

genotypst: A bioinformatics Typst package for sequence analysis and visualization

genotypst is a bioinformatics package for Typst that enables analysis and visualization of biological data. It provides functionality for parsing FASTA and Newick files and generating publication-ready visualizations, including multiple sequence alignments, sequence logos, genome maps, and phylogenetic trees.

Contents

Working with sequence data	2
Loading data	2
FASTA rendering	2
Pairwise sequence alignment	2
Rendering pairwise alignments	3
Dynamic programming matrix visualization	3
Scoring matrices	4
Multiple sequence alignments	5
Sequence logos	6
Color palettes	7
Amino acid palette	7
Nucleic acid palettes	7
Visualizing genomic loci with genome maps	8
Working with phylogenetic trees	9
Visualizing trees	9
Customizing visualizations	10
Font selection	10
Bibliography	12

Working with sequence data

genotypst provides functions to parse sequence data and produce different visualizations.

Loading data

The `parse-fasta` function reads FASTA data and returns a dictionary mapping sequence identifiers to their corresponding sequences.

```
#let sequences = parse-fasta(read("/docs/data/dna.fna"))

(
  seq_1: "AAGGGACACTGATTTCTCCCACAGCTGGCCGTGGACCGTAGTGTTCAGAACGCCACACAC",
  seq_2: "GCAATGGAGACAACATAGCCAACTACCTACTAGATGCCCTAGATCTGCCGCA",
  seq_3: "GGAACCTGGCGTTACAGACAGTTGTGAGCCACCACATGGGCCTGGGATTAAATTATAAAGCTCCTC",
)
```

FASTA rendering

Use `render-fasta` to display sequences in the standard FASTA format.

```
#render-fasta(sequences, max-width: 50)

>seq_1
AAGGGACACTGATTTCTCCCACAGCTGGCCGTGGACCGTAGTGTTC
AGAACGCCACACAC
>seq_2
GCAATGGAGACAACATAGCCAACTACCTACTAGATGCCCTAGATCTGCCG
CA
>seq_3
GGAACCTGGCGTTACAGACAGTTGTGAGCCACCACATGGGCCTGGGATT
AATATTATAAAGCTCCTC
```

In this example, `max-width` controls how many characters appear per line (default is 60).

Pairwise sequence alignment

Pairwise alignment is a method for comparing biological sequences, allowing quantification of sequence similarity and identification of evolutionary relationships between the residues of two sequences. genotypst supports both global alignment (end-to-end)¹ and local alignment (best-matching subsequences)², using dynamic programming with a user-defined scoring scheme (match/mismatch or a substitution matrix) and gap penalties. The `align-seq-pair` function performs pairwise alignment using either match/mismatch scores or a substitution matrix (see the scoring matrices section below).

```

#let dna_pair_alignment = align-seq-pair(
  "ACT",
  "ACGT",
  match-score: 3,
  mismatch-score: -1,
  gap-penalty: -2,
  mode: "global",
)

#let protein_pair_alignment = align-seq-pair(
  "MAVHLTPEEKS",
  "HLTPEE",
  matrix: "BLOSUM62",
  gap-penalty: -4,
  mode: "local",
)

```

The DNA example uses a custom match/mismatch model for a global alignment, while the protein example uses BLOSUM62 for a local alignment. The alignment scores are 7 and 28.

Rendering pairwise alignments

Pairwise alignments can be rendered using the `render-pair-alignment` function. The example below uses the local protein alignment from the previous section, showing the unaligned regions using a light gray color.

```

#render-pair-alignment(
  protein_pair_alignment.seq-1,
  protein_pair_alignment.seq-2,
  protein_pair_alignment.traceback-paths.at(0),
  unaligned-color: luma(75%),
)

```

A local protein alignment diagram showing two sequences: MGRHMTYPEEKS and HLT-PEE. The sequences are aligned as follows:

```

MGRHMTYPEEKS
| | | |
HLT-PEE

```

The first sequence has a light gray background, indicating unaligned regions. The second sequence has a white background with black text for aligned positions.

