
PyBEL Documentation

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GETTING STARTED

1 Overview	3
1.1 Background on Systems Biology Modeling	3
1.2 Design Considerations	3
1.3 Implementation	4
1.4 Extensions to BEL	4
1.5 Things to Consider	5
2 Installation	9
2.1 Extras	9
2.2 Caveats	10
2.3 Upgrading	10
3 Data Model	11
3.1 Constants	11
3.2 Graph	12
3.3 Nodes	21
3.4 Unqualified Edges	28
3.5 Edges	30
4 Example Networks	35
5 Summary	39
6 Operations	45
7 Filters	47
8 Transformations	49
9 Grouping	51
10 Pipeline	53
10.1 Transformation Decorators	55
10.2 Exceptions	56
11 Input and Output	57
11.1 Import	58
11.2 Transport	60
11.3 Visualization	68
11.4 Analytical Services	69
11.5 Web Services	72

11.6 Databases	76
11.7 Lossy Export	77
12 Manager	79
12.1 Manager API	79
12.2 Manager Components	81
13 Models	85
14 Cookbook	93
14.1 Configuration	93
15 Command Line Interface	95
15.1 pybel	95
16 Constants	103
17 Parsers	111
17.1 BEL Parser	111
17.2 Metadata Parser	115
17.3 Control Parser	118
17.4 Concept Parser	120
17.5 Sub-Parsers	122
18 Internal Domain Specific Language	123
18.1 Primitives	123
18.2 Named Entities	124
18.3 Central Dogma	125
18.4 Fusions	130
18.5 Utilities	133
19 Logging Messages	135
19.1 Errors	135
19.2 Parse Exceptions	135
20 References	145
20.1 Related Publications	145
20.2 Software using PyBEL	145
20.3 BEL Content	146
21 Roadmap	147
21.1 PyBEL	147
21.2 Bio2BEL	147
21.3 Data2BEL	148
21.4 PyBEL Tools	148
21.5 BEL Commons	148
22 Current Issues	149
22.1 Speed	149
22.2 Namespaces	149
22.3 Testing	149
23 Technology	151
23.1 Versioning	151
23.2 Testing in PyBEL	151
23.3 Deployment	152

24 Indices and Tables	155
Bibliography	157
Python Module Index	159
Index	161

Biological Expression Language (BEL) is a domain-specific language that enables the expression of complex molecular relationships and their context in a machine-readable form. Its simple grammar and expressive power have led to its successful use in the to describe complex disease networks with several thousands of relationships.

PyBEL is a pure Python software package that parses BEL documents, validates their semantics, and facilitates data interchange between common formats and database systems like JSON, CSV, Excel, SQL, CX, and Neo4J. Its companion package, [PyBEL-Tools](#), contains a library of functions for analysis of biological networks. For result-oriented guides, see the [PyBEL-Notebooks](#) repository.

Installation is as easy as getting the code from [PyPI](#) with `python3 -m pip install pybel`. See the *installation* documentation.

For citation information, see the [references](#) page.

PyBEL is tested on Python 3.5+ on Mac OS and Linux using [Travis CI](#) as well as on Windows using [AppVeyor](#).

See also:

- Specified by [BEL 1.0](#) and [BEL 2.0](#)
- Documented on [Read the Docs](#)
- Versioned on [GitHub](#)
- Tested on [Travis CI](#)
- Distributed by [PyPI](#)

OVERVIEW

1.1 Background on Systems Biology Modeling

1.1.1 Biological Expression Language (BEL)

Biological Expression Language (BEL) is a domain specific language that enables the expression of complex molecular relationships and their context in a machine-readable form. Its simple grammar and expressive power have led to its successful use to describe complex disease networks with several thousands of relationships. For a detailed explanation, see the [BEL 1.0](#) and [2.0](#) specifications.

1.1.2 OpenBEL Links

- [OpenBEL on Google Groups](#)
- [OpenBEL Wiki](#)
- [OpenBEL on GitHub](#)
- [Chat on Gitter](#)

1.2 Design Considerations

1.2.1 Missing Namespaces and Improper Names

The use of openly shared controlled vocabularies (namespaces) within BEL facilitates the exchange and consistency of information. Finding the correct namespace:name pair is often a difficult part of the curation process.

1.2.2 Outdated Namespaces

OpenBEL provides a variety of namespaces covering each of the BEL function types. These namespaces are generated by code found at <https://github.com/OpenBEL/resource-generator> and distributed at <http://resources.openbel.org/belframework/>.

This code has not been maintained to reflect the changes in the underlying resources, so this repository has been forked and updated at <https://github.com/pybel/resource-generator> to reflect the most recent versions of the underlying namespaces. The files are now distributed using the Fraunhofer SCAI [Artifactory](#) server.

1.2.3 Generating New Namespaces

In some cases, it is appropriate to design a new namespace, using the [custom namespace specification](#) provided by the OpenBEL Framework. Packages for generating namespace, annotation, and knowledge resources have been grouped in the [Bio2BEL](#) organization on GitHub.

1.2.4 Synonym Issues

Due to the huge number of terms across many namespaces, it's difficult for curators to know the domain-specific synonyms that obscure the controlled/preferred term. However, the issue of synonym resolution and semantic searching has already been generally solved by the use of ontologies. Besides just a controlled vocabulary, they also a hierarchical model of knowledge, synonyms with cross-references to databases and other ontologies, and other information semantic reasoning. Ontologies in the biomedical domain can be found at [OBO](#) and [EMBL-EBI OLS](#).

Additionally, as a tool for curators, the EMBL Ontology Lookup Service (OLS) allows for semantic searching. Simple queries for the terms 'mitochondrial dysfunction' and 'amyloid beta-peptides' immediately returned results from relevant ontologies, and ended a long debate over how to represent these objects within BEL. EMBL-EBI also provides a programmatic API to the OLS service, for searching terms (<http://www.ebi.ac.uk/ols/api/search?q=folic%20acid>) and suggesting resolutions (<http://www.ebi.ac.uk/ols/api/suggest?q=folic+acid>)

1.3 Implementation

PyBEL is implemented using the PyParsing module. It provides flexibility and incredible speed in parsing compared to regular expression implementation. It also allows for the addition of parsing action hooks, which allow the graph to be checked semantically at compile-time.

It uses SQLite to provide a consistent and lightweight caching system for external data, such as namespaces, annotations, ontologies, and SQLAlchemy to provide a cross-platform interface. The same data management system is used to store graphs for high-performance querying.

1.4 Extensions to BEL

The PyBEL compiler is fully compliant with both BEL v1.0 and v2.0 and automatically upgrades legacy statements. Additionally, PyBEL includes several additions to the BEL specification to enable expression of important concepts in molecular biology that were previously missing and to facilitate integrating new data types. A short example is the inclusion of protein oxidation in the default BEL namespace for protein modifications. Other, more elaborate additions are outlined below.

1.4.1 Syntax for Epigenetics

PyBEL introduces the gene modification function, `gmod()`, as a syntax for encoding epigenetic modifications. Its usage mirrors the `pmod()` function for proteins and includes arguments for methylation.

For example, the methylation of NDUFB6 was found to be negatively correlated with its expression in a study of insulin resistance and Type II diabetes. This can now be expressed in BEL such as in the following statement:

```
g(HGNC:NDUFB6, gmod(Me)) negativeCorrelation r(HGNC:NDUFB6)
```

References:

- <https://www.ncbi.nlm.nih.gov/pubmed/17948130>

- <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4655260/>

Note: This syntax is currently under consideration as BEP-0006.

1.4.2 Definition of Namespaces as Regular Expressions

BEL imposes the constraint that each identifier must be qualified with an enumerated namespace to enable semantic interoperability and data integration. However, enumerating a namespace with potentially billions of names, such as dbSNP, poses a computational issue. PyBEL introduces syntax for defining namespaces with a consistent pattern using a regular expression to overcome this issue. For these namespaces, semantic validation can be performed in post-processing against the underlying database. The dbSNP namespace can be defined with a syntax familiar to BEL annotation definitions with regular expressions as follows:

```
DEFINE NAMESPACE dbSNP AS PATTERN "rs[0-9]+"
```

Note: This syntax was proposed with BEP-0005 and has been officially accepted as part of the BEL 2.1 specification.

1.4.3 Definition of Resources using OWL

Previous versions of PyBEL until 0.11.2 had an alternative namespace definition. Now it is recommended to either generate namespace files with reproducible build scripts following the Bio2BEL framework, or to directly add them to the database with the Bio2BEL `bio2bel.manager.namespace_manager.NamespaceManagerMixin` extension.

1.5 Things to Consider

1.5.1 Do All Statements Need Supporting Text?

Yes! All statements must be minimally qualified with a citation and evidence (now called SupportingText in BEL 2.0) to maintain provenance. Statements without evidence can't be traced to their source or evaluated independently from the curator, so they are excluded.

1.5.2 Multiple Annotations

All single annotations are considered as single element sets. When multiple annotations are present, all are unioned and attached to a given edge.

```
SET Citation = {"PubMed", "Example Article", "12345"}
SET ExampleAnnotation1 = {"Example Value 11", "Example Value 12"}
SET ExampleAnnotation2 = {"Example Value 21", "Example Value 22"}
p(HGNC:YFG1) -> p(HGNC:YFG2)
```

1.5.3 Namespace and Annotation Name Choices

*.belns and *.belanno configuration files include an entry called “Keyword” in their respective [Namespace] and [AnnotationDefinition] sections. To maintain understandability between BEL documents, PyBEL warns when the names given in *.bel documents do not match their respective resources. For now, capitalization is not considered, but in the future, PyBEL will also warn when capitalization is not properly stylized, like forgetting the lowercase ‘h’ in “ChEMBL”.

1.5.4 Why Not Nested Statements?

BEL has different relationships for modeling direct and indirect causal relations.

Direct

- $A \Rightarrow B$ means that A directly increases B through a physical process.
- $A =| B$ means that A directly decreases B through a physical process.

Indirect

The relationship between two entities can be coded in BEL, even if the process is not well understood.

- $A \rightarrow B$ means that A indirectly increases B . There are hidden elements in X that mediate this interaction through a pathway direct interactions $A (\Rightarrow \text{or } =|) X_1 (\Rightarrow \text{or } =|) \dots X_n (\Rightarrow \text{or } =|) B$, or through a set of multiple pathways that constitute a network.
- $A -| B$ means that A indirectly decreases B . Like for $A \rightarrow B$, this process involves hidden components with varying activities.

Increasing Nested Relationships

BEL also allows object of a relationship to be another statement.

- $A \Rightarrow (B \Rightarrow C)$ means that A increases the process by which B increases C . The example in the BEL Spec $p(\text{HGNC:GATA1}) \Rightarrow (\text{act}(p(\text{HGNC:ZBTB16})) \Rightarrow r(\text{HGNC:MPL}))$ represents GATA1 directly increasing the process by which ZBTB16 directly increases MPL. Before, directly increasing was used to specify physical contact, so it’s reasonable to conclude that $p(\text{HGNC:GATA1}) \Rightarrow \text{act}(p(\text{HGNC:ZBTB16}))$. The specification cites examples when B is an activity that only is affected in the context of A and C . This complicated enough that it is both impractical to standardize during curation, and impractical to represent in a network.
- $A \rightarrow (B \Rightarrow C)$ can be interpreted by assuming that A indirectly increases B , and because of monotonicity, conclude that $A \rightarrow C$ as well.
- $A \Rightarrow (B \rightarrow C)$ is more difficult to interpret, because it does not describe which part of process $B \rightarrow C$ is affected by A or how. Is it that $A \Rightarrow B$, and $B \Rightarrow C$, so we conclude $A \rightarrow C$, or does it mean something else? Perhaps A impacts a different portion of the hidden process in $B \rightarrow C$. These statements are ambiguous enough that they should be written as just $A \Rightarrow B$, and $B \rightarrow C$. If there is no literature evidence for the statement $A \rightarrow C$, then it is not the job of the curator to make this inference. Identifying statements of this might be the goal of a bioinformatics analysis of the BEL network after compilation.
- $A \rightarrow (B \rightarrow C)$ introduces even more ambiguity, and it should not be used.
- $A \Rightarrow (B =| C)$ states A increases the process by which B decreases C . One interpretation of this statement might be that $A \Rightarrow B$ and $B =| C$. An analysis could infer $A -| C$. Statements in the form of $A \rightarrow (B =| C)$ can also be resolved this way, but with added ambiguity.

Decreasing Nested Relationships

While we could agree on usage for the previous examples, the decrease of a nested statement introduces an unreasonable amount of ambiguity.

- $A = | (B \Rightarrow C)$ could mean A decreases B , and B also increases C . Does this mean A decreases C , or does it mean that C is still increased, but just not as much? Which of these statements takes precedence? Or do their effects cancel? The same can be said about $A - | (B \Rightarrow C)$, and with added ambiguity for indirect increases $A - | (B -> C)$
- $A = | (B = | C)$ could mean that A decreases B and B decreases C . We could conclude that A increases C , or could we again run into the problem of not knowing the precedence? The same is true for the indirect versions.

Recommendations for Use in PyBEL

After considering the ambiguity of nested statements to be a great risk to clarity, and PyBEL disables the usage of nested statements by default. See the Input and Output section for different parser settings. At Fraunhofer SCAI, curators resolved these statements to single statements to improve the precision and readability of our BEL documents.

While most statements in the form $A \text{ rel1 } (B \text{ rel2 } C)$ can be reasonably expanded to $A \text{ rel1 } B$ and $B \text{ rel2 } C$, the few that cannot are the difficult-to-interpret cases that we need to be careful about in our curation and later analyses.

