

¹ Peptacular: A Python package for amino acid sequence analysis

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⁵ Summary

⁶ Mass spectrometry-based proteomics relies on computational methods to identify and
⁷ characterize amino acid (AA) sequences. These sequences, which range from short peptides to
⁸ complete proteins, exhibit considerable chemical complexity arising from post-translational
⁹ modifications (PTMs), variable charge states, neutral losses, and isotopic patterns ([Angel et](#)
¹⁰ [al., 2012; Smith & Kelleher, 2013](#)). **Peptacular** is a fully type-annotated pure Python library
¹¹ designed around ProForma 2.1 notation for working with such complexity. The library provides
¹² functionality for modifying AA sequences, calculating mass, m/z, isotopic distributions,
¹³ physicochemical properties, digesting via common enzymes, fragmenting, and more.

¹⁴ Statement of Need

¹⁵ Historically, the proteomics field has lacked standardization for representing AA sequences.
¹⁶ Individual software tools have implemented proprietary notations, which has created barriers to
¹⁷ data integration and reanalysis across platforms. ProForma notation ([LeDuc et al., 2018](#)) was
¹⁸ developed to address this challenge by providing a unified representation system. However,
¹⁹ adoption has remained limited, partly due to insufficient support in widely-used computational
²⁰ tools and libraries. **Peptacular** addresses this gap by providing an accessible and efficient
²¹ application programming interface (API) for working with AA sequences in a standardized
²² manner. The library was built from the ground up to support ProForma notation and implements
²³ nearly all features specified in ProForma 2.1.

²⁴ Modern proteomics experiments routinely identify hundreds of thousands of peptide-spectrum
²⁵ matches (PSMs). These results are typically exported as tabular files and processed using
²⁶ Python, particularly with data manipulation libraries such as **pandas** ([Team, 2025](#)) and **polars**
²⁷ ([Vink et al., 2025](#)). Given the scale of these datasets, efficient computational methods for
²⁸ processing these tables are essential. **Peptacular** addresses this requirement by providing a
²⁹ functional API that operates directly on tabular data structures while automatically supporting
³⁰ parallelization to improve computational performance.

³¹ State of the Field

³² Several existing Python packages provide functionality for working with AA sequences and
³³ ProForma parsing. **Pyteomics** ([Goloborodko et al., 2013](#)) offers considerable overlap with
³⁴ **Peptacular** but does not support the complete ProForma specification and lacks built-in
³⁵ parallelization capabilities. **BioPython** ([Cock et al., 2009](#)) provides basic support for calculating
³⁶ properties of AA sequences; however, its broader scope (encompassing DNA and RNA) results
³⁷ in a more complex API, and it does not include ProForma support. **RustyMS** ([Schulte et al.,](#)
³⁸ [n.d.](#)) Python bindings offer comprehensive ProForma support with the performance benefits
³⁹ of a Rust backend, though this architecture introduces additional API complexity and omits

40 certain features available in Peptacular. PyOpenMS (Röst et al., 2013) supports AA sequence
41 operations but lacks ProForma compatibility, requires specific Python versions, and addresses
42 a substantially broader scope than AA sequence manipulation alone.

43 The development of Peptacular was motivated by the need for a package that combines
44 comprehensive AA sequence support with ProForma notation in a pure Python implementation
45 compatible with any Python environment. The decision to implement this library from
46 scratch, rather than extending existing projects, was based on several considerations. First,
47 many established packages have expanded to support various features beyond processing AA
48 sequences, making targeted contributions more challenging. Second, because many of these
49 projects predate ProForma notation, ProForma support was incorporated retrospectively, which
50 constrains the extent of implementation and complicates API design. Finally, not all current
51 packages are compatible with the latest Python versions, particularly free-threaded versions of
52 Python with the Global Interpreter Lock (GIL) disabled.