Figure 1. Local protein alignment with unaligned regions shown in light gray.

Dynamic programming matrix visualization

Dynamic programming is the core procedure used by the pairwise alignment algorithm: it fills a matrix of optimal scores for all prefix pairs of the two sequences, where each cell stores the best score achievable at that position and arrows indicate the traceback directions that can lead to an optimal alignment. The `render-dp-matrix` function renders the DP matrix of a given alignment, overlaying the traceback path used to produce the final alignment.

```
// Because there may be multiple optimal alignments, `traceback-paths`  

// is an array. We use `.at(0)` to visualize the path of the first alignment.  

#render-dp-matrix(  

  dna_pair_alignment.seq-1,  

  dna_pair_alignment.seq-2,  

  dna_pair_alignment.dp-matrix.values,  

  path: dna_pair_alignment.traceback-paths.at(0),  

  arrows: dna_pair_alignment.dp-matrix.arrows,  

)  

)
```

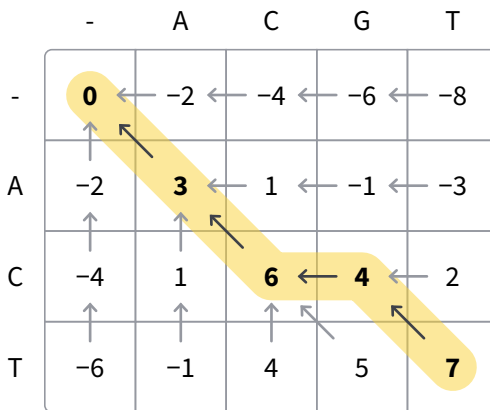


Figure 2. Dynamic programming matrix for the DNA alignment, with the optimal path highlighted.

Scoring matrices

Substitution matrices assign scores for aligning residues in pairwise and multiple sequence alignments, rewarding likely substitutions and penalizing unlikely ones. They represent substitution preferences as log-odds scores derived from observed evolutionary changes. In this way, they guide alignment algorithms by quantifying how plausible it is for one residue to replace another over time, helping produce biologically meaningful alignments. `genotypst` provides scoring matrices from the BLOSUM³ and PAM⁴ families for protein sequences, as well as the EDNAFULL matrix for DNA and RNA sequences.

Use `get-scoring-matrix` to retrieve a matrix data and `get-score-from-matrix` to query scores for specific residue pairs.

```
#let blosum62 = get-scoring-matrix("BLOSUM62")
#let ar_score = get-score-from-matrix(blosum62, "L", "D")
#text[The substitution score for L vs D in BLOSUM62 is: #ar_score.]
```

The substitution score for L vs D in BLOSUM62 is: -4.

You can render the entire scoring matrix using `render-scoring-matrix` function.

```
#render-scoring-matrix(blosum62)
```

Figure 3. BLOSUM62 scoring matrix.

Multiple sequence alignments

The `render-msa` function displays multiple sequence alignments with optional residue coloring and conservation bars.

In the example below:

- colors: true enables residue coloring based on biochemical properties.
 - conservation: true adds conservation bars above the alignment.
 - start: 100 and end: 160 limit the display to a specific region of interest (residues 100 to 160).

```
#let protein_msa = parse-fasta(read("/docs/data/msa.afa"))

#render-msa(
  protein_msa,
  start: 100,
  end: 160,
  colors: true,
  conservation: true,
)
```



Figure 4. MSA visualization for positions 100–160, with residue coloring and conservation bars enabled.

Residue coloring represents amino acid physicochemical properties. The sequence alphabet (amino acid, DNA, or RNA) is determined automatically and a suitable color palette is applied.

The bars above the alignment indicate the degree of conservation at each column.