1.5.5 Why Not RDF?

Current bel2rdf serialization tools build URLs with the OpenBEL Framework domain as a namespace, rather than respect the original namespaces of original entities. This does not follow the best practices of the semantic web, where URL's representing an object point to a real page with additional information. For example, UniProt does an exemplary job of this. Ultimately, using non-standard URLs makes harmonizing and data integration difficult.

Additionally, the RDF format does not easily allow for the annotation of edges. A simple statement in BEL that one protein up-regulates another can be easily represented in a triple in RDF, but when the annotations and citation from the BEL document need to be included, this forces RDF serialization to use approaches like representing the statement itself as a node. RDF was not intended to represent this type of information, but more properly for locating resources (hence its name). Furthermore, many blank nodes are introduced throughout the process. This makes RDF incredibly difficult to understand or work with. Later, writing queries in SPARQL becomes very difficult because the data format is complicated and the language is limited. For example, it would be incredibly complicated to write a query in SPARQL to get the objects of statements from publications by a certain author.

INSTALLATION

The latest stable code can be installed from [PyPI](#) with:

```
$ python3 -m pip install pybel
```

The most recent code can be installed from the source on [GitHub](#) with:

```
$ python3 -m pip install git+https://github.com/pybel/pybel.git
```

For developers, the repository can be cloned from [GitHub](#) and installed in editable mode with:

```
$ git clone https://github.com/pybel/pybel.git
$ cd pybel
$ python3 -m pip install -e .
```

2.1 Extras

The `setup.py` makes use of the `extras_require` argument of `setuptools.setup()` in order to make some heavy packages that support special features of PyBEL optional to install, in order to make the installation more lean by default. A single extra can be installed from PyPI like `python3 -m pip install pybel[neo4j]` or multiple can be installed using a list like `python3 -m pip install pybel[neo4j,indra]`. Likewise, for developer installation, extras can be installed in editable mode with `python3 -m pip install -e .[neo4j]` or multiple can be installed using a list like `python3 -m pip install -e .[neo4j,indra]`. The available extras are:

2.1.1 neo4j

This extension installs the `py2neo` package to support upload and download to Neo4j databases.

See also:

- [`pybel.to_neo4j\(\)`](#)

2.1.2 `indra`

This extra installs support for `indra`, the integrated network dynamical reasoner and assembler. Because it also represents biology in BEL-like statements, many statements from PyBEL can be converted to INDRA, and visa-versa. This package also enables the import of BioPAX, SBML, and SBGN into BEL.

See also:

- `pybel.from_biopax()`
- `pybel.from_indra_statements()`
- `pybel.from_indra_pickle()`
- `pybel.to_indra()`

2.1.3 `jupyter`

This extra installs support for visualizing BEL graphs in Jupyter notebooks.

See also:

- `pybel.io.jupyter.to_html()`
- `pybel.io.jupyter.to_jupyter()`

2.2 Caveats

- PyBEL extends the `networkx` for its core data structure. Many of the graphical aspects of `networkx` depend on `matplotlib`, which is an optional dependency.
- If `HTMLlib5` is installed, the test that's supposed to fail on a web page being missing actually tries to parse it as RDFa, and doesn't fail. Disregard this.

2.3 Upgrading

During the current development cycle, programmatic access to the definition and graph caches might become unstable. If you have any problems working with the database, try removing it with one of the following commands:

1. Running `pybel manage drop` (unix)
2. Running `python3 -m pybel manage drop` (windows)
3. Removing the folder `~/.pybel`

PyBEL will build a new database and populate it on the next run.

DATA MODEL

The `pybel.struct` module houses functions for handling the main data structure in PyBEL.

Because BEL expresses how biological entities interact within many different contexts, with descriptive annotations, PyBEL represents data as a directed multi-graph by sub-classing the `networkx.MultiDiGraph`. Each node is an instance of a subclass of the `pybel.dsl.BaseEntity` and each edge has a stable key and associated data dictionary for storing relevant contextual information.

The graph contains metadata for the PyBEL version, the BEL script metadata, the namespace definitions, the annotation definitions, and the warnings produced in analysis. Like any `networkx` graph, all attributes of a given object can be accessed through the `graph` property, like in: `my_graph.graph['my_key']`. Convenient property definitions are given for these attributes that are outlined in the documentation for `pybel.BELGraph`.

This allows for much easier programmatic access to answer more complicated questions, which can be written with python code. Because the data structure is the same in Neo4J, the data can be directly exported with `pybel.to_neo4j()`. Neo4J supports the Cypher querying language so that the same queries can be written in an elegant and simple way.

3.1 Constants

These documents refer to many aspects of the data model using constants, which can be found in the top-level module `pybel.constants`.

Terms describing abundances, annotations, and other internal data are designated in `pybel.constants` with full-caps, such as `pybel.constants.FUNCTION` and `pybel.constants.PROTEIN`.

For normal usage, we suggest referring to values in dictionaries by these constants, in case the hard-coded strings behind these constants change.

3.1.1 Function Nomenclature

The following table shows PyBEL's internal mapping from BEL functions to its own constants. This can be accessed programmatically via `pybel.parser.language.abundance_labels`.

BEL Function	PyBEL Constant	PyBEL DSL
a(), abundance()	<code>pybel.constants.ABUNDANCE</code>	<code>pybel.dsl.Abundance</code>
g(), geneAbundance()	<code>pybel.constants.GENE</code>	<code>pybel.dsl.Gene</code>
r(), rnaAbundance()	<code>pybel.constants.RNA</code>	<code>pybel.dsl.Rna</code>
m(), microRNAAbundance()	<code>pybel.constants.MIRNA</code>	<code>pybel.dsl.MicroRna</code>
p(), proteinAbundance()	<code>pybel.constants.PROTEIN</code>	<code>pybel.dsl.Protein</code>
bp(), biologicalProcess()	<code>pybel.constants.BIOPROCESS</code>	<code>pybel.dsl.BiologicalProcess</code>
path(), pathology()	<code>pybel.constants.PATHOLOGY</code>	<code>pybel.dsl.Pathology</code>
complex(), complexAbundance()	<code>pybel.constants.COMPLEX</code>	<code>pybel.dsl.ComplexAbundance</code>
composite(), compositeAbundance()	<code>pybel.constants.COMPOSITE</code>	<code>pybel.dsl.CompositeAbundance</code>
rxn(), reaction()	<code>pybel.constants.REACTION</code>	<code>pybel.dsl.Reaction</code>

3.2 Graph

```
class pybel.BELGraph(name=None, version=None, description=None, authors=None, contact=None, license=None, copyright=None, disclaimer=None, path=None)
```

An extension to `networkx.MultiDiGraph` to represent BEL.

Initialize a BEL graph with its associated metadata.

Parameters

- **name** (`Optional[str]`) – The graph’s name
- **version** (`Optional[str]`) – The graph’s version. Recommended to use semantic versioning or YYYYMMDD format.
- **description** (`Optional[str]`) – A description of the graph
- **authors** (`Optional[str]`) – The authors of this graph
- **contact** (`Optional[str]`) – The contact email for this graph
- **license** (`Optional[str]`) – The license for this graph
- **copyright** (`Optional[str]`) – The copyright for this graph
- **disclaimer** (`Optional[str]`) – The disclaimer for this graph

add (other)

Copy this graph and join it with another graph with it using `pybel.struct.left_full_join()`.

Parameters `other` (`BELGraph`) – Another BEL graph

Return type `BELGraph`

Example usage:

```
>>> import pybel  
>>> g = pybel.from_bel_script('...')
```

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```
>>> h = pybel.from_bel_script('...')

>>> k = g + h
```

`__iadd__(other)`Join another graph into this one, in-place, using `pybel.struct.left_full_join()`.**Parameters** `other` (`BELGraph`) – Another BEL graph**Return type** `BELGraph`

Example usage:

```
>>> import pybel
>>> g = pybel.from_bel_script('...')
>>> h = pybel.from_bel_script('...')
>>> g += h
```

`__and__(other)`

Create a deep copy of this graph and left outer joins another graph.

Uses `pybel.struct.left_outer_join()`.**Parameters** `other` (`BELGraph`) – Another BEL graph**Return type** `BELGraph`

Example usage:

```
>>> import pybel
>>> g = pybel.from_bel_script('...')
>>> h = pybel.from_bel_script('...')
>>> k = g & h
```

`__iand__(other)`Join another graph into this one, in-place, using `pybel.struct.left_outer_join()`.**Parameters** `other` (`BELGraph`) – Another BEL graph**Return type** `BELGraph`

Example usage:

```
>>> import pybel
>>> g = pybel.from_bel_script('...')
>>> h = pybel.from_bel_script('...')
>>> g &= h
```

`property path`

The graph's path, if it was derived from a BEL document.

Return type `Optional[str]`**`property name`**

The graph's name.

Hint: Can be set with the `SET DOCUMENT Name = "..."` entry in the source BEL script.**Return type** `Optional[str]`

property version

The graph's version.

Hint: Can be set with the `SET DOCUMENT Version = "..."` entry in the source BEL script.

Return type `Optional[str]`

property description

The graph's description.

Hint: Can be set with the `SET DOCUMENT Description = "..."` entry in the source BEL document.

Return type `Optional[str]`

property authors

The graph's authors.

Hint: Can be set with the `SET DOCUMENT Authors = "..."` entry in the source BEL document.

Return type `Optional[str]`

property contact

The graph's contact information.

Hint: Can be set with the `SET DOCUMENT ContactInfo = "..."` entry in the source BEL document.

Return type `Optional[str]`

property license

The graph's license.

Hint: Can be set with the `SET DOCUMENT Licenses = "..."` entry in the source BEL document

Return type `Optional[str]`

property copyright

The graph's copyright.

Hint: Can be set with the `SET DOCUMENT Copyright = "..."` entry in the source BEL document

Return type `Optional[str]`

property disclaimer

The graph's disclaimer.

Hint: Can be set with the SET DOCUMENT Disclaimer = "... " entry in the source BEL document.

Return type `Optional[str]`

property namespace_url

The mapping from the keywords used in this graph to their respective BEL namespace URLs.

Hint: Can be appended with the DEFINE NAMESPACE [key] AS URL "[value]" entries in the definitions section of the source BEL document.

Return type `Dict[str, str]`

property defined_namespace_keywords

The set of all keywords defined as namespaces in this graph.

Return type `Set[str]`

property namespace_pattern

The mapping from the namespace keywords used to create this graph to their regex patterns.

Hint: Can be appended with the DEFINE NAMESPACE [key] AS PATTERN "[value]" entries in the definitions section of the source BEL document.

Return type `Dict[str, str]`

property annotation_url

The mapping from the annotation keywords used to create this graph to the URLs of the BELANNO files.

Hint: Can be appended with the DEFINE ANNOTATION [key] AS URL "[value]" entries in the definitions section of the source BEL document.

Return type `Dict[str, str]`

property annotation_pattern

The mapping from the annotation keywords used to create this graph to their regex patterns as strings.

Hint: Can be appended with the DEFINE ANNOTATION [key] AS PATTERN "[value]" entries in the definitions section of the source BEL document.

Return type `Dict[str, str]`

property annotation_list

The mapping from the keywords of locally defined annotations to their respective sets of values.

Hint: Can be appended with the `DEFINE ANNOTATION [key] AS LIST {"[value]", ...}` entries in the definitions section of the source BEL document.

Return type `Dict[str, Set[str]]`

property defined_annotation_keywords

Get the set of all keywords defined as annotations in this graph.

Return type `Set[str]`

property pybel_version

The version of PyBEL with which this graph was produced as a string.

Return type `str`

property warnings

A list of warnings associated with this graph.

Return type `List[Tuple[Optional[str], BELParserWarning, Mapping]]`

number_of_warnings()

Return the number of warnings.

Return type `int`

number_of_citations()

Return the number of citations contained within the graph.

Return type `int`

number_of_authors()

Return the number of authors contained within the graph.

Return type `int`

get_authors()

Get the authors for the citations in the graph.

Return type `Set[str]`

add_unqualified_edge(*u*, *v*, *relation*)

Add a unique edge that has no annotations.

Parameters

- **u** (`BaseEntity`) – The source node
- **v** (`BaseEntity`) – The target node
- **relation** (`str`) – A relationship label from `pybel.constants`

Return type `str`

Returns The key for this edge (a unique hash)

add_transcription(*gene*, *rna*)

Add a transcription relation from a gene to an RNA or miRNA node.

Parameters

- **gene** (`Gene`) – A gene node

- **rna** (`Union[Rna, MicroRna]`) – An RNA or microRNA node

Return type `str`

add_translation (`rna, protein`)

Add a translation relation from a RNA to a protein.

Parameters

- **rna** (`Rna`) – An RNA node
- **protein** (`Protein`) – A protein node

Return type `str`

add_equivalence (`u: pybel.dsl.node_classes BaseEntity, v: pybel.dsl.node_classes BaseEntity,`

`*args, **kwargs) → str`

Add two equivalence relations for the nodes.

add_orthology (`u: pybel.dsl.node_classes BaseEntity, v: pybel.dsl.node_classes BaseEntity, *args,`

`**kwargs) → str`

Add two orthology relations for the nodes such that `u orthologousTo v` and `v orthologousTo u`.

add_is_a (`u: pybel.dsl.node_classes BaseEntity, v: pybel.dsl.node_classes BaseEntity, *, relation: str`

`= 'isA')` → `str`

Add an `isA` relationship such that `u isA v`.

add_part_of (`u: pybel.dsl.node_classes BaseEntity, v: pybel.dsl.node_classes BaseEntity, *, relation:`

`str = 'partOf')` → `str`

Add a `partOf` relationship such that `u partOf v`.

add_has_variant (`u: pybel.dsl.node_classes BaseEntity, v: pybel.dsl.node_classes BaseEntity, *, relation: str = 'hasVariant')` → `str`

Add a `hasVariant` relationship such that `u hasVariant v`.

add_has_reactant (`u: pybel.dsl.node_classes BaseEntity, v: pybel.dsl.node_classes BaseEntity, *, relation: str = 'hasReactant')` → `str`

Add a `hasReactant` relationship such that `u hasReactant v`.

add_has_product (`u: pybel.dsl.node_classes BaseEntity, v: pybel.dsl.node_classes BaseEntity, *, relation: str = 'hasProduct')` → `str`

Add a `hasProduct` relationship such that `u hasProduct v`.

add_qualified_edge (`u, v, *, relation, evidence, citation, annotations=None, subject_modifier=None, object_modifier=None, **attr`)

Add a qualified edge.

Qualified edges have a relation, evidence, citation, and optional annotations, subject modifications, and object modifications.

Parameters

- **u** – The source node
- **v** – The target node
- **relation** (`str`) – The type of relation this edge represents
- **evidence** (`str`) – The evidence string from an article
- **citation** (`Union[str, Tuple[str, str], CitationDict]`) – The citation data dictionary for this evidence. If a string is given, assumes it's a PubMed identifier and auto-fills the citation type.
- **annotations** (`Union[Mapping[str, str], Mapping[str, Set[str]], Mapping[str, Mapping[str, bool]], None]`) – The annotations data dictionary

- **subject_modifier** (`Optional[Mapping]`) – The modifiers (like activity) on the subject node. See data model documentation.
- **object_modifier** (`Optional[Mapping]`) – The modifiers (like activity) on the object node. See data model documentation.

Return type `str`

Returns The hash of the edge

add_binds (`u, v, *, evidence, citation, annotations=None, **attr`)

Add a “binding” relationship between the two entities such that `u => complex(u, v)`.

Return type `str`

add_increases (`u, v, *, relation: str = 'increases', evidence: str, citation: Union[str, Tuple[str, str], pybel.utils.CitationDict], annotations: Union[Mapping[str, str], Mapping[str, Set[str]], Mapping[str, Mapping[str, bool]], None] = None, subject_modifier: Optional[Mapping] = None, object_modifier: Optional[Mapping] = None, **attr`) → `str`

Wrap `add_qualified_edge()` for the `pybel.constants.INCREASES` relation.

add_directly_increases (`u, v, *, relation: str = 'directlyIncreases', evidence: str, citation: Union[str, Tuple[str, str], pybel.utils.CitationDict], annotations: Union[Mapping[str, str], Mapping[str, Set[str]], Mapping[str, Mapping[str, bool]], None] = None, subject_modifier: Optional[Mapping] = None, object_modifier: Optional[Mapping] = None, **attr`) → `str`

Add a `pybel.constants.DIRECTLY_INCREASES` with `add_qualified_edge()`.

add_decreases (`u, v, *, relation: str = 'decreases', evidence: str, citation: Union[str, Tuple[str, str], pybel.utils.CitationDict], annotations: Union[Mapping[str, str], Mapping[str, Set[str]], Mapping[str, Mapping[str, bool]], None] = None, subject_modifier: Optional[Mapping] = None, object_modifier: Optional[Mapping] = None, **attr`) → `str`

Add a `pybel.constants.DECREASES` relationship with `add_qualified_edge()`.

add_directly_decreases (`u, v, *, relation: str = 'directlyDecreases', evidence: str, citation: Union[str, Tuple[str, str], pybel.utils.CitationDict], annotations: Union[Mapping[str, str], Mapping[str, Set[str]], Mapping[str, Mapping[str, bool]], None] = None, subject_modifier: Optional[Mapping] = None, object_modifier: Optional[Mapping] = None, **attr`) → `str`

Add a `pybel.constants.DIRECTLY_DECREASES` relationship with `add_qualified_edge()`.

add_association (`u: pybel.dsl.node_classes BaseEntity, v: pybel.dsl.node_classes BaseEntity, *args, **kwargs`) → `str`

Add a `pybel.constants.ASSOCIATION` relationship with `add_qualified_edge()`.

add_regulates (`u, v, *, relation: str = 'regulates', evidence: str, citation: Union[str, Tuple[str, str], pybel.utils.CitationDict], annotations: Union[Mapping[str, str], Mapping[str, Set[str]], Mapping[str, Mapping[str, bool]], None] = None, subject_modifier: Optional[Mapping] = None, object_modifier: Optional[Mapping] = None, **attr`) → `str`

Add a `pybel.constants.REGULATES` relationship with `add_qualified_edge()`.

add_correlation (`u: pybel.dsl.node_classes BaseEntity, v: pybel.dsl.node_classes BaseEntity, *args, **kwargs`) → `str`

Add a `pybel.constants.CORRELATION` relationship with `add_qualified_edge()`.

add_no_correlation (`u: pybel.dsl.node_classes BaseEntity, v: pybel.dsl.node_classes BaseEntity, *args, **kwargs`) → `str`

Add a `pybel.constants.NO_CORRELATION` relationship with `add_qualified_edge()`.

