

# Package: glyparse (via r-universe)

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**Title** Parsing Glycan Structure Text Representations

**Version** 0.6.0

**Description** Provides functions to parse glycan structure text representations into 'glyrepr' glycan structures. Currently, it supports StrucGP-style, pGlyco-style, IUPAC-condensed, IUPAC-extended, IUPAC-short, WURCS, Linear Code, and GlycoCT format. It also provides an automatic parser to detect the format and parse the structure string.

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**Suggests** knitr, rmarkdown, glue, testthat (>= 3.0.0)

**Config/testthat/edition** 3

**Encoding** UTF-8

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**RoxygenNote** 7.3.3

**URL** <https://glycoverse.github.io/glyparse/>,

<https://github.com/glycoverse/glyparse>

**Imports** checkmate, cli, dplyr, glyrepr (>= 0.9.0), igrph, purrr, rlang, rstackdeque, stringr, vctrs

**Depends** R (>= 4.1)

**VignetteBuilder** knitr

**BugReports** <https://github.com/glycoverse/glyparse/issues>

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**Repository** <https://glycoverse.r-universe.dev>

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auto_parse	<i>Automatic Structure Parsing</i>
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### Description

Detect the structure string type and use the appropriate parser to parse automatically. Mixed types are supported.

Supported types:

1. GlycoCT
2. IUPAC-condensed
3. IUPAC-extended
4. IUPAC-short
5. WURCS
6. Linear Code
7. pGlyco
8. StrucGP

### Usage

```
auto_parse(x, on_failure = "error")
```

### Arguments

x	A character vector of structure strings. NA values are allowed and will be returned as NA structures.
on_failure	How to handle parsing failures. "error" aborts when a structure cannot be parsed. "na" returns NA at invalid positions.

### Value

A `glyrepr::glycan_structure()` object.

## Examples

```
# Single structure
x <- "Gal(b1-3)GlcNAc(b1-4)Glc(a1-)" # IUPAC-condensed
auto_parse(x)

# Mixed types
x <- c(
  "Gal(b1-3)GlcNAc(b1-4)Glc(a1-)", # IUPAC-condensed
  "Neu5Aca3Gal3(Fuca6)GlcNAcb-" # IUPAC-short
)
auto_parse(x)
```

---

parse\_glycoct

*Parse GlycoCT Structures*

---

## Description

This function parses GlycoCT strings into a `glyrepr::glycan_structure()`. GlycoCT is a format used by databases like GlyTouCan and GlyGen.

## Usage

```
parse_glycoct(x, on_failure = "error")
```

## Arguments

<code>x</code>	A character vector of GlycoCT strings. NA values are allowed and will be returned as NA structures.
<code>on_failure</code>	How to handle parsing failures. "error" aborts when a structure cannot be parsed. "na" returns NA at invalid positions.

## Details

GlycoCT format consists of two parts:

- RES: Contains monosaccharides (lines starting with 'b:') and substituents (lines starting with 's:')
- LIN: Contains linkage information between residues

For more information about GlycoCT format, see the `glycoct.md` documentation.

## Value

A `glyrepr::glycan_structure()` object.

## Examples

```
glycoct <- paste0(
  "RES\n",
  "1b:a-dgal-HEX-1:5\n",
  "2s:n-acetyl\n",
  "3b:b-dgal-HEX-1:5\n",
  "LIN\n",
  "1:1d(2+1)2n\n",
  "2:1o(3+1)3d"
)
parse_glycoct(glycoct)
```

---

parse\_iupac\_condensed *Parse IUPAC-condensed Structures*

---

## Description

This function parses IUPAC-condensed strings into a `glyrepr:glycan_structure()`. For more information about IUPAC-condensed notation, see [doi:10.1351/pac199668101919](https://doi.org/10.1351/pac199668101919).

## Usage

```
parse_iupac_condensed(x, on_failure = "error")
```

## Arguments

<code>x</code>	A character vector of IUPAC-condensed strings. NA values are allowed and will be returned as NA structures.
<code>on_failure</code>	How to handle parsing failures. "error" aborts when a structure cannot be parsed. "na" returns NA at invalid positions.

## Details

The IUPAC-condensed notation is a compact form of IUPAC-extended notation. It is used by the [GlyConnect](#) database. It contains the following information:

- Monosaccharide name, e.g. "Gal", "GlcNAc", "Neu5Ac".
- Substituent, e.g. "9Ac", "4Ac", "3Me", "?S".
- Linkage, e.g. "b1-3", "a1-2", "a1-?".

An example of IUPAC-condensed string is "Gal(b1-3)GlcNAc(b1-4)Glc(a1-".

The reducing-end monosaccharide can be with or without anomer information. For example, the two strings below are all valid:

- "Neu5Ac(a2-"
- "Neu5Ac"

In the first case, the anomer is "a2". In the second case, the anomer is "?2".

**Value**

A `glyrepr::glycan_structure()` object.

**See Also**

`parse_iupac_short()`, `parse_iupac_extended()`

**Examples**

```
iupac <- "Gal(b1-3)GlcNAc(b1-4)Glc(a1-"  
parse_iupac_condensed(iupac)
```

---

parse\_iupac\_extended *Parse IUPAC-extended Structures*

---

**Description**

Parse IUPAC-extended-style structure characters into a `glyrepr::glycan_structure()`. For more information about IUPAC-extended format, see [doi:10.1351/pac199668101919](https://doi.org/10.1351/pac199668101919).

**Usage**

```
parse_iupac_extended(x, on_failure = "error")
```

**Arguments**

<code>x</code>	A character vector of IUPAC-extended strings. NA values are allowed and will be returned as NA structures.
<code>on_failure</code>	How to handle parsing failures. "error" aborts when a structure cannot be parsed. "na" returns NA at invalid positions.