Sequence logos

Sequence logos⁵ summarize conservation patterns within a sequence alignment and are commonly used to visualize binding sites, motifs, and functional domains. In a sequence logo, the total height of each stack represents the information content (in bits) at that position, while the height of individual letters reflects their relative frequencies.

In the example below, we visualize the same region as the MSA of the previous section (positions 100 to 160).

```
#render-sequence-logo(protein_msa, start: 100, end: 160)
```



Figure 5. Sequence logo for positions 100–160, showing conservation and residue frequency.

Like `render-msa`, `render-sequence-logo` automatically applies the appropriate color palette based on the sequence alphabet.

Color palettes

`genotypst` uses predefined color palettes to assign colors to sequence residues.

Amino acid palette

Amino acids are colored according to their physicochemical properties. Grouping residues by color helps reveal the chemical nature of conserved positions (e.g., whether a position is consistently hydrophobic or charged), which is often important for understanding protein structure, function, and evolution.

Hydrophobic

Alanine	Ala	A	#4d78ff
Histidine	His	H	#4d78ff
Isoleucine	Ile	I	#4d78ff
Leucine	Leu	L	#4d78ff
Methionine	Met	M	#4d78ff
Valine	Val	V	#4d78ff

Polar

Serine	Ser	S	#00c990
Threonine	Thr	T	#00c990
Glutamine	Gln	Q	#00c990
Asparagine	Asn	N	#00c990

Aromatic

Phenylalanine	Phe	F	#bac1d2
Tryptophan	Trp	W	#bac1d2
Tyrosine	Tyr	Y	#bac1d2

Negatively charged

Aspartic acid	Asp	D	#ff07b8
Glutamic acid	Glu	E	#ff07b8

Positively charged

Lysine	Lys	K	#e51f3e
Arginine	Arg	R	#e51f3e

Cysteine

Cysteine	Cys	C	#494e5b
----------	-----	---	---------

Glycine

Glycine	Gly	G	#f59116
---------	-----	---	---------

Proline

Proline	Pro	P	#dce100
---------	-----	---	---------

Nucleic acid palettes

The DNA and RNA palettes assign a distinct color to each nucleotide.

DNA palette

Adenine	A	#00c990
Cytosine	C	#4d78ff
Guanine	G	#ff07b8
Thymine	T	#f59116

RNA palette

Adenine	A	#00c990
Cytosine	C	#4d78ff
Guanine	G	#ff07b8
Uracil	U	#f59116

Visualizing genomic loci with genome maps

Genome maps enable visualization of the genes and other genomic elements within a locus, highlighting their order, orientation, and length. `genotypst` provides a `render-genome-map` function that produces a genome map from an array of dictionaries, each representing a genomic feature that will be plotted:

- `start` (required): Start coordinate.
- `end` (required): End coordinate.
- `strand`: Feature orientation (1 or "+" for the positive strand, -1 or "-" for the negative strand). `none` draws an undirected block.
- `label`: Feature label
- `color`: Fill color.

```
#let f_plasmid_locus = (
  (start: 65556, end: 66065, strand: -1, label: [_ygfA_]),
  (start: 66118, end: 66407, label: [_oriT_], color: rgb("#696975")),
  (start: 66479, end: 66862, strand: 1, label: [_traM_], color: rgb("#62B9F2")),
  (start: 66977, end: 67055, strand: -1, label: [_finP_]),
  (start: 67049, end: 67738, strand: 1, label: [_traJ_], color: rgb("#F7ED6C")),
  (start: 67837, end: 68232, strand: 1, label: [_traY_]),
  (start: 68265, end: 68630, strand: 1, label: [_traA_]),
  (start: 68645, end: 68956, strand: 1, label: [_traL_]),
  (start: 68978, end: 69544, strand: 1, label: [_traE_]),
)

#render-genome-map(
  f_plasmid_locus,
  coordinate-axis: true,
  unit: "bp",
)
```