```
add_positive_correlation(u: pybel.dsl.node_classes BaseEntity, v: pybel.dsl.node_classes BaseEntity, *args, **kwargs) → str
    Add a pybel.constants.POSITIVE_CORRELATION relationship with add_qualified_edge().

add_negative_correlation(u: pybel.dsl.node_classes BaseEntity, v: pybel.dsl.node_classes BaseEntity, *args, **kwargs) → str
    Add a pybel.constants.NEGATIVE_CORRELATION relationship with add_qualified_edge().

add_causes_no_change(u, v, *, relation: str = 'causesNoChange', evidence: str, citation: Union[str, Tuple[str, str], pybel.utils.CitationDict], annotations: Union[Mapping[str, str], Mapping[str, Set[str]], Mapping[str, Mapping[str, bool]], None] = None, subject_modifier: Optional[Mapping] = None, object_modifier: Optional[Mapping] = None, **attr) → str
    Add a pybel.constants.CAUSES_NO_CHANGE relationship with add_qualified_edge().

add_inhibits(u, v, *, relation: str = 'decreases', evidence: str, citation: Union[str, Tuple[str, str], pybel.utils.CitationDict], annotations: Union[Mapping[str, str], Mapping[str, Set[str]], Mapping[str, Mapping[str, bool]], None] = None, subject_modifier: Optional[Mapping] = None, object_modifier: Optional[Mapping] = {'modifier': 'Activity'}, **attr) → str
    Add an "inhibits" relationship.

    A more specific version of add_decreases() that automatically populates the object modifier with an activity.

add_activates(u, v, *, relation: str = 'increases', evidence: str, citation: Union[str, Tuple[str, str], pybel.utils.CitationDict], annotations: Union[Mapping[str, str], Mapping[str, Set[str]], Mapping[str, Mapping[str, bool]], None] = None, subject_modifier: Optional[Mapping] = None, object_modifier: Optional[Mapping] = {'modifier': 'Activity'}, **attr) → str
    Add an "activates" relationship.

    A more specific version of add_increases() that automatically populates the object modifier with an activity.

add_phosphorylates(u, v, code: Optional[str] = None, position: Optional[int] = None, *, evidence: str, citation: Union[str, Mapping[str, str]], annotations: Union[Mapping[str, str], Mapping[str, Set[str]], Mapping[str, Mapping[str, bool]], None] = None, subject_modifier: Optional[Mapping] = None, object_modifier: Optional[Mapping] = None, **attr)
    Add an increase of modified object with phosphorylation.

add_directly_phosphorylates(u, v, code: Optional[str] = None, position: Optional[int] = None, *, evidence: str, citation: Union[str, Mapping[str, str]], annotations: Union[Mapping[str, str], Mapping[str, Set[str]], Mapping[str, Mapping[str, bool]], None] = None, subject_modifier: Optional[Mapping] = None, object_modifier: Optional[Mapping] = None, **attr)
    Add a direct increase of modified object with phosphorylation.

add_dephosphorylates(u, v, code: Optional[str] = None, position: Optional[int] = None, *, evidence: str, citation: Union[str, Mapping[str, str]], annotations: Union[Mapping[str, str], Mapping[str, Set[str]], Mapping[str, Mapping[str, bool]], None] = None, subject_modifier: Optional[Mapping] = None, object_modifier: Optional[Mapping] = None, **attr)
    Add a decrease of modified object with phosphorylation.
```

```
add_directly_dephosphorylates(u, v, code: Optional[str] = None, position: Optional[int] = None, *, evidence: str, citation: Union[str, Mapping[str, str]], annotations: Union[Mapping[str, str], Mapping[str, Set[str]], Mapping[str, Mapping[str, bool]]], None = None, subject_modifier: Optional[Mapping] = None, object_modifier: Optional[Mapping] = None, **attr)
```

Add a direct decrease of modified object with phosphorylation.

```
add_node_from_data(node)
```

Add an entity to the graph.

Return type `None`

```
add_reaction(reactants, products)
```

Add a reaction directly to the graph.

```
has_edge_citation(u, v, key)
```

Check if the given edge has a citation.

Return type `bool`

```
has_edge_evidence(u, v, key)
```

Check if the given edge has an evidence.

Return type `bool`

```
get_edge_citation(u, v, key)
```

Get the citation for a given edge.

Return type `Optional[CitationDict]`

```
get_edge_evidence(u, v, key)
```

Get the evidence for a given edge.

Return type `Optional[str]`

```
get_edge_annotations(u, v, key)
```

Get the annotations for a given edge.

Return type `Optional[Mapping[str, Mapping[str, bool]]]`

```
static node_to_bel(n)
```

Serialize a node as BEL.

Return type `str`

```
static edge_to_bel(u, v, edge_data, sep=None, use_identifiers=False)
```

Serialize a pair of nodes and related edge data as a BEL relation.

Return type `str`

```
iter_equivalent_nodes(node)
```

Iterate over nodes that are equivalent to the given node, including the original.

Return type `Iterable[BaseEntity]`

```
get_equivalent_nodes(node)
```

Get a set of equivalent nodes to this node, excluding the given node.

Return type `Set[BaseEntity]`

```
node_has_namespace(node, namespace)
```

Check if the node have the given namespace.

This also should look in the equivalent nodes.

Return type `bool`

summary_dict()
Return a dictionary that summarizes the graph.

Return type `Mapping[str, float]`

summary_str()
Return a string that summarizes the graph.

Return type `str`

summarize(file=None)
Print a summary of the graph.

Return type `None`

3.3 Nodes

Nodes (or *entities*) in a `pybel.BELGraph` represent physical entities' abundances. Most contain information about the identifier for the entity using a namespace/name pair. The PyBEL parser converts BEL terms to an internal representation using an internal domain specific language (DSL) that allows for writing BEL directly in Python.

For example, after the BEL term `p (HGNC:GSK3B)` is parsed, it is instantiated as a Python object using the DSL function corresponding to the `p()` function in BEL, `pybel.dsl.Protein`, like:

```
from pybel.dsl import Protein
gsk3b_protein = Protein(namespace='HGNC', name='GSK3B')
```

`pybel.dsl.Protein`, like the others mentioned before, inherit from `pybel.dsl BaseEntity`, which itself inherits from `dict`. Therefore, the resulting object can be used like a dict that looks like:

```
from pybel.constants import *
{
    FUNCTION: PROTEIN,
    NAMESPACE: 'HGNC',
    NAME: 'GSK3B',
}
```

Alternatively, it can be used in more exciting ways, outlined later in the documentation for `pybel.dsl`.

3.3.1 Variants

The addition of a variant tag results in an entry called ‘variants’ in the data dictionary associated with a given node. This entry is a list with dictionaries describing each of the variants. All variants have the entry ‘kind’ to identify whether it is a post-translational modification (PTM), gene modification, fragment, or HGVS variant.

Warning: The canonical ordering for the elements of the `VARIANTS` list correspond to the sorted order of their corresponding node tuples using `pybel.parser.canonicalize.sort_dict_list()`. Rather than directly modifying the `BELGraph`'s structure, use `pybel.BELGraph.add_node_from_data()`, which takes care of automatically canonicalizing this dictionary.

HGVS Variants.

For example, the BEL term `p (HGNC:GSK3B, var (p.Gly123Arg))` is translated to the following internal DSL:

```
from pybel.dsl import Protein, Hgvs
gsk3b_variant = Protein(namespace='HGNC', name='GSK3B', variants=Hgvs('p.Gly123Arg'))
```

Further, the shorthand for protein substitutions, `pybel.dsl.ProteinSubstitution`, can be used to produce the same result, as it inherits from `pybel.dsl.Hgvs`:

```
from pybel.dsl import Protein, ProteinSubstitution
gsk3b_variant = Protein(namespace='HGNC', name='GSK3B', variants=ProteinSubstitution(
    'Gly', 123, 'Arg'))
```

Either way, the resulting object can be used like a dict that looks like:

```
from pybel.constants import *
{
    FUNCTION: PROTEIN,
    NAMESPACE: 'HGNC',
    NAME: 'GSK3B',
    VARIANTS: [
        {
            KIND: HGVS,
            IDENTIFIER: 'p.Gly123Arg',
        },
    ],
}
```

See also:

- BEL 2.0 specification on [variants](#)
- HGVS conventions
- PyBEL module `pybel.parser.modifiers.get_hgvs_language`

3.3.2 Gene Substitutions

Gene Substitutions.

Gene substitutions are legacy statements defined in BEL 1.0. BEL 2.0 recommends using HGVS strings. Luckily, the information contained in a BEL 1.0 encoding, such as `g (HGNC:APP, sub (G, 275341, C))` can be automatically translated to the appropriate HGVS `g (HGNC:APP, var (c.275341G>C))`, assuming that all substitutions are using the reference coding gene sequence for numbering and not the genomic reference. The previous statements both produce the underlying data:

```
from pybel.constants import *
{
    FUNCTION: GENE,
    NAMESPACE: 'HGNC',
    NAME: 'APP',
    VARIANTS: [
        {
            KIND: HGVS,
            IDENTIFIER: 'c.275341G>C',
        },
    ],
}
```

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```
],
}
```

See also:

- BEL 2.0 specification on [gene substitutions](#)
- PyBEL module `pybel.parser.modifiers.get_gene_substitution_language`

3.3.3 Gene Modifications

Gene Modifications.

