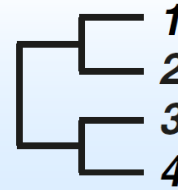
MUSCLE, MAFFT, Geneious

programy pro seřazení součástí mnoha populačněgenetických balíků

Align-free methods



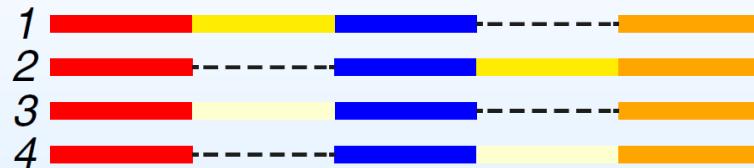
homologous sequence set



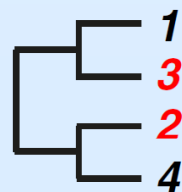
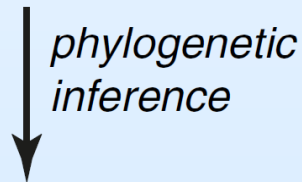
reference phylogeny

A

classical approach



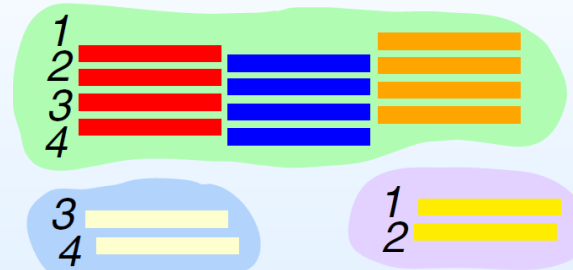
multiple sequence alignment



phylogenetic tree

B

alternative approach



alignment-free method