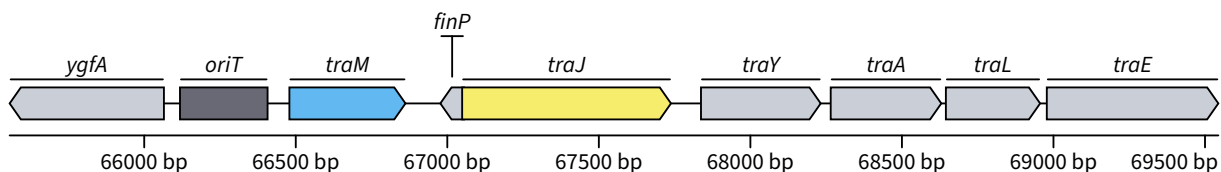


Figure 6. Genome map showing the genes within the 65,556–69,544 bp region of the F plasmid of *Escherichia coli* K-12 (GenBank: AP001918.1).

Working with phylogenetic trees

genotypst includes functions to parse and render phylogenetic trees. Trees can be created by parsing Newick-formatted strings with `parse-newick` or by manually constructing nested dictionary structures.

```
#let parsed_newick_tree = parse-newick(  
  "((Leaf A':0.2,'Leaf B':0.1)'Internal node':0.3,'Leaf C':0.6)Root;"  
)  
  
#let manual_tree = (  
  rooted: true,  
  name: "Root",  
  length: none,  
  children: (  
    (  
      name: "Internal node",  
      length: 0.3,  
      children: (  
        (name: "Leaf A", length: 0.2, children: none),  
        (name: "Leaf B", length: 0.1, children: none),  
      ),  
    ),  
    (name: "Leaf C", length: 0.6, children: none),  
  ),  
)
```

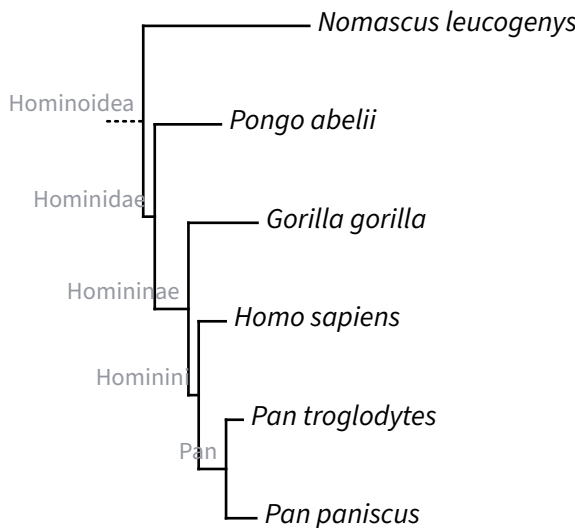
Visualizing trees

genotypst can produce visualizations of phylogenetic trees. To illustrate this, we will read and render a Newick file containing a phylogeny of the *Hominoidea* superfamily, which was extracted from the Ensembl Compara species tree⁶.

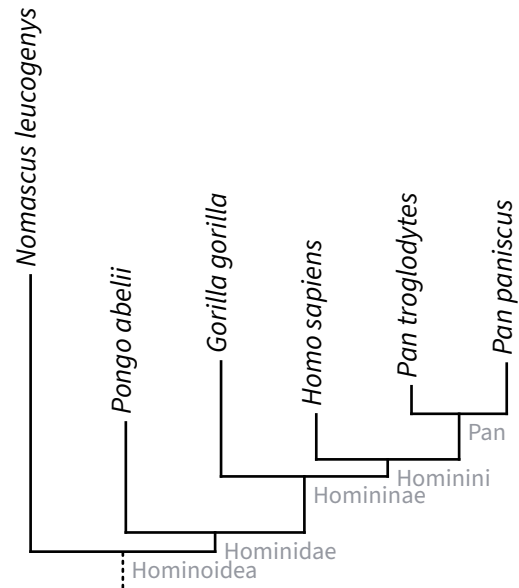
```
#let hominoidea_tree = parse-newick(read("/docs/data/hominoidea.nwk"))
```