PyBEL introduces the gene modification tag, `gmod()`, to allow for the encoding of epigenetic modifications. Its syntax follows the same style as the `pmod()` tags for proteins, and can include the following values:

- M
- Me
- methylation
- A
- Ac
- acetylation

For example, the node `g (HGNC:GSK3B, gmod (M))` is represented with the following:

```
from pybel.constants import *

{
    FUNCTION: GENE,
    NAMESPACE: 'HGNC',
    NAME: 'GSK3B',
    VARIANTS: [
        {
            KIND: GMOD,
            IDENTIFIER: {
                NAMESPACE: BEL_DEFAULT_NAMESPACE,
                NAME: 'Me',
            },
        },
    ],
}
```

The addition of this function does not preclude the use of all other standard functions in BEL; however, other compilers probably won't support these standards. If you agree that this is useful, please contribute to discussion in the OpenBEL community.

See also:

- PyBEL module `pybel.parser.modifiers.get_gene_modification_language()`

3.3.4 Protein Substitutions

Protein Substitutions.

Protein substitutions are legacy statements defined in BEL 1.0. BEL 2.0 recommends using HGVS strings. Luckily, the information contained in a BEL 1.0 encoding, such as `p(HGNC:APP, sub(R, 275, H))` can be automatically translated to the appropriate HGVS `p(HGNC:APP, var(p.Arg275His))`, assuming that all substitutions are using the reference protein sequence for numbering and not the genomic reference. The previous statements both produce the underlying data:

```
from pybel.constants import *

{
    FUNCTION: GENE,
    NAMESPACE: 'HGNC',
    NAME: 'APP',
    VARIANTS: [
        {
            KIND: HGVS,
            IDENTIFIER: 'p.Arg275His',
        },
    ],
}
```

See also:

- BEL 2.0 specification on [protein substitutions](#)
- PyBEL module `pybel.parser.modifiers.get_protein_substitution_language`

3.3.5 Protein Modifications

Protein Modifications.

The addition of a post-translational modification (PTM) tag results in an entry called ‘variants’ in the data dictionary associated with a given node. This entry is a list with dictionaries describing each of the variants. All variants have the entry ‘kind’ to identify whether it is a PTM, gene modification, fragment, or HGVS variant. The ‘kind’ value for PTM is ‘pmod’.

Each PMOD contains an identifier, which is a dictionary with the namespace and name, and can optionally include the position (‘pos’) and/or amino acid code (‘code’).

For example, the node `p(HGNC:GSK3B, pmod(P, S, 9))` is represented with the following:

```
from pybel.constants import *

{
    FUNCTION: PROTEIN,
    NAMESPACE: 'HGNC',
    NAME: 'GSK3B',
    VARIANTS: [
        {
            KIND: PMOD,
            IDENTIFIER: {
                NAMESPACE: BEL_DEFAULT_NAMESPACE
                NAME: 'Ph',
            },
            PMOD_CODE: 'Ser',
        },
    ],
}
```

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```

        PMOD_POSITION: 9,
    },
],
}

```

As an additional example, in `p(HGNC:MAPK1, pmod(Ph, Thr, 202), pmod(Ph, Tyr, 204))`, MAPK is phosphorylated twice to become active. This results in the following:

```

{
    FUNCTION: PROTEIN,
    NAMESPACE: 'HGNC',
    NAME: 'MAPK1',
    VARIANTS: [
        {
            KIND: PMOD,
            IDENTIFIER: {
                NAMESPACE: BEL_DEFAULT_NAMESPACE
                NAME: 'Ph',
            },
            PMOD_CODE: 'Thr',
            PMOD_POSITION: 202
        },
        {
            KIND: PMOD,
            IDENTIFIER: {
                NAMESPACE: BEL_DEFAULT_NAMESPACE
                NAME: 'Ph',
            },
            PMOD_CODE: 'Tyr',
            PMOD_POSITION: 204
        }
    ]
}

```

See also:

- BEL 2.0 specification on [protein modifications](#)
- PyBEL module `pybel.parser.modifiers.get_protein_modification_language`

3.3.6 Protein Truncations

Truncations.

Truncations in the legacy BEL 1.0 specification are automatically translated to BEL 2.0 with HGVS nomenclature. `p(HGNC:AKT1, trunc(40))` becomes `p(HGNC:AKT1, var(p.40*))` and is represented with the following dictionary:

```

from pybel.constants import *

{
    FUNCTION: PROTEIN,
    NAMESPACE: 'HGNC',
    NAME: 'AKT1',
}

```

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```
VARIANTS: [
    {
        KIND: HGVS,
        IDENTIFIER: 'p.40*',
    },
]
```

Unfortunately, the HGVS nomenclature requires the encoding of the terminal amino acid which is exchanged for a stop codon, and this information is not required by BEL 1.0. For this example, the proper encoding of the truncation at position also includes the information that the 40th amino acid in the AKT1 is Cys. Its BEL encoding should be `p(HGNC:AKT1, var(p.Cys40*))`. Temporary support has been added to compile these statements, but it's recommended they are upgraded by reexamining the supporting text, or looking up the amino acid sequence.

See also:

- BEL 2.0 specification on [truncations](#)
- PyBEL module `pybel.parser.modifiers.get_truncation_language`

3.3.7 Protein Fragments

Fragments.

The addition of a fragment results in an entry called `pybel.constants.VARIANTS` in the data dictionary associated with a given node. This entry is a list with dictionaries describing each of the variants. All variants have the entry `pybel.constants.KIND` to identify whether it is a PTM, gene modification, fragment, or HGVS variant. The `pybel.constants.KIND` value for a fragment is `pybel.constants.FRAGMENT`.

Each fragment contains an identifier, which is a dictionary with the namespace and name, and can optionally include the position ('pos') and/or amino acid code ('code').

For example, the node `p(HGNC:GSK3B, frag(45_129))` is represented with the following:

```
from pybel.constants import *

{
    FUNCTION: PROTEIN,
    NAMESPACE: 'HGNC',
    NAME: 'GSK3B',
    VARIANTS: [
        {
            KIND: FRAGMENT,
            FRAGMENT_START: 45,
            FRAGMENT_STOP: 129,
        },
    ],
}
```

Additionally, nodes can have an asterick (*) or question mark (?) representing unbound or unknown fragments, respectively.

A fragment may also be unknown, such as in the node `p(HGNC:GSK3B, frag(?))`. This is represented with the key `pybel.constants.FRAGMENT_MISSING` and the value of '?' like:

```
from pybel.constants import *
```

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```
{
    FUNCTION: PROTEIN,
    NAMESPACE: 'HGNC',
    NAME: 'GSK3B',
    VARIANTS: [
        {
            KIND: FRAGMENT,
            FRAGMENT_MISSING: '?',
        },
    ],
}
```

See also:

- BEL 2.0 specification on [proteolytic fragments](#) (2.2.3)
- PyBEL module `pybel.parser.modifiers.get_fragment_language`

3.3.8 Fusions

Fusions.

Gene, RNA, miRNA, and protein fusions are all represented with the same underlying data structure. Below it is shown with uppercase letters referring to constants from `pybel.constants` and. For example, g(HGNC:BCR, fus(HGNC:JAK2, 1875, 2626)) is represented as:

```
from pybel.constants import *

{
    FUNCTION: GENE,
    FUSION: {
        PARTNER_5P: {NAMESPACE: 'HGNC', NAME: 'BCR'},
        PARTNER_3P: {NAMESPACE: 'HGNC', NAME: 'JAK2'},
        RANGE_5P: {
            FUSION_REFERENCE: 'c',
            FUSION_START: '?',
            FUSION_STOP: 1875,
        },
        RANGE_3P: {
            FUSION_REFERENCE: 'c',
            FUSION_START: 2626,
            FUSION_STOP: '?',
        },
    },
}
```

See also:

- BEL 2.0 specification on [fusions](#) (2.6.1)
- PyBEL module `pybel.parser.modifiers.get_fusion_language`
- PyBEL module `pybel.parser.modifiers.get_legacy_fusion_language`

3.4 Unqualified Edges

Unqualified edges are automatically inferred by PyBEL and do not contain citations or supporting evidence.

3.4.1 Variant and Modifications' Parent Relations

All variants, modifications, fragments, and truncations are connected to their parent entity with an edge having the relationship `hasParent`.

For `p(HGNC:GSK3B, var(p.Gly123Arg))`, the following edge is inferred:

```
p(HGNC:GSK3B, var(p.Gly123Arg)) hasParent p(HGNC:GSK3B)
```

All variants have this relationship to their reference node. BEL does not specify relationships between variants, such as the case when a given phosphorylation is necessary to make another one. This knowledge could be encoded directly like BEL, since PyBEL does not restrict users from manually asserting unqualified edges.

3.4.2 List Abundances

Complexes and composites that are defined by lists. As of version 0.9.0, they contain a list of the data dictionaries that describe their members. For example `complex(p(HGNC:FOS), p(HGNC:JUN))` becomes:

```
from pybel.constants import *

{
    FUNCTION: COMPLEX,
    MEMBERS: [
        {
            FUNCTION: PROTEIN,
            NAMESPACE: 'HGNC',
            NAME: 'FOS',
        },
        {
            FUNCTION: PROTEIN,
            NAMESPACE: 'HGNC',
            NAME: 'JUN',
        }
    ]
}
```

The following edges are also inferred:

```
complex(p(HGNC:FOS), p(HGNC:JUN)) hasMember p(HGNC:FOS)
complex(p(HGNC:FOS), p(HGNC:JUN)) hasMember p(HGNC:JUN)
```

See also:

BEL 2.0 specification on complex abundances

Similarly, `composite(a(CHEBI:malonate), p(HGNC:JUN))` becomes:

```
from pybel.constants import *

{
    FUNCTION: COMPOSITE,
    MEMBERS: [

```

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```
{
    FUNCTION: ABUNDANCE,
    NAMESPACE: 'CHEBI',
    NAME: 'malonate',
}, {
    FUNCTION: PROTEIN,
    NAMESPACE: 'HGNC',
    NAME: 'JUN',
}
]
```

The following edges are inferred:

```
composite(a(CHEBI:malonate), p(HGNC:JUN)) hasComponent a(CHEBI:malonate)
composite(a(CHEBI:malonate), p(HGNC:JUN)) hasComponent p(HGNC:JUN)
```

Warning: The canonical ordering for the elements of the `pybel.constants.MEMBERS` list correspond to the sorted order of their corresponding node tuples using `pybel.parser.canonicalize.sort_dict_list()`. Rather than directly modifying the BELGraph's structure, use `BELGraph.add_node_from_data()`, which takes care of automatically canonicalizing this dictionary.

See also:

BEL 2.0 specification on composite abundances

3.4.3 Reactions

The usage of a reaction causes many nodes and edges to be created. The following example will illustrate what is added to the network for

```
rxn(reactants(a(CHEBI:"(3S)-3-hydroxy-3-methylglutaryl-CoA"), a(CHEBI:"NADPH"), \
a(CHEBI:"hydron")), products(a(CHEBI:"mevalonate"), a(CHEBI:"NADP (+)")))
```

As of version 0.9.0, the reactants' and products' data dictionaries are included as sub-lists keyed REACTANTS and PRODUCTS. It becomes:

```
from pybel.constants import *

{
    FUNCTION: REACTION
    REACTANTS: [
        {
            FUNCTION: ABUNDANCE,
            NAMESPACE: 'CHEBI',
            NAME: '(3S)-3-hydroxy-3-methylglutaryl-CoA'
        },
        {
            FUNCTION: ABUNDANCE,
            NAMESPACE: 'CHEBI',
            NAME: 'NADPH',
        },
        {
            FUNCTION: ABUNDANCE,
            NAMESPACE: 'CHEBI',
            NAME: 'hydron',
        }
    ]
}
```

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```
        NAME: 'hydron',
    }
],
PRODUCTS: [
{
    FUNCTION: ABUNDANCE,
    NAMESPACE: 'CHEBI',
    NAME: 'mevalonate',
},
{
    FUNCTION: ABUNDANCE,
    NAMESPACE: 'CHEBI',
    NAME: 'NADP(+)',
}
]
```

Warning: The canonical ordering for the elements of the REACTANTS and PRODUCTS lists correspond to the sorted order of their corresponding node tuples using `pybel.parser.canonicalize.sort_dict_list()`. Rather than directly modifying the BELGraph's structure, use `BELGraph.add_node_from_data()`, which takes care of automatically canonicalizing this dictionary.

The following edges are inferred, where X represents the previous reaction, for brevity:

```
X hasReactant a(CHEBI:"(3S)-3-hydroxy-3-methylglutaryl-CoA")
X hasReactant a(CHEBI:"NADPH")
X hasReactant a(CHEBI:"hydron")
X hasProduct a(CHEBI:"mevalonate")
X hasProduct a(CHEBI:"NADP(+))")
```

See also:

[BEL 2.0 specification on reactions](#)

3.5 Edges

3.5.1 Design Choices

In the OpenBEL Framework, modifiers such as activities (kinaseActivity, etc.) and transformations (translocations, degradations, etc.) were represented as their own nodes. In PyBEL, these modifiers are represented as a property of the edge. In reality, an edge like `sec(p(HGNC:A)) -> activity(p(HGNC:B), ma(kinaseActivity))` represents a connection between HGNC:A and HGNC:B. Each of these modifiers explains the context of the relationship between these physical entities. Further, querying a network where these modifiers are part of a relationship is much more straightforward. For example, finding all proteins that are upregulated by the kinase activity of another protein now can be directly queried by filtering all edges for those with a subject modifier whose modification is molecular activity, and whose effect is kinase activity. Having fewer nodes also allows for a much easier display and visual interpretation of a network. The information about the modifier on the subject and activity can be displayed as a color coded source and terminus of the connecting edge.

The compiler in OpenBEL framework created nodes for molecular activities like `kin(p(HGNC:YFG))` and induced an edge like `p(HGNC:YFG) actsIn kin(p(HGNC:YFG))`. For transformations, a statement like `tloc(p(HGNC:YFG), GOCC:intracellular, GOCC:"cell membrane")` also induced