**Details**

The function accepts both a Unicode format (using the Greek letters alpha/beta and the arrow symbol ->) and a plain-text format (using the strings "alpha", "beta", and "->"). For example, both "`\u03b2-D-Galp-(1\u21923)-\u03b1-D-GalpNAc-(1\u21922`" and "`beta-D-Galp-(1->3)-alpha-D-GalpNAc-(1->`" are valid inputs.

**Value**

A `glyrepr::glycan_structure()` object.

**See Also**

`parse_iupac_condensed()`, `parse_iupac_short()`

## Examples

```
iupac <- "\u03b2-D-Galp-(1\u21923)-\u03b2-D-GalpNAc-(1\u2192)"
parse_iupac_extended(iupac)
parse_iupac_extended("beta-D-Galp-(1->3)-alpha-D-GalpNAc-(1->")
```

---

parse\_iupac\_short      *Parse IUPAC-short Structures*

---

## Description

Parse IUPAC-short-style structure characters into a `glyrepr::glycan_structure()`. For more information about IUPAC-short format, see [doi:10.1351/pac199668101919](https://doi.org/10.1351/pac199668101919).

## Usage

```
parse_iupac_short(x, on_failure = "error")
```

## Arguments

x	A character vector of IUPAC-short strings. NA values are allowed and will be returned as NA structures.
on_failure	How to handle parsing failures. "error" aborts when a structure cannot be parsed. "na" returns NA at invalid positions.

## Details

The IUPAC-short notation is a compact form of IUPAC-condensed notation. It is rarely used in database, but appears a lot in literature for its conciseness. Compared with IUPAC-condensed notation, IUPAC-short notation ignore the anomer positions, assuming they are known for common monosaccharides. For example, "Neu5Aca3Gala-" assumes the anomer of Neu5Ac is C2 (a2-3 linked). Also, the parentheses around linkages are omitted, and parentheses are used to indicate branching, e.g. "Neu5Aca3Gala3(Fuca3)GlcNAcb-".

In the first case, the anomer is "a2". In the second case, the anomer is "?2".

## Value

A `glyrepr::glycan_structure()` object.

## See Also

`parse_iupac_condensed()`, `parse_iupac_extended()`

## Examples

```
iupac <- "Neu5Aca3Gala3(Fuca6)GlcNAcb-"
parse_iupac_short(iupac)
```

---

parse\_linear\_code      *Parse Linear Code Structures*

---

### Description

Parse Linear Code structures into a `glyrepr::glycan_structure()`. To know more about Linear Code, see [this article](#).

### Usage

```
parse_linear_code(x, on_failure = "error")
```

### Arguments

x	A character vector of Linear Code strings. NA values are allowed and will be returned as NA structures.
on_failure	How to handle parsing failures. "error" aborts when a structure cannot be parsed. "na" returns NA at invalid positions.

### Value

A `glyrepr::glycan_structure()` object.

### Examples

```
linear_code <- "Ma3(Ma6)Mb4GNb4GNb"  
parse_linear_code(linear_code)
```

---

parse\_pglyco\_struct      *Parse pGlyco Structures*

---

### Description

Parse pGlyco-style structure characters into a `glyrepr::glycan_structure()`. See example below for the structure format.

### Usage

```
parse_pglyco_struct(x, on_failure = "error")
```

### Arguments

x	A character vector of pGlyco-style structure strings. NA values are allowed and will be returned as NA structures.
on_failure	How to handle parsing failures. "error" aborts when a structure cannot be parsed. "na" returns NA at invalid positions.

**Value**

A `glyrepr::glycan_structure()` object.

**Examples**

```
glycan <- parse_pglyco_struc("(N(F)(N(H(H(N))(H(N(H))))))")  
print(glycan, verbose = TRUE)
```

---

parse\_strucgp\_struc    *Parse StrucGP Structures*

---

**Description**

Parse StrucGP-style structure characters into a `glyrepr::glycan_structure()`. See example below for the structure format.

**Usage**

```
parse_strucgp_struc(x, on_failure = "error")
```

**Arguments**

<code>x</code>	A character vector of StrucGP-style structure strings. NA values are allowed and will be returned as NA structures.
<code>on_failure</code>	How to handle parsing failures. "error" aborts when a structure cannot be parsed. "na" returns NA at invalid positions.

**Value**

A `glyrepr::glycan_structure()` object.

**Examples**

```
glycan <- parse_strucgp_struc("A2B2C1D1E2F1fedD1E2edcbB5ba")  
print(glycan, verbose = TRUE)
```

---

`parse_wurcs`*Parse WURCS Structures*

---

**Description**

This function parses WURCS strings into a `glyrepr::glycan_structure()`. Currently, only WURCS 2.0 is supported. For more information about WURCS, see [WURCS](#).

**Usage**

```
parse_wurcs(x, on_failure = "error")
```

**Arguments**

<code>x</code>	A character vector of WURCS strings. NA values are allowed and will be returned as NA structures.
<code>on_failure</code>	How to handle parsing failures. "error" aborts when a structure cannot be parsed. "na" returns NA at invalid positions.

**Value**

A `glyrepr::glycan_structure()` object.

**Examples**

```
wurcs <- paste0(
  "WURCS=2.0/3,5,4/",
  "[a2122h-1b_1-5_2*NCC/3=0][a1122h-1b_1-5][a1122h-1a_1-5]/",
  "1-1-2-3-3/a4-b1_b4-c1_c3-d1_c6-e1"
)
parse_wurcs(wurcs)
```

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