53 Software Design

54 Peptacular provides two primary APIs: a functional API and an object-oriented API. The
55 object-oriented API employs a factory pattern to modify Annotation objects, enabling precise
56 control over annotations. The functional API provides functions that operate directly on
57 serialized sequences and annotations, with automatic parallelization when multiple inputs are
58 provided.

59 The built-in parallelization supports three execution backends: sequential, threaded, and
60 process-based. Sequential execution runs on a single core. Threading is limited by the GIL in
61 most Python versions; however, as Python development progresses toward making the GIL
62 optional, threading will likely become increasingly viable. Process-based parallelization is the
63 default backend, as it has the best compatibility between systems. Users can globally configure
64 the spawning mechanism for processes or threads, selecting from fork, spawn, or forkserver
65 modes. Processes are cached to avoid the startup cost of creating new processes.

66 Peptacular employs lazy evaluation to minimize memory overhead and improve performance.
67 Modifications remain in serialized form until they are explicitly required for calculations. The
68 serialization and parsing of objects utilize extensive caching, as proteomics datasets commonly
69 contain only a subset of repeated modifications. Additionally, Peptacular uses conditional
70 initialization, whereby modification-related data structures are instantiated only when the
71 corresponding modification type is needed. This approach reduces the memory footprint of
72 annotation objects and accelerates their creation and copying.

73 Modification reference data, including masses, compositions, and identifiers from Unimod
74 (Creasy & Cottrell, 2004), PSI-MOD (Hupo-Psi, n.d.-a), RESID (*RESID Database [PIR -*
75 *Protein Information Resource]*, n.d.), XLMod (Hupo-Psi, n.d.-b), and GNOme (Glygen-Glycan-
76 Data, n.d.), are provided by the Tacular (P. Garrett, 2026) package. This package embeds
77 the data directly within itself as Python modules rather than storing them as external files.
78 Only valid modifications are included in the embedded data; a modification is considered valid
79 if it possesses at least one of the following properties: average mass, monoisotopic mass, or
80 chemical formula. This design eliminates file I/O overhead during the parsing of supported
81 ontologies. The primary limitation of this approach is that updates to the ontologies are not
82 immediately available and require a package update. However, Tacular includes a pipeline to
83 rebuild from the latest ontology versions.

84 Peptacular is fully type-annotated and includes a `py.typed` marker, enabling static type checking
85 with tools. This provides enhanced IDE support through intelligent autocomplete and inline
86 documentation, while allowing users to catch type-related errors before runtime. The package
87 maintains >70% test coverage and employs continuous integration via GitHub Actions.

88 Research impact statement

89 Since its initial release, Peptacular has demonstrated measurable adoption. The package has
 90 accumulated over 33k downloads from PyPI, with sustained weekly download rates exceeding
 91 200 installations. It has been recognized as one of 3 python packages to support Proforma
 92 notation by the PSI group. Additionally, Peptacular was used to generate figures within a
 93 textbook chapter ([P. T. Garrett et al., 2025](#)).

94 Example Usage

95 Object-Based API

```
import peptacular as pt

# Parse a sequence into a ProFormaAnnotation
peptide: pt.ProFormaAnnotation = pt.parse("PEM[Oxidation]TIDE")

# Calculate mass and m/z
mass: float = peptide.mass() # 849.342
mz: float = peptide.mz(charge=2) # 425.678

# Factory pattern
print(peptide.set_charge(2).set_peptide_name("Peptacular").serialize())
# (>Peptacular)PEM[Oxidation]TIDE/2
```

96 Functional-Based API

```
import peptacular as pt

peptides = ['[Acetyl]-PEPTIDES', '<C13>ARE', 'SICK/2']

# Calculate mass and m/z for all peptides
masses: list[float] = pt.mass(peptides) # [928.4026, 374.1914, 451.2454]
mzs: list[float] = pt.mz(peptides, charge=2) # [465.2086, 188.103, 225.6227]
```

97 Mathematics

98 Peptacular implements algorithms for molecular mass calculations and isotopic pattern
 99 prediction. The following sections formalize the mathematical framework underlying these
 100 calculations.