```
tloc(p(HGNC:YFG), GOCC:intracellular, GOCC:"cell membrane") translocates
p(HGNC:YFG).
```

In PyBEL, we recognize that these modifications are actually annotations to the type of relationship between the subject's entity and the object's entity. `p(HGNC:ABC) -> tloc(p(HGNC:YFG), GOCC:intracellular, GOCC:"cell membrane")` is about the relationship between `p(HGNC:ABC)` and `p(HGNC:YFG)`, while the information about the translocation qualifies that the object is undergoing an event, and not just the abundance. This is a confusion with the use of `proteinAbundance` as a keyword, and perhaps is why many people prefer to use just the keyword `p`

3.5.2 Example Edge Data Structure

Because this data is associated with an edge, the node data for the subject and object are not included explicitly. However, information about the activities, modifiers, and transformations on the subject and object are included. Below is the “skeleton” for the edge data model in PyBEL:

```
from pybel.constants import *

{
    SUBJECT: {
        # ... modifications to the subject node. Only present if non-empty.
    },
    RELATION: POSITIVE_CORRELATION,
    OBJECT: {
        # ... modifications to the object node. Only present if non-empty.
    },
    EVIDENCE: ...,
    CITATION : {
        CITATION_TYPE: CITATION_TYPE_PUBMED,
        CITATION_REFERENCE: ...,
        CITATION_DATE: 'YYYY-MM-DD',
        CITATION_AUTHORS: 'Jon Snow|John Doe',
    },
    ANNOTATIONS: {
        'Disease': {
            'Colorectal Cancer': True,
        },
        # ... additional annotations as tuple[str, dict[str,bool]] pairs
    },
}
```

Each edge must contain the `RELATION`, `EVIDENCE`, and `CITATION` entries. The `CITATION` must minimally contain `CITATION_TYPE` and `CITATION_REFERENCE` since these can be used to look up additional metadata.

Note: Since version 0.10.2, annotations now always appear as dictionaries, even if only one value is present.

3.5.3 Activities

Modifiers are added to this structure as well. Under this schema, `p(HGNC:GSK3B, pmod(P, S, 9)) pos act(p(HGNC:GSK3B), ma(kin))` becomes:

```
from pybel.constants import *

{
    RELATION: POSITIVE_CORRELATION,
    OBJECT: {
        MODIFIER: ACTIVITY,
        EFFECT: {
            NAME: 'kin',
            NAMESPACE: BEL_DEFAULT_NAMESPACE,
        }
    },
    CITATION: { ... },
    EVIDENCE: ...,
    ANNOTATIONS: { ... },
}
```

Activities without molecular activity annotations do not contain an `pybel.constants.EFFECT` entry: Under this schema, `p(HGNC:GSK3B, pmod(P, S, 9)) pos act(p(HGNC:GSK3B))` becomes:

```
from pybel.constants import *

{
    RELATION: POSITIVE_CORRELATION,
    OBJECT: {
        MODIFIER: ACTIVITY
    },
    CITATION: { ... },
    EVIDENCE: ...,
    ANNOTATIONS: { ... },
}
```

3.5.4 Locations

Locations.

Location data also is added into the information in the edge for the node (subject or object) for which it was annotated. `p(HGNC:GSK3B, pmod(P, S, 9), loc(GO:lysosome)) pos act(p(HGNC:GSK3B), ma(kin))` becomes:

```
from pybel.constants import *

{
    SUBJECT: {
        LOCATION: {
            NAMESPACE: 'GO',
            NAME: 'lysosome',
        }
    },
    RELATION: POSITIVE_CORRELATION,
    OBJECT: {
        MODIFIER: ACTIVITY,
```

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```

EFFECT: {
    NAMESPACE: BEL_DEFAULT_NAMESPACE
    NAME: 'kin',
}
},
EVIDENCE: ...,
CITATION: { ... },
}

```

The addition of the `location()` element in BEL 2.0 allows for the unambiguous expression of the differences between the process of hypothetical HGNC:A moving from one place to another and the existence of hypothetical HGNC:A in a specific location having different effects. In BEL 1.0, this action had its own node, but this introduced unnecessary complexity to the network and made querying more difficult. This calls for thoughtful consideration of the following two statements:

- `tloc(p(HGNC:A), fromLoc(GO:intracellular), toLoc(GO:"cell membrane")) -> p(HGNC:B)`
- `p(HGNC:A, location(GO:"cell membrane")) -> p(HGNC:B)`

See also:

- BEL 2.0 specification on [cellular location](#) (2.2.4)
- PyBEL module `pybel.parser.modifiers.get_location_language`

3.5.5 Translocations

Translocations have their own unique syntax. `p(HGNC:YFG1) -> sec(p(HGNC:YFG2))` becomes:

```

from pybel.constants import *

{
    RELATION: INCREASES,
    OBJECT: {
        MODIFIER: TRANSLOCATION,
        EFFECT: {
            FROM_LOC: {
                NAMESPACE: 'GO',
                NAME: 'intracellular',
            },
            TO_LOC: {
                NAMESPACE: 'GO',
                NAME: 'extracellular space',
            }
        }
    },
    CITATION: { ... },
    EVIDENCE: ...,
    ANNOTATIONS: { ... },
}

```

See also:

[BEL 2.0 specification on translocations](#)

3.5.6 Degradations

Degradations are more simple, because there's no :pybel.constants.EFFECT entry. $p(\text{HGNC:YFG1}) \rightarrow \text{deg}(p(\text{HGNC:YFG2}))$ becomes:

```
from pybel.constants import *

{
    RELATION: INCREASES,
    OBJECT: {
        MODIFIER: DEGRADATION,
    },
    CITATION: { ... },
    EVIDENCE: ...,
    ANNOTATIONS: { ... },
}
```

See also:

[BEL 2.0 specification on degradations](#)

**CHAPTER
FOUR**

EXAMPLE NETWORKS

This directory contains example networks, precompiled as BEL graphs that are appropriate to use in examples.

An example describing EGF's effect on cellular processes.

```
SET Citation = {"PubMed", "Clin Cancer Res 2003 Jul 9(7) 2416-25", "12855613"}  
SET Evidence = "This induction was not seen either when LNCaP cells were treated with flutamide or conditioned medium were pretreated with antibody to the epidermal growth factor (EGF)"  
SET Species = 9606  
  
tscript(p(HGNC:AR)) increases p(HGNC:EGF)  
  
UNSET ALL  
  
SET Citation = {"PubMed", "Int J Cancer 1998 Jul 3 77(1) 138-45", "9639405"}  
SET Evidence = "DU-145 cells treated with 5000 U/ml of IFNgamma and IFN alpha, both reduced EGF production with IFN gamma reduction more significant."  
SET Species = 9606  
  
p(HGNC:IFNA1) decreases p(HGNC:EGF)  
p(HGNC:IFNG) decreases p(HGNC:EGF)  
  
UNSET ALL  
  
SET Citation = {"PubMed", "DNA Cell Biol 2000 May 19(5) 253-63", "10855792"}  
SET Evidence = "Although found predominantly in the cytoplasm and, less abundantly, in the nucleus, VCP can be translocated from the nucleus after stimulation with epidermal growth factor."  
SET Species = 9606  
  
p(HGNC:EGF) increases tloc(p(HGNC:VCP), GO:nucleus, GO:cytoplasm)  
  
UNSET ALL  
  
SET Citation = {"PubMed", "J Clin Oncol 2003 Feb 1 21(3) 447-52", "12560433"}  
SET Evidence = "Valosin-containing protein (VCP; also known as p97) has been shown to be associated with antiapoptotic function and metastasis via activation of the nuclear factor-kappaB signaling pathway."  
SET Species = 9606  
  
cat(p(HGNC:VCP)) increases tscript(complex(p(HGNC:NFKB1), p(HGNC:NFKB2), p(HGNC:REL), p(HGNC:RELA), p(HGNC:RELB)))  
tscript(complex(p(HGNC:NFKB1), p(HGNC:NFKB2), p(HGNC:REL), p(HGNC:RELA), p(HGNC:RELB))) decreases bp(MESHPP:Apoptosis)
```

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UNSET ALL

pybel.examples.egf_graph

Curation of the article “Genetics ignite focus on microglial inflammation in Alzheimer’s disease”.

```
SET Citation = {"PubMed", "26438529"}
SET Evidence = "Sialic acid binding activates CD33, resulting in phosphorylation of_
↪the CD33
immunoreceptor tyrosine-based inhibitory motif (ITIM) domains and activation of the_
↪SHP-1 and
SHP-2 tyrosine phosphatases [66, 67]."

complex(p(HGNC:CD33), a(CHEBI:"sialic acid")) -> p(HGNC:CD33, pmod(P))
act(p(HGNC:CD33, pmod(P))) => act(p(HGNC:PTPN6), ma(phos))
act(p(HGNC:CD33, pmod(P))) => act(p(HGNC:PTPN11), ma(phos))

UNSET {Evidence, Species}

SET Evidence = "These phosphatases act on multiple substrates, including Syk, to_
↪inhibit immune
activation [68, 69]. Hence, CD33 activation leads to increased SHP-1 and SHP-2_
↪activity that antagonizes Syk,
inhibiting ITAM-signaling proteins, possibly including TREM2/DAP12 (Fig. 1, [70, 71])._
↪"

SET Species = 9606

act(p(HGNC:PTPN6)) =| act(p(HGNC:SYK))
act(p(HGNC:PTPN11)) =| act(p(HGNC:SYK))
act(p(HGNC:SYK)) -> act(p(HGNC:TREM2))
act(p(HGNC:SYK)) -> act(p(HGNC:TYROBP))

UNSET ALL
```

pybel.examples.sialic_acid_graph

An example describing a single evidence about BRAF.

```
SET Citation = {"PubMed", "11283246"}
SET Evidence = "Expression of both dominant negative forms, RasN17 and Rap1N17, in_
↪UT7-Mpl cells decreased
thrombopoietin-mediated Elk1-dependent transcription. This suggests that both Ras and_
↪Rap1 contribute to
thrombopoietin-induced ELK1 transcription."

SET Species = 9606

p(HGNC:THPO) increases kin(p(HGNC:BRAF))
p(HGNC:THPO) increases kin(p(HGNC:RAF1))
kin(p(HGNC:BRAF)) increases tscript(p(HGNC:ELK1))

UNSET ALL
```

pybel.examples.braf_example_graph

An example describing statins.

pybel.examples.statin_graph

An example describing a translocation.

```
SET Citation = {"PubMed", "16170185"}  
SET Evidence = "These modifications render Ras functional and capable of localizing  
to the lipid-rich inner surface of the cell membrane. The first and most critical  
modification, farnesylation, which is principally catalyzed by protein FTase, adds  
a 15-carbon hydrophobic farnesyl isoprenyl tail to the carboxyl terminus of Ras."  
SET TextLocation = Review  
  
cat(complex(p(HGNC:FNTA),p(HGNC:FNTB))) directlyIncreases p(SFAM:"RAS Family",pmod(F))  
p(SFAM:"RAS Family",pmod(F)) directlyIncreases tloc(p(SFAM:"RAS Family"),MESHCS:  
"Intracellular Space",MESHCS:"Cell Membrane")
```

pybel.examples.ras_tloc_graph

SUMMARY

Summary functions for BEL graphs.

`pybel.struct.summary.get_syntax_errors(graph)`

List the syntax errors encountered during compilation of a BEL script.

Return type `List[Tuple[Optional[str], BELParserWarning, Mapping]]`

`pybel.struct.summary.count_error_types(graph)`

Count the occurrence of each type of error in a graph.

Return type `Counter[str]`

Returns A Counter of {error type: frequency}

`pybel.struct.summary.count_naked_names(graph)`

Count the frequency of each naked name (names without namespaces).

Return type `Counter[str]`

Returns A Counter from {name: frequency}

`pybel.struct.summary.get_naked_names(graph)`

Get the set of naked names in the graph.

Return type `Set[str]`

`pybel.struct.summary.calculate_incorrect_name_dict(graph)`

Get missing names grouped by namespace.

Return type `Mapping[str, List[str]]`

`pybel.struct.summary.calculate_error_by_annotation(graph, annotation)`

Group error names by a given annotation.

Return type `Mapping[str, List[str]]`

`pybel.struct.summary.get_functions(graph)`

Get the set of all functions used in this graph.

Parameters `graph` (`pybel.BELGraph`) – A BEL graph

Return type `Set[str]`

Returns A set of functions

`pybel.struct.summary.count_functions(graph)`

Count the frequency of each function present in a graph.

Parameters `graph` (`pybel.BELGraph`) – A BEL graph

Return type `Counter[str]`

Returns A Counter from {function: frequency}

`pybel.struct.summary.count_namespaces(graph)`

Count the frequency of each namespace across all nodes (that have namespaces).

Parameters `graph` (`pybel.BELGraph`) – A BEL graph

Returns A Counter from {namespace: frequency}

Return type `collections.Counter`

`pybel.struct.summary.get_namespaces(graph)`

Get the set of all namespaces used in this graph.