To render the tree, use the `render-tree` function. By default, it produces a horizontal rectangular dendrogram, but a vertical layout can be specified using the `orientation: "vertical"` option.

```
#render-tree(hominoidea_tree, tip-label-italics: true, orientation:  
"horizontal")  
#render-tree(hominoidea_tree, tip-label-italics: true, orientation: "vertical")
```



Tree with horizontal orientation



Tree with vertical orientation

Customizing visualizations

Font selection

By default, `render-fasta` and `render-msa` inherit the monospaced font used for raw text in your document. To use a different font, wrap the rendering function in a context block with a custom font for raw text.

```

#let dna_msa = (
  "seq1": "AGTCTCAAGATAACTTCGAAACAACTTC",
  "seq2": "AGTTCCAAGTGGATTGGAATTGAACCTT",
  "seq3": "ACTCT-CGGATGGATTGGATACAAACTTT",
  "seq4": "AGTCT---GATTGATGTGGATACAAACTTC",
  "seq5": "AGTCT--GGGTGGATTGG-AACAAATTT",
  "seq6": "CAGTGCTCCCTGGTGGTGG-ACCATCTTAC",
  "seq7": "AGTCTCAAGACGGATACTG--ATGCCCTAT",
)

#context {
  show raw: set text(font: "Maple Mono")
  render-msa(dna_msa)
}

```

seq1	AGTCTCAAGATAACTTCGAAACAAACTTC	seq1	AGTCTCAAGATAACTTCGAAACAAACTTC
seq2	AGTTTCCAAGTGGATTGGATTGAACTTT	seq2	AGTTTCCAAGTGGATTGGATTGAACTTT
seq3	ACTCT-CGGATGGATTCGGATACAAACTTT	seq3	ACTCT-CGGATGGATTCGGATACAAACTTT
seq4	AGTCT---GATTGATGTGGATACAAACTTC	seq4	AGTCT---GATTGATGTGGATACAAACTTC
seq5	AGTCT--GGGTGGATTGG-AACAAATTTC	seq5	AGTCT--GGGTGGATTGG-AACAAATTTC
seq6	CAGTGCTCCCTGGTGGTGG-ACCATCTTAC	seq6	CAGTGCTCCCTGGTGGTGG-ACCATCTTAC
seq7	AGTCTCAAGACGGATACTG--ATGCCCTAT	seq7	AGTCTCAAGACGGATACTG--ATGCCCTAT

Default document font for raw text

Custom font (Maple Mono)

Pairwise alignments, dynamic programming matrices, scoring matrices, sequence logos, genome maps, and trees are rendered using the default document font, rather than the monospaced font for raw text. To specify a custom font for these visualizations, use a `show_text` rule instead. These renderers also expose styling parameters such as `unaligned-color`, `cell-size`, `path-color`, `arrow-color`, `color-map`, and `scale-limit`.

```
#context {
  show text: set text(font: "Libertinus Serif")
  render-pair-alignment(
    protein_pair_alignment.seq-1,
    protein_pair_alignment.seq-2,
    protein_pair_alignment.traceback-paths.at(0),
  )
}
```

M G R H M T Y P E E K S
| | | ||
H L T - P E E

Default document font

M G R H M T Y P E E K S
| | | ||
H L T - P E E

Custom font (Libertinus Serif)