101 Base Mass

102 The base mass M_{base} of a peptide sequence with modifications is calculated as the sum of all
 103 constituent components:

$$104 M_{base} = \sum_{i=1}^n m_i + M_N + M_C + M_S + M_I + M_R + M_U + \mathbb{1}_{\text{precursor}} \cdot M_L$$

105 where:

- 106 ■ n is the sequence length
- 107 ■ m_{AA_i} is the monoisotopic (or average) mass of amino acid at position i
- 108 ■ M_N is the total mass of N-terminal modifications
- 109 ■ M_C is the total mass of C-terminal modifications
- 110 ■ M_S is the total mass of static/fixed modifications

- 111 ▪ M_I is the total mass of position-specific modifications
 - 112 ▪ M_R is the total mass of modifications within defined sequence intervals
 - 113 ▪ M_U is the total mass of modifications with unknown positions
 - 114 ▪ M_L is the total mass of labile modifications
 - 115 ▪ $\mathbb{1}_{\text{precursor}}$ is an indicator function: 1 for precursor ions, 0 for fragment ions
- 116 Labile modifications are only included in precursor ion mass calculations and excluded from all
 117 fragment ion types.

118 Neutral Mass

119 The neutral mass M_{neutral} of a fragment ion is calculated by combining the base mass with
 120 ion-type, isotope modifications, neutral deltas.

$$121 \quad M_{\text{neutral}} = M_{\text{base}} + M_{\text{ion}} + M_{\text{isotope}} + M_{\text{ndelta}}$$

122 where:

- 123 ▪ M_{base} is the peptide base mass from the previous section
- 124 ▪ M_{ion} is the ion-type-specific mass offset
- 125 ▪ M_{isotope} is the mass shift from a specific isotopic species
- 126 ▪ M_{ndelta} is the mass change from neutral losses/gains

127 Mass-to-charge Ratio

128 The mass-to-charge ratio is calculated by incorporating charge carriers and electron mass
 129 corrections to the neutral mass:

$$130 \quad \frac{m}{z} = \frac{M_{\text{neutral}} + M_{\text{adduct}} - z \cdot m_e}{z}$$

131 where:

- 132 ▪ M_{neutral} is the neutral fragment mass
- 133 ▪ M_{adduct} is the total mass of charge carriers
- 134 ▪ z is the total charge state
- 135 ▪ $m_e = 0.0005485799$ Da (electron mass)

136 Isotopic Distribution

137 The isotopic distribution of a peptide is calculated by convolving the isotopic patterns of all
 138 constituent elements. For a peptide with elemental composition $\{E_1 : n_1, E_2 : n_2, \dots, E_k : n_k\}$,
 139 the isotopic distribution is:

$$140 \quad P(\text{total}) = P(E_1)^{n_1} \otimes P(E_2)^{n_2} \otimes \dots \otimes P(E_k)^{n_k}$$

141 where $P(E_i)$ is the natural isotopic distribution of element E_i , n_i is the count of that element,
 142 and \otimes represents the convolution operation.

143 Averagine Model

144 When the exact elemental composition is unknown, the averagine model estimates composition
 145 from molecular mass using empirically-derived ratios. The composition is calculated as:

$$146 \quad n_E = r_E \cdot M_{\text{neutral}} + n_{E,\text{ion}}$$

147 where:

- 148 ▪ n_E is the estimated count of element E
- 149 ▪ r_E is the averagine ratio (atoms per dalton) for element E
- 150 ▪ M_{neutral} is the neutral peptide mass
- 151 ▪ $n_{E,\text{ion}}$ is the elemental contribution from the ion type

¹⁵² The averagine ratios (atoms/Da) derived from the human proteome are:

- ¹⁵³ ▪ C: 0.044179
- ¹⁵⁴ ▪ H: 0.069749
- ¹⁵⁵ ▪ N: 0.012344
- ¹⁵⁶ ▪ O: 0.013352
- ¹⁵⁷ ▪ S: 0.000400

¹⁵⁸ Figures

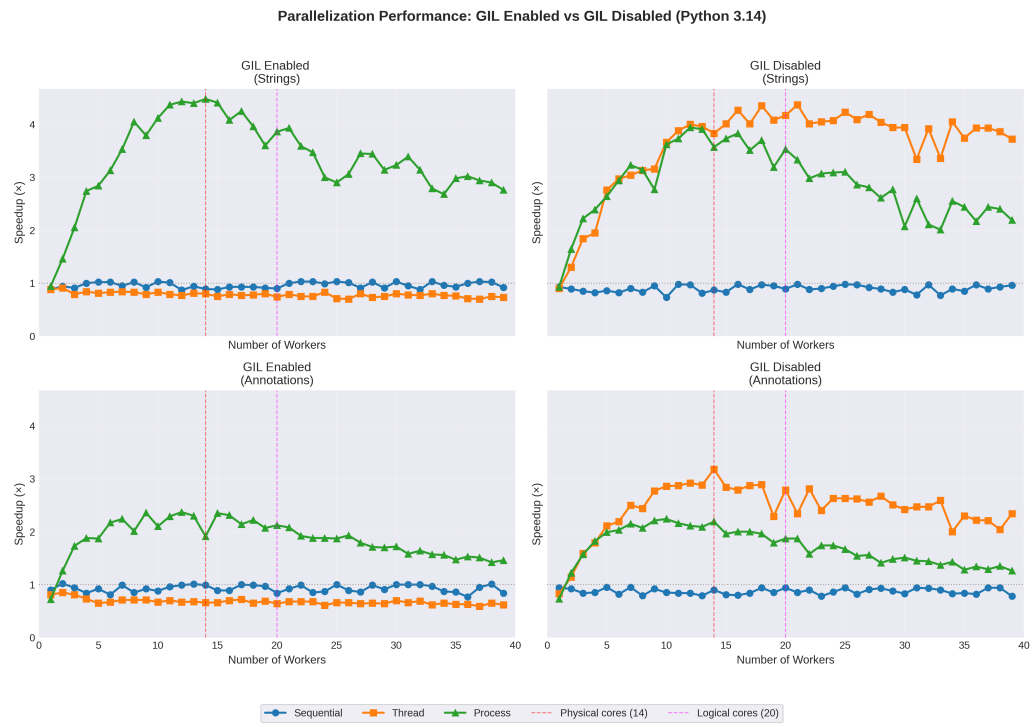
¹⁵⁹ **Table 1: Proforma 2.1 Compliance**