Parameters `graph` (`pybel.BELGraph`) – A BEL graph

Return type `Set[str]`

Returns A set of namespaces

`pybel.struct.summary.count_names_by_namespace(graph, namespace)`

Get the set of all of the names in a given namespace that are in the graph.

Parameters

- `graph` (`pybel.BELGraph`) – A BEL graph
- `namespace` (`str`) – A namespace keyword

Return type `Counter[str]`

Returns A counter from {name: frequency}

Raises `IndexError` – if the namespace is not defined in the graph.

`pybel.struct.summary.get_names_by_namespace(graph, namespace)`

Get the set of all of the names in a given namespace that are in the graph.

Parameters

- `graph` (`pybel.BELGraph`) – A BEL graph
- `namespace` (`str`) – A namespace keyword

Return type `Set[str]`

Returns A set of names belonging to the given namespace that are in the given graph

Raises `IndexError` – if the namespace is not defined in the graph.

`pybel.struct.summary.get_unused_namespaces(graph)`

Get the set of all namespaces that are defined in a graph, but are never used.

Parameters `graph` (`pybel.BELGraph`) – A BEL graph

Return type `Set[str]`

Returns A set of namespaces that are included but not used

`pybel.struct.summary.count_variants(graph)`

Count how many of each type of variant a graph has.

Parameters `graph` (`pybel.BELGraph`) – A BEL graph

Return type `Counter[str]`

`pybel.struct.summary.get_names(graph)`

Get all names for each namespace.

Return type `dict[str, set[str]]`

`pybel.struct.summary.count_pathologies(graph)`

Count the number of edges in which each pathology is incident.

Parameters `graph` (`pybel.BELGraph`) – A BEL graph

Return type `Counter[BaseEntity]`

`pybel.struct.summary.get_top_pathologies(graph, n=15)`

Get the top highest relationship-having edges in the graph by BEL.

Parameters

- `graph` (`pybel.BELGraph`) – A BEL graph
- `n` (`Optional[int]`) – The number of top connected pathologies to return. If None, returns all nodes

Return type `List[Tuple[BaseEntity, int]]`

`pybel.struct.summary.get_top_hubs(graph, *, n=15)`

Get the top hubs in the graph by BEL.

Parameters

- `graph` (`pybel.BELGraph`) – A BEL graph
- `n` (`Optional[int]`) – The number of top hubs to return. If None, returns all nodes

Return type `List[Tuple[BaseEntity, int]]`

`pybel.struct.summary.iterate_pubmed_identifiers(graph)`

Iterate over all PubMed identifiers in a graph.

Parameters `graph` (`pybel.BELGraph`) – A BEL graph

Return type `Iterable[str]`

Returns An iterator over the PubMed identifiers in the graph

`pybel.struct.summary.get_pubmed_identifiers(graph)`

Get the set of all PubMed identifiers cited in the construction of a graph.

Parameters `graph` (`pybel.BELGraph`) – A BEL graph

Return type `Set[str]`

Returns A set of all PubMed identifiers cited in the construction of this graph

`pybel.struct.summary.iter_annotation_value_pairs(graph)`

Iterate over the key/value pairs, with duplicates, for each annotation used in a BEL graph.

Parameters `graph` (`pybel.BELGraph`) – A BEL graph

Return type `Iterable[Tuple[str, str]]`

`pybel.struct.summary.iter_annotation_values(graph, annotation)`

Iterate over all of the values for an annotation used in the graph.

Parameters

- `graph` (`pybel.BELGraph`) – A BEL graph
- `annotation` (`str`) – The annotation to grab

Return type `Iterable[str]`

```
pybel.struct.summary.get_annotation_values_by_annotation(graph)
```

Get the set of values for each annotation used in a BEL graph.

Parameters `graph` (pybel.BELGraph) – A BEL graph

Return type Mapping[str, Set[str]]

Returns A dictionary of {annotation key: set of annotation values}

```
pybel.struct.summary.get_annotation_values(graph, annotation)
```

Get all values for the given annotation.

Parameters

- `graph` (pybel.BELGraph) – A BEL graph

- `annotation` (str) – The annotation to summarize

Return type Set[str]

Returns A set of all annotation values

```
pybel.struct.summary.count_relations(graph)
```

Return a histogram over all relationships in a graph.

Parameters `graph` (pybel.BELGraph) – A BEL graph

Return type Counter

Returns A Counter from {relation type: frequency}

```
pybel.struct.summary.get_annotations(graph)
```

Get the set of annotations used in the graph.

Parameters `graph` (pybel.BELGraph) – A BEL graph

Return type Set[str]

Returns A set of annotation keys

```
pybel.struct.summary.count_annotations(graph)
```

Count how many times each annotation is used in the graph.

Parameters `graph` (pybel.BELGraph) – A BEL graph

Return type Counter

Returns A Counter from {annotation key: frequency}

```
pybel.struct.summary.get_unused_annotations(graph)
```

Get the set of all annotations that are defined in a graph, but are never used.

Parameters `graph` (pybel.BELGraph) – A BEL graph

Return type Set[str]

Returns A set of annotations

```
pybel.struct.summary.get_unused_list_annotation_values(graph)
```

Get all of the unused values for list annotations.

Parameters `graph` (pybel.BELGraph) – A BEL graph

Return type Mapping[str, Set[str]]

Returns A dictionary of {str annotation: set of str values that aren't used}

```
pybel.struct.summary.get_metaedge_to_key(graph)
```

Get all edge types.

Return type `Mapping[Tuple[str, Optional[Tuple], Optional[Tuple]], Set[Tuple[BaseEntity, BaseEntity, str]]]`

CHAPTER
SIX

OPERATIONS

This page outlines operations that can be done to BEL graphs.

`pybel.struct.left_full_join(g, h)`
Add all nodes and edges from h to g, in-place for g.

Parameters

- `g` (`pybel.BELGraph`) – A BEL graph
- `h` (`pybel.BELGraph`) – A BEL graph

Example usage:

```
>>> import pybel
>>> g = pybel.from_bel_script('...')
>>> h = pybel.from_bel_script('...')
>>> left_full_join(g, h)
```

Return type

None

`pybel.struct.left_outer_join(g, h)`
Only add components from the h that are touching g.

Algorithm:

1. Identify all weakly connected components in h
2. Add those that have an intersection with the g

Parameters

- `g` (`BELGraph`) – A BEL graph
- `h` (`BELGraph`) – A BEL graph

Example usage:

```
>>> import pybel
>>> g = pybel.from_bel_script('...')
>>> h = pybel.from_bel_script('...')
>>> left_outer_join(g, h)
```

Return type

None

`pybel.struct.union(graphs, use_tqdm=False)`

Take the union over a collection of graphs into a new graph.

Assumes iterator is longer than 2, but not infinite.

Parameters

- **graphs** (`iter[BELGraph]`) – An iterator over BEL graphs. Can't be infinite.
- **use_tqdm** (`bool`) – Should a progress bar be displayed?

Returns A merged graph

Return type `BELGraph`

Example usage:

```
>>> import pybel
>>> g = pybel.from_bel_script('...')
>>> h = pybel.from_bel_script('...')
>>> k = pybel.from_bel_script('...')
>>> merged = union([g, h, k])
```

**CHAPTER
SEVEN**

FILTERS

This module contains functions for filtering node and edge iterables.

It relies heavily on the concepts of [functional programming](#) and the concept of [predicates](#).

**CHAPTER
EIGHT**

TRANSFORMATIONS

This module contains functions that mutate or make transformations on a network.

GROUPING

Functions for grouping BEL graphs into sub-graphs.

`pybel.struct.grouping.get_subgraphs_by_annotation(graph, annotation, sentinel=None)`

Stratify the given graph into sub-graphs based on the values for edges' annotations.

Parameters

- **graph** (`pybel.BELGraph`) – A BEL graph
- **annotation** (`str`) – The annotation to group by
- **sentinel** (`Optional[str]`) – The value to stick unannotated edges into. If none, does not keep undefined.

Return type `dict[str,pybel.BELGraph]`

`pybel.struct.grouping.get_subgraphs_by_citation(graph)`

Stratify the graph based on citations.

Return type `dict[tuple[str,str],pybel.BELGraph]`

PIPELINE

```
class pybel.Pipeline(protocol=None)
    Build and runs analytical pipelines on BEL graphs.
```

Example usage:

```
>>> from pybel import BELGraph
>>> from pybel.struct.pipeline import Pipeline
>>> from pybel.struct.mutation import enrich_protein_and_rna_origins, prune_
   protein_rna_origins
>>> graph = BELGraph()
>>> example = Pipeline()
>>> example.append(enrich_protein_and_rna_origins)
>>> example.append(prune_protein_rna_origins)
>>> result = example.run(graph)
```

Initialize the pipeline with an optional pre-defined protocol.

Parameters `protocol` (`Optional[Iterable[Dict]]`) – An iterable of dictionaries describing how to transform a network

static `from_functions` (`functions`)
Build a pipeline from a list of functions.

Parameters `functions` (`iter[((pybel.BELGraph) -> pybel.BELGraph) or ((pybel.BELGraph) -> None) or str]`) – A list of functions or names of functions

Example with function:

```
>>> from pybel.struct.pipeline import Pipeline
>>> from pybel.struct.mutation import remove_associations
>>> pipeline = Pipeline.from_functions([remove_associations])
```

Equivalent example with function names:

```
>>> from pybel.struct.pipeline import Pipeline
>>> pipeline = Pipeline.from_functions(['remove_associations'])
```

Lookup by name is possible for built in functions, and those that have been registered correctly using one of the four decorators:

1. `pybel.struct.pipeline.transformation()`,
2. `pybel.struct.pipeline.in_place_transformation()`,
3. `pybel.struct.pipeline.uni_transformation()`,

4. `pybel.struct.pipeline.uni_in_place_transformation()`,

Return type `Pipeline`

append (`name, *args, **kwargs`)

Add a function (either as a reference, or by name) and arguments to the pipeline.

Parameters

- **name** (`str` or `(pybel.BELGraph -> pybel.BELGraph)`) – The name of the function
- **args** – The positional arguments to call in the function
- **kwargs** – The keyword arguments to call in the function

Return type `Pipeline`

Returns This pipeline for fluid query building

Raises `MissingPipelineFunctionError` – If the function is not registered

extend (`protocol`)

Add another pipeline to the end of the current pipeline.

Parameters `protocol` (`Union[Iterable[Dict], Pipeline]`) – An iterable of dictionaries (or another Pipeline)

Return type `Pipeline`

Returns This pipeline for fluid query building

Example: `>>> p1 = Pipeline.from_functions(['enrich_protein_and_rna_origins']) >>> p2 = Pipeline.from_functions(['remove_pathologies']) >>> p1.extend(p2)`

run (`graph, universe=None`)

Run the contained protocol on a seed graph.

Parameters

- **graph** (`pybel.BELGraph`) – The seed BEL graph
- **universe** (`pybel.BELGraph`) – Allows just-in-time setting of the universe in case it wasn't set before. Defaults to the given network.

Returns The new graph is returned if not applied in-place

Return type `pybel.BELGraph`

to_json ()

Return this pipeline as a JSON list.

Return type `List`

dumps (`**kwargs`)

Dump this pipeline as a JSON string.

Return type `str`

dump (`file, **kwargs`)

Dump this protocol to a file in JSON.

Return type `None`

static from_json (`data`)

Build a pipeline from a JSON list.

Return type Pipeline

static load(*file*)
Load a protocol from JSON contained in file.

Return type Pipeline

Returns The pipeline represented by the JSON in the file

Raises *MissingPipelineFunctionError* – If any functions are not registered

static loads(*s*)
Load a protocol from a JSON string.

Parameters *s* (`str`) – A JSON string

Return type Pipeline

Returns The pipeline represented by the JSON in the file

Raises *MissingPipelineFunctionError* – If any functions are not registered

static union(*pipelines*)
Take the union of multiple pipelines.

Parameters *pipelines* (`Iterable[Pipeline]`) – A list of pipelines

Return type Pipeline

Returns The union of the results from multiple pipelines

static intersection(*pipelines*)
Take the intersection of the results from multiple pipelines.

Parameters *pipelines* (`Iterable[Pipeline]`) – A list of pipelines

Return type Pipeline

Returns The intersection of results from multiple pipelines

10.1 Transformation Decorators

This module contains the functions for decorating transformation functions.

A transformation function takes in a `pybel.BELGraph` and either returns None (in-place) or a new `pybel.BELGraph` (out-of-place).

`pybel.struct.pipeline.decorators.in_place_transformation(func)`
A decorator for functions that modify BEL graphs in-place

`pybel.struct.pipeline.decorators.uni_in_place_transformation(func)`
A decorator for functions that require a “universe” graph and modify BEL graphs in-place

`pybel.struct.pipeline.decorators.uni_transformation(func)`
A decorator for functions that require a “universe” graph and create new BEL graphs from old BEL graphs

`pybel.struct.pipeline.decorators.transformation(func)`
A decorator for functions that create new BEL graphs from old BEL graphs

`pybel.struct.pipeline.decorators.get_transformation(name)`
Get a transformation function and error if its name is not registered.

Parameters *name* (`str`) – The name of a function to look up

Returns A transformation function

Raises `MissingPipelineFunctionError` – If the given function name is not registered

10.2 Exceptions

Exceptions for the `pybel.struct.pipeline` module.

`exception pybel.struct.pipeline.exc.MissingPipelineFunctionError`

Raised when trying to run the pipeline with a function that isn't registered.

`exception pybel.struct.pipeline.exc.MetaValueError`

Raised when getting an invalid meta value.

`exception pybel.struct.pipeline.exc.MissingUniverseError`

Raised when running a universe function without a universe being present.