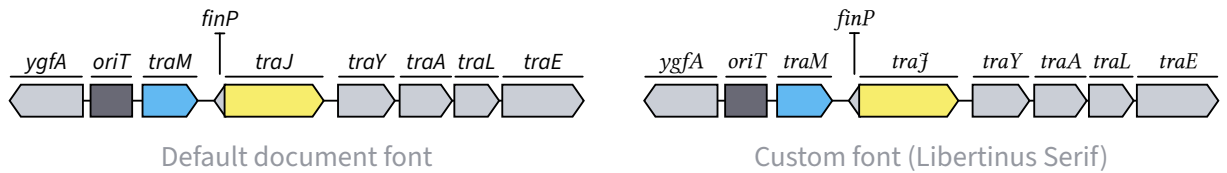
```
#context {
  show text: set text(font: "Libertinus Serif")
  render-sequence-logo(dna_msa)
}
```

Default document font

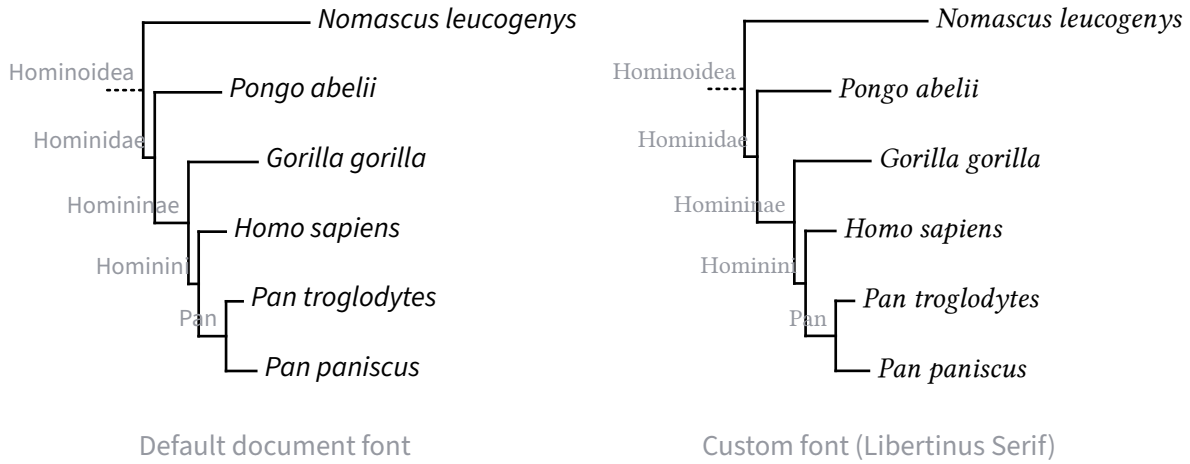
A G T c T C G A T G A T T G G A A A C A A A C T T T A C

Custom font (Libertinus Serif)

```
#context {
  show text: set text(font: "Libertinus Serif")
  render-genome-map(f_plasmid_locus)
}
```



```
#let hominoidea_tree = parse-newick(read("/docs/data/hominoidea.nwk"))
#context {
  show text: set text(font: "Libertinus Serif")
  #render-tree(hominoidea_tree, tip-label-italics: true)
}
```



Bibliography

1. Needleman, S. B. & Wunsch, C. D. A general method applicable to the search for similarities in the amino acid sequence of two proteins. *Journal of Molecular Biology* **48**, 443–453 (1970).

2. Smith, T. & Waterman, M. Identification of common molecular subsequences. *Journal of Molecular Biology* **147**, 195–197 (1981).
3. Henikoff, S. & Henikoff, J. G. Amino acid substitution matrices from protein blocks. *Proceedings of the National Academy of Sciences* **89**, 10915–10919 (1992).
4. Dayhoff, M., Schwartz, R. & Orcutt, B. A model of evolutionary change in proteins. *Atlas of protein sequence and structure* (1979).
5. Schneider, T. D. & Stephens, R. Sequence logos: a new way to display consensus sequences. *Nucleic Acids Research* **18**, 6097–6100 (1990).
6. Herrero, J. *et al.* Ensembl comparative genomics resources. *Database* **2016**, bav96 (2016).