? Feature	Example	\$
Y Amino acids (+UO)	AAHCFKUOT	6.1
Y Unimod names	PEM[Oxidation]AT	6.2.1
Y PSI-MOD names	PEM[monohydroxylated residue]AT	6.2.1
Y Unimod numbers	PEM[UNIMOD:35]AT	6.2.2
Y PSI-MOD numbers	PEM[MOD:00425]AT	6.2.2
Y Delta masses	PEM[+15.995]AT	6.2.3
Y N-terminal modifications	[Carbamyl]-QPEPTIDE	6.3
Y C-terminal modifications	PEPTIDE-[Methyl]	6.3
Y Labile modifications	{Glycan:Hex}EM[U:Oxidation]EV	6.4
Y Multiple modifications	MPGNW[Oxidation][Carboxymethyl]PESQE	6.5
Y Information tag	ELV[INFO:AnyString]IS	6.6
Y Ambiguous amino acids	BZJX	7.1
Y Prefixed delta masses	PEM[U:+15.995]AT	7.2
Y Mass gap	PEX[+147.035]AT	7.3
Y Formulas	PEM[Formula:0]AT, PEM[Formula:[1701]]AT	7.4
Y Mass with interpretation	PEM[+15.995\ Oxidation]AT	7.5
Y Unknown mod position	[Oxidation]?PEMAT	7.6.1
Y Set of positions	PEP[Oxidation#1]M[#1]AT	7.6.2
Y Range of positions	PRT(ESFRMS)[+19.0523]ISK	7.6.3
Y Position scores	PEP[Oxidation#1(0.95)]M[#1(0.05)]AT	7.6.4
Y Range position scores	(PEP)[Oxidation#1(0.95)]M[#1(0.05)]AT	7.6.5
Y Amino acid ambiguity	(?VCH)AT	7.7
Y Modification prefixes	PEPM[U:Oxidation]AS[M:0-phospho-L-serine]	7.8
Y RESID modifications	EM[R:L-methionine sulfone]EM[RESID:AA0581]	8.1
Y Names	(>Heavy chain)EVQLVESG	8.2
Y XL-MOD modifications	EVT[X:Aryl azide]LEK[XLMOD:00114]SEFD	9.1
N Cross-linkers (intrachain)	EVT[X:Aryl azide#XL1]LEK[#XL1]SEFD	9.2.1
N Cross-linkers (interchain)	EVT[X:Aryl azide#XL1]L//EK[#XL1]SEFD	9.2.2
N Branches	ED[MOD:00093#BRANCH]//D[#BRANCH]ATR	9.3
Y GNO modifications	NEEYN[GNO:G59626AS]K	10.1
Y Glycan compositions	NEEYN[Glycan:Hex5HexNAc4NeuAc1]K	10.2
Y Charged formulas	SEQUEN[Formula:Zn1:z+2]CE	11.1
Y Controlling placement	PTI(MERMERME)[+32\ Position:E]PTIDE	11.2
Y Global isotope	<13C>CARBON	11.3.1
Y Fixed modifications	<[Oxidation]@M>ATPEMILTCMGCLK	11.3.2
Y Chimeric spectra	NEEYN+SEQUEN	11.4
Y Charges	SEQUEN/2, SEQUEN/[Na:z+1,H:z+1]	11.5
N Ion notation	SEQUEN-[b-type-ion]	11.6

¹⁶⁰ **Table 1** depicts the level of ProForma support in Peptacular. The package currently supports

161 all ProForma 2.1 features for linear peptides. Cross-linked peptides (both inter- and intrachain)
 162 and branched structures are not supported. Ion notation is also not supported, as it conflicts
 163 with mzPAF (Peak Annotation Format) notation for fragment ions.

164 **Figure 1: Parallelization Performance - GIL Enabled vs GIL Disabled (Python 3.14t)**



165 **Figure 1:** Parallelization performance comparison for calculating the mass of randomly generated modified peptides with lengths ranging from 10 to 30 amino acids. The benchmark compares serialized annotations (strings) and annotation objects across different parallelization methods, varying numbers of workers, and both GIL-enabled and GIL-disabled configurations. The baseline for speedup calculations is single-worker sequential-based execution ($0.336s \pm 0.011s$ for serialized strings, $0.178s \pm 0.004s$ for annotation objects).
 166 Benchmark environment: Intel i7-12700H (14 cores, 20 threads), 64GB RAM, Python 3.14t.

165 **AI usage disclosure**

166 Generative AI models were employed to support the development of this software package.
 167 Specifically, Claude Sonnet 4.5, Gemini 2.0 Pro, and GitHub Copilot's autocomplete extension
 168 were utilized for code generation, test development, debugging assistance, and documentation
 169 preparation. These tools were accessed through the Copilot extension in Visual Studio Code.
 170 Additionally, Type.ai was used to assist in manuscript preparation. All AI-generated content
 171 was subsequently reviewed and verified for accuracy.

172 **Availability**

173 Peptacular is distributed through PyPI (<https://pypi.org/project/peptacular/>) and available
 174 as open-source software on GitHub (<https://github.com/tacular-omics/peptacular>).
 175 Documentation is accessible at <https://peptacular.readthedocs.io>. The software is released
 176 under the MIT license.

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