`exception pybel.struct.pipeline.exc.DeprecationMappingError`

Raised when applying the deprecation function annotation and the given name already is being used.

`exception pybel.struct.pipeline.exc.PipelineNameError`

Raised when a second function tries to use the same name.

INPUT AND OUTPUT

Input and output functions for BEL graphs.

PyBEL provides multiple lossless interchange options for BEL. Lossy output formats are also included for convenient export to other programs. Notably, a *de facto* interchange using Resource Description Framework (RDF) to match the ability of other existing software is excluded due the immaturity of the BEL to RDF mapping.

`pybel.load(path, **kwargs)`

Read a BEL graph.

Parameters

- **path** (`str`) – The path to a BEL graph in any of the formats with extensions described below
- **kwargs** – The keyword arguments are passed to the importer function

Return type `BELGraph`

Returns A BEL graph.

This is the universal loader, which means any file path can be given and PyBEL will look up the appropriate load function. Allowed extensions are:

- bel
- bel.nodelink.json
- bel.cx.json
- bel.jgif.json

The previous extensions also support gzipping. Other allowed extensions that don't support gzip are:

- bel.pickle / bel.gpickle / bel.pkl
- indra.json

`pybel.dump(graph, path, **kwargs)`

Write a BEL graph.

Parameters

- **graph** (`BELGraph`) – A BEL graph
- **path** (`str`) – The path to which the BEL graph is written.
- **kwargs** – The keyword arguments are passed to the exporter function

This is the universal loader, which means any file path can be given and PyBEL will look up the appropriate writer function. Allowed extensions are:

- bel

- bel.nodelink.json
- bel.unodelink.json
- bel.cx.json
- bel.jgif.json
- bel.graphdati.json

The previous extensions also support gzipping. Other allowed extensions that don't support gzip are:

- bel.pickle / bel.gpickle / bel.pkl
- indra.json
- tsv
- gsea

Return type None

11.1 Import

11.1.1 Parsing Modes

The PyBEL parser has several modes that can be enabled and disabled. They are described below.

Allow Naked Names

By default, this is set to `False`. The parser does not allow identifiers that are not qualified with namespaces (*naked names*), like in `p(YFG)`. A proper namespace, like `p(HGNC:YFG)` must be used. By setting this to `True`, the parser becomes permissive to naked names. In general, this is bad practice and this feature will be removed in the future.

Allow Nested

By default, this is set to `False`. The parser does not allow nested statements is disabled. See *overview*. By setting this to `True` the parser will accept nested statements one level deep.

Citation Clearing

By default, this is set to `True`. While the BEL specification clearly states how the language should be used as a state machine, many BEL documents do not conform to the strict SET/UNSET rules. To guard against annotations accidentally carried from one set of statements to the next, the parser has two modes. By default, in citation clearing mode, when a `SET CITATION` command is reached, it will clear all other annotations (except the `STATEMENT_GROUP`, which has higher priority). This behavior can be disabled by setting this to `False` to re-enable strict parsing.

11.1.2 Reference

```
pybel.from_bel_script(path, **kwargs)
```

Load a BEL graph from a file resource. This function is a thin wrapper around `from_lines()`.

Parameters `path` (`Union[str, TextIO]`) – A path or file-like

The remaining keyword arguments are passed to `pybel.io.line_utils.parse_lines()`, which populates a `BELGraph`.

Return type `BELGraph`

```
pybel.from_bel_script_url(url, **kwargs)
```

Load a BEL graph from a URL resource.

Parameters `url` (`str`) – A valid URL pointing to a BEL document

The remaining keyword arguments are passed to `pybel.io.line_utils.parse_lines()`.

Return type `BELGraph`

```
pybel.to_bel_script(graph, path, use_identifiers=True)
```

Write the `BELGraph` as a canonical BEL script.

Parameters

- `graph` (`BELGraph`) – the BEL Graph to output as a BEL Script
- `path` (`Union[str, TextIO]`) – A path or file-like.
- `use_identifiers` (`bool`) – Enables extended BEP-0008 syntax

Return type `None`

11.1.3 Hetionet

Importer for Hetionet JSON.

```
pybel.from_hetionet_json(hetionet_dict, use_tqdm=True)
```

Convert a Hetionet dictionary to a BEL graph.

Return type `BELGraph`

```
pybel.from_hetionet_file(file)
```

Get Hetionet from a JSON file.

Return type `BELGraph`

```
pybel.from_hetionet_gz(path)
```

Get Hetionet from its JSON GZ file.

Return type `BELGraph`

```
pybel.get_hetionet()
```

Get Hetionet from GitHub, cache, and convert to BEL.

Return type `BELGraph`

11.2 Transport

All transport pairs are reflective and data-preserving.

11.2.1 Bytes

Conversion functions for BEL graphs with bytes and Python pickles.

`pybel.from_bytes(bytes_graph, check_version=True)`
Read a graph from bytes (the result of pickling the graph).

Parameters

- `bytes_graph` (`bytes`) – File or filename to write
- `check_version` (`bool`) – Checks if the graph was produced by this version of PyBEL

Return type `BELGraph`

`pybel.to_bytes(graph, protocol=4)`
Convert a graph to bytes with pickle.

Note that the pickle module has some incompatibilities between Python 2 and 3. To export a universally importable pickle, choose 0, 1, or 2.

Parameters

- `graph` (`BELGraph`) – A BEL network
- `protocol` (`int`) – Pickling protocol to use. Defaults to `HIGHEST_PROTOCOL`.

See also:

<https://docs.python.org/3.6/library/pickle.html#data-stream-format>

Return type `bytes`

`pybel.from_pickle(path, check_version=True)`
Read a graph from a pickle file.

Parameters

- `path` (`Union[str, BinaryIO]`) – File or filename to read. Filenames ending in `.gz` or `.bz2` will be uncompressed.
- `check_version` (`bool`) – Checks if the graph was produced by this version of PyBEL

Return type `BELGraph`

`pybel.to_pickle(graph, path, protocol=4)`
Write this graph to a pickle object with `networkx.write_gpickle()`.

Note that the pickle module has some incompatibilities between Python 2 and 3. To export a universally importable pickle, choose 0, 1, or 2.

Parameters

- `graph` (`BELGraph`) – A BEL graph
- `path` (`Union[str, BinaryIO]`) – A path or file-like
- `protocol` (`int`) – Pickling protocol to use. Defaults to `HIGHEST_PROTOCOL`.

See also:

<https://docs.python.org/3.6/library/pickle.html#data-stream-format>

Return type None

11.2.2 Node-Link JSON

Conversion functions for BEL graphs with node-link JSON.

`pybel.from_nodelink(graph_json_dict, check_version=True)`

Build a graph from node-link JSON Object.

Return type BELGraph

`pybel.to_nodelink(graph)`

Convert this graph to a node-link JSON object.

Parameters `graph` (BELGraph) – BEL Graph

Return type Mapping[str, Any]

`pybel.from_nodelink_jsons(graph_json_str, check_version=True)`

Read a BEL graph from a node-link JSON string.

Return type BELGraph

`pybel.to_nodelink_jsons(graph, **kwargs)`

Dump this graph as a node-link JSON object to a string.

Return type str

`pybel.from_nodelink_file(path, check_version=True)`

Build a graph from the node-link JSON contained in the given file.

Parameters `path` (Union[str, TextIO]) – A path or file-like

Return type BELGraph

`pybel.to_nodelink_file(graph, path, **kwargs)`

Write this graph as node-link JSON to a file.

Parameters

- `graph` (BELGraph) – A BEL graph
- `path` (Union[str, TextIO]) – A path or file-like

Return type None

`pybel.from_nodelink_gz(path)`

Read a graph as node-link JSON from a gzip file.

Return type BELGraph

`pybel.to_nodelink_gz(graph, path, **kwargs)`

Write a graph as node-link JSON to a gzip file.

Return type None

11.2.3 Cyberinfrastructure Exchange

This module wraps conversion between `pybel.BELGraph` and the Cyberinfrastructure Exchange (CX) JSON.

CX is an aspect-oriented network interchange format encoded in JSON with a format inspired by the JSON-LD encoding of Resource Description Framework (RDF). It is primarily used by the Network Data Exchange (NDEEx) and more recent versions of Cytoscape.

See also:

- The NDEEx Data Model Specification
- [Cytoscape.js](#)
- CX Support for Cytoscape.js on the Cytoscape [App Store](#)

`pybel.from_cx(cx)`

Rebuild a BELGraph from CX JSON output from PyBEL.

Parameters `cx` (`List[Dict]`) – The CX JSON object for this graph

Return type `BELGraph`

`pybel.to_cx(graph)`

Convert a BEL Graph to a CX JSON object for use with [NDEEx](#).

See also:

- [NDEEx Python Client](#)

Return type `List[Dict]`

`pybel.from_cx_jsons(graph_json_str)`

Read a BEL graph from a CX JSON string.

Return type `BELGraph`

`pybel.to_cx_jsons(graph, **kwargs)`

Dump this graph as a CX JSON object to a string.

Return type `str`

`pybel.from_cx_file(path)`

Read a file containing CX JSON and converts to a BEL graph.

Parameters `path` (`Union[str, TextIO]`) – A readable file or file-like containing the CX JSON for this graph

Return type `BELGraph`

Returns A BEL Graph representing the CX graph contained in the file

`pybel.to_cx_file(graph, path, indent=2, **kwargs)`

Write a BEL graph to a JSON file in CX format.

Parameters

- `graph` (`BELGraph`) – A BEL graph
- `path` (`Union[str, TextIO]`) – A writable file or file-like
- `indent` (`Optional[int]`) – How many spaces to use to pretty print. Change to None for no pretty printing

The example below shows how to output a BEL graph as CX to an open file.

```
from pybel.examples import sialic_acid_graph
from pybel import to_cx_file
with open('graph.bel.cx.json', 'w') as file:
    to_cx_file(sialic_acid_graph, file)
```

The example below shows how to output a BEL graph as CX to a file at a given path.

```
from pybel.examples import sialic_acid_graph
from pybel import to_cx_file
to_cx_file(sialic_acid_graph, 'graph.bel.cx.json')
```

If you have a big graph, you might consider storing it as a gzipped JGIF file by using `to_cx_gz()`.

Return type None

`pybel.from_cx_gz(path)`

Read a graph as CX JSON from a gzip file.

Return type `BELGraph`

`pybel.to_cx_gz(graph, path, **kwargs)`

Write a graph as CX JSON to a gzip file.

Return type None

11.2.4 JSON Graph Interchange Format

Conversion functions for BEL graphs with JGIF JSON.

The JSON Graph Interchange Format (JGIF) is specified similarly to the Node-Link JSON. Interchange with this format provides compatibility with other software and repositories, such as the Causal Biological Network Database.

`pybel.from_jgif(graph_jgif_dict, parser_kwargs=None)`

Build a BEL graph from a JGIF JSON object.

Parameters `graph_jgif_dict (dict)` – The JSON object representing the graph in JGIF format

Return type `BELGraph`

`pybel.to_jgif(graph)`

Build a JGIF dictionary from a BEL graph.

Parameters `graph (pybel.BELGraph)` – A BEL graph

Returns A JGIF dictionary

Return type `dict`

Warning: Untested! This format is not general purpose and is therefore time is not heavily invested. If you want to use Cytoscape.js, we suggest using `pybel.to_cx()` instead.

The example below shows how to output a BEL graph as a JGIF dictionary.

```
import os
from pybel.examples import sialic_acid_graph
graph_jgif_json = pybel.to_jgif(sialic_acid_graph)
```

If you want to write the graph directly to a file as JGIF, see func:`to_jgif_file`.

```
pybel.from_jgif_jsons(graph_json_str)
```

Read a BEL graph from a JGIF JSON string.

Return type BELGraph

```
pybel.to_jgif_jsons(graph, **kwargs)
```

Dump this graph as a JGIF JSON object to a string.

Return type str

```
pybel.from_jgif_file(path)
```

Build a graph from the JGIF JSON contained in the given file.

Parameters path (Union[str, TextIO]) – A path or file-like

Return type BELGraph

```
pybel.to_jgif_file(graph, file, **kwargs)
```

Write JGIF to a file.

Parameters

- **graph** (BELGraph) – A BEL graph
- **file** (Union[str, TextIO]) – A writable file or file-like

The example below shows how to output a BEL graph as JGIF to an open file.

```
from pybel.examples import sialic_acid_graph
from pybel import to_jgif_file
with open('graph.bel.jgif.json', 'w') as file:
    to_jgif_file(sialic_acid_graph, file)
```

The example below shows how to output a BEL graph as JGIF to a file at a given path.

```
from pybel.examples import sialic_acid_graph
from pybel import to_jgif_file
to_jgif_file(sialic_acid_graph, 'graph.bel.jgif.json')
```

If you have a big graph, you might consider storing it as a gzipped JGIF file by using `to_jgif_gz()`.

Return type None

```
pybel.from_jgif_gz(path)
```

Read a graph as JGIF JSON from a gzip file.

Return type BELGraph

```
pybel.to_jgif_gz(graph, path, **kwargs)
```

Write a graph as JGIF JSON to a gzip file.

Return type None

```
pybel.post_jgif(graph, url, **kwargs)
```

Post the JGIF to a given URL.

Return type Response

```
pybel.from_cbn_jgif(graph_jgif_dict)
```

Build a BEL graph from CBN JGIF.

Map the JGIF used by the Causal Biological Network Database to standard namespace and annotations, then builds a BEL graph using `pybel.from_jgif()`.

Parameters `graph_jgif_dict` (`dict`) – The JSON object representing the graph in JGIF format

Return type `BELGraph`

Example: .. code-block:: python

```
import requests from pybel import from_cbn_jgif apoptosis_url = 'http://causalbionet.com/Networks/GetJSONGraphFile?networkId=810385422' graph_jgif_dict = requests.get(apoptosis_url).json() graph = from_cbn_jgif(graph_jgif_dict)
```

Warning: Handling the annotations is not yet supported, since the CBN documents do not refer to the resources used to create them. This may be added in the future, but the annotations must be stripped from the graph before uploading to the network store using `pybel.struct.mutation.strip_annotations()`.

`pybel.from_cbn_jgif_file(path)`

Build a graph from a file containing the CBN variant of JGIF.

Parameters `path` (`Union[str, TextIO]`) – A path or file-like

Return type `BELGraph`

11.2.5 GraphDati

Conversion functions for BEL graphs with GraphDati.

Note that these are not exact I/O - you can't currently use them as a round trip because the input functions expect the GraphDati format that's output by BioDati.

`pybel.to_graphdati(graph, *, use_identifiers=True, skip_unqualified=True, use_tqdm=False, metadata_extras=None)`

Export a GraphDati list using the nanopub.

Parameters

- `graph` – A BEL graph
- `use_identifiers` (`bool`) – use OBO-style identifiers
- `use_tqdm` (`bool`) – Show a progress bar while generating nanopubs
- `skip_unqualified` (`bool`) – Should unqualified edges be output as nanopubs? Defaults to false.
- `metadata_extras` (`Optional[Mapping[str, Any]]`) – Extra information to pass into the metadata part of nanopubs

Return type `List[Mapping[str, Mapping[str, Any]]]`

`pybel.from_graphdati(j, use_tqdm=True)`

Convert data from the “normal” network format.

Warning: BioDati crashes when requesting the `full` network format, so this isn't yet explicitly supported

Return type `BELGraph`

```
pybel.to_graphdati_file(graph, path, use_identifiers=True, **kwargs)
```

Write this graph as GraphDati JSON to a file.

Parameters

- **graph** (BELGraph) – A BEL graph
- **path** (Union[str, TextIO]) – A path or file-like

Return type None

```
pybel.from_graphdati_file(path)
```

Load a file containing GraphDati JSON.

Parameters **path** (Union[str, TextIO]) – A path or file-like

Return type BELGraph

```
pybel.to_graphdati_gz(graph, path, **kwargs)
```

Write a graph as GraphDati JSON to a gzip file.

Return type None

```
pybel.from_graphdati_gz(path)
```

Read a graph as GraphDati JSON from a gzip file.

Return type BELGraph

```
pybel.to_graphdati_jsons(graph, **kwargs)
```

Dump this graph as a GraphDati JSON object to a string.

Parameters **graph** (BELGraph) – A BEL graph

Return type str

```
pybel.from_graphdati_jsons(s)
```

Load a graph from a GraphDati JSON string.

Parameters **graph** – A BEL graph

Return type BELGraph

```
pybel.to_graphdati_jsonl(graph, file, use_identifiers=True, use_tqdm=True)
```

Write this graph as a GraphDati JSON lines file.

Parameters **graph** – A BEL graph

```
pybel.to_graphdati_jsonl_gz(graph, path, **kwargs)
```

Write a graph as GraphDati JSONL to a gzip file.

Parameters **graph** (BELGraph) – A BEL graph

Return type None

11.2.6 INDRA

Conversion functions for BEL graphs with INDRA.

After assembling a model with [INDRA](#), a list of `indra.statements.Statement` can be converted to a `pybel.BELGraph` with `indra.assemblers.pybel.PybelAssembler`.

```

from indra.assemblers.pybel import PybelAssembler
import pybel

stmts = [
    # A list of INDRA statements
]

pba = PybelAssembler(
    stmts,
    name='Graph Name',
    version='0.0.1',
    description='Graph Description'
)
graph = pba.make_model()

# Write to BEL file
pybel.to_bel_path(belgraph, 'simple_pybel.bel')

```

Warning: These functions are hard to unit test because they rely on a whole set of java dependencies and will likely not be for a while.

`pybel.from_indra_statements(stmts, name=None, version=None, description=None, authors=None, contact=None, license=None, copyright=None, disclaimer=None)`

Import a model from indra.

Parameters

- `stmts` (`List[indra.statements.Statement]`) – A list of statements
- `name` (`Optional[str]`) – The graph's name
- `version` (`Optional[str]`) – The graph's version. Recommended to use semantic versioning or YYYYMMDD format.
- `description` (`Optional[str]`) – The description of the graph
- `authors` (`Optional[str]`) – The authors of this graph
- `contact` (`Optional[str]`) – The contact email for this graph
- `license` (`Optional[str]`) – The license for this graph
- `copyright` (`Optional[str]`) – The copyright for this graph
- `disclaimer` (`Optional[str]`) – The disclaimer for this graph

Return type `pybel.BELGraph`

`pybel.from_indra_statements_json(stmts_json, **kwargs)`

Get a BEL graph from INDRA statements JSON.

Return type `BELGraph`

Other kwargs are passed to `from_indra_statements()`.

`pybel.from_indra_statements_json_file(file, **kwargs)`

Get a BEL graph from INDRA statements JSON file.

Return type `BELGraph`

Other kwargs are passed to `from_indra_statements()`.

`pybel.to_indra_statements(graph)`

Export this graph as a list of INDRA statements using the `indra.sources.pybel.PybelProcessor`.

Parameters `graph` (`pybel.BELGraph`) – A BEL graph

Return type `list[indra.statements.Statement]`

`pybel.to_indra_statements_json(graph)`

Export this graph as INDRA JSON list.

Parameters `graph` (`pybel.BELGraph`) – A BEL graph

Return type `List[Mapping[str, Any]]`

`pybel.to_indra_statements_json_file(graph, path, indent=2, **kwargs)`

Export this graph as INDRA statement JSON.

Parameters

- `graph` (`pybel.BELGraph`) – A BEL graph

- `path` (`Union[str, TextIO]`) – A writable file or file-like

Other kwargs are passed to `json.dump()`.

`pybel.from_biopax(path, **kwargs)`

Import a model encoded in Pathway Commons BioPAX via `indra`.

Parameters `path` (`str`) – Path to a BioPAX OWL file

Return type `pybel.BELGraph`

Other kwargs are passed to `from_indra_statements()`.

Warning: Not compatible with all BioPAX! See INDRA documentation.

11.3 Visualization

11.3.1 Jupyter

Support for displaying BEL graphs in Jupyter notebooks.

`pybel.io.jupyter.to_jupyter(graph, width=1000, height=650, color_map=None)`

Display a BEL graph inline in a Jupyter notebook.

To use successfully, make run as the last statement in a cell inside a Jupyter notebook.

Parameters

- `graph` (`BELGraph`) – A BEL graph

- `width` (`int`) – The width of the visualization window to render

- `height` (`int`) – The height of the visualization window to render

- `color_map` (`Optional[Mapping[str, str]]`) – A dictionary from PyBEL internal node functions to CSS color strings like `#FFEE00`. Defaults to `default_color_map`

Returns An IPython notebook Javascript object

Return type `IPython.display.Javascript`

11.4 Analytical Services

11.4.1 PyNPA

Exporter for PyNPA.

See also:

<https://github.com/pynpa>

`pybel.to_npa_directory(graph, directory, **kwargs)`

Write the BEL file to two files in the directory for pynpa.

Return type `None`

`pybel.to_npa_dfs(graph, cartesian_expansion=False, nomenclature_method_first_layer=None, nomenclature_method_second_layer=None, direct_tf_only=False)`

Export the BEL graph as two lists of triples for the pynpa.

Parameters

- **graph** (`BELGraph`) – A BEL graph
- **cartesian_expansion** (`bool`) – If true, applies cartesian expansion on both reactions (reactants x products) as well as list abundances using `list_abundance_cartesian_expansion()` and `reaction_cartesian_expansion()`
- **nomenclature_method_first_layer** (`Optional[str]`) – Either “curie”, “name” or “inodes. Defaults to “curie”.
- **nomenclature_method_second_layer** (`Optional[str]`) – Either “curie”, “name” or “inodes. Defaults to “curie”.

1. Pick out all transcription factor relationships. Protein X is a transcription factor for gene Y IFF $\text{complex(p(X), g(Y))} \rightarrow r(Y)$
2. Get all other interactions between any gene/rna/protein that are directed causal for the PPI layer

Return type `Tuple[DataFrame, DataFrame]`

11.4.2 HiPathia

Convert a BEL graph to HiPathia inputs.

Input

SIF File

- Text file with three columns separated by tabs.
- Each row represents an interaction in the pathway. First column is the source node, third column the target node, and the second is the type of relation between them.
- Only activation and inhibition interactions are allowed.
- The name of the nodes in this file will be stored as the IDs of the nodes.
- The nodes IDs should have the following structure: N (dash) pathway ID (dash) node ID.

- HiPathia distinguish between two types of nodes: simple and complex.

Simple nodes:

- Simple nodes may include many genes, but only one is needed to perform the function of the node. This could correspond to a protein family of enzymes that all have the same function - only one of them needs to be present for the action to take place. Simple nodes are defined within
- Node IDs from simple nodes do not include any space, i.e. N-hsa04370-11.

Complex nodes:

- Complex nodes include different simple nodes and represent protein complexes. Each simple node within the complex represents one protein in the complex. This node requires the presence of all their simple nodes to perform its function.
- Node IDs from complex nodes are the juxtaposition of the included simple node IDs, separated by spaces, i.e. N-hsa04370-10 26.

ATT File

Text file with twelve (12) columns separated by tabulars. Each row represents a node (either simple or complex).

The columns included are:

1. `ID`: Node ID as explained above.
2. `label`: Name to be shown in the picture of the pathway en HGNC. Generally, the gene name of the first included EntrezID gene is used as label. For complex nodes, we juxtapose the gene names of the first genes of each simple node included (see `genesList` column below).
3. `X`: The X-coordinate of the position of the node in the pathway.
4. `Y`: The Y-coordinate of the position of the node in the pathway.
5. `color`: The default color of the node.
6. `shape`: The shape of the node. “rectangle” should be used for genes and “circle” for metabolites.
7. `type`: The type of the node, either “gene” for genes or “compound” for metabolites. For complex nodes, the type of each of their included simple nodes is juxtaposed separated by commas, i.e. gene,gene.
8. `label.cex`: Amount by which plotting label should be scaled relative to the default.
9. `label.color`: Default color of the node.
10. `width`: Default width of the node.
11. `height`: Default height of the node.
12. `genesList`: List of genes included in each node, with EntrezID:
 - Simple nodes: EntrezIDs of the genes included, separated by commas (“,”) and no spaces, i.e. 56848,8877 for node N-hsa04370-11.
 - Complex nodes: GenesList of the simple nodes included, separated by a slash (“/”) and no spaces, and in the same order as in the node ID. For example, node N-hsa04370-10 26 includes two simple nodes: 10 and 26. Its `genesList` column is 5335,5336,/,9047, meaning that the genes included in node 10 are 5335 and 5336, and the gene included in node 26 is 9047.

`pybel.to_hipathia(graph, directory)`
Export HiPathia artifacts for the graph.

Return type None

```
pybel.to_hipathia_dfs(graph)
```

Get the ATT and SIF dataframes.

1. Identify nodes: 1. Identify all proteins 2. Identify all protein families 3. Identify all complexes with just a protein or a protein family in them
2. Identify interactions between any of those things that are causal
3. Profit!

Return type Tuple[DataFrame, DataFrame]

```
pybel.from_hipathia_paths(name, att_path, sif_path)
```

Get a BEL graph from HiPathia files.

Return type BELGraph

```
pybel.from_hipathia_dfs(name, att_df, sif_df)
```

Get a BEL graph from HiPathia dataframes.

Return type BELGraph

11.4.3 SPIA

An exporter for signaling pathway impact analysis (SPIA) described by [Tarcă2009].

See also:

<https://bioconductor.org/packages/release/bioc/html/SPIA.html>

```
pybel.to_spia_dfs(graph)
```

Create an excel sheet ready to be used in SPIA software.

Parameters `graph` (BELGraph) – BELGraph

Return type Mapping[str, DataFrame]

Returns dictionary with matrices

```
pybel.to_spia_excel(graph, path)
```

Write the BEL graph as an SPIA-formatted excel sheet at the given path.

Return type None

```
pybel.to_spia_tsvs(graph, directory)
```

Write the BEL graph as a set of SPIA-formatted TSV files in a given directory.

Return type None

11.4.4 Machine Learning

Export functions for Machine Learning.

While BEL is a fantastic medium for storing metadata and high granularity information on edges, machine learning algorithms can not consume BEL graphs directly. This module provides functions that make inferences and interpretations of BEL graphs in order to interface with machine learning platforms. One example where we've done this is BioKEEN, which uses this module to convert BEL graphs into a format for knowledge graph embeddings.

```
pybel.to_tsv(graph, path, *, use_tqdm=False, sep='\t', raise_on_none=False)
```

Write the graph as a TSV.

Parameters

- **graph** (BELGraph) – A BEL graph
- **path** (Union[str, TextIO]) – A path or file-like
- **use_tqdm** (bool) – Should a progress bar be shown?
- **sep** – The separator to use
- **raise_on_none** (bool) – Should an exception be raised if no triples are returned?

Raises NoTriplesValueError

Return type None

`pybel.to_edgelist(graph, path, *, use_tqdm=False, sep='\t', raise_on_none=False)`

Write the graph as an edgelist.

Parameters

- **graph** (BELGraph) – A BEL graph
- **path** (Union[str, TextIO]) – A path or file-like
- **use_tqdm** (bool) – Should a progress bar be shown?
- **sep** – The separator to use
- **raise_on_none** (bool) – Should an exception be raised if no triples are returned?

Raises NoTriplesValueError

Return type None

11.5 Web Services

11.5.1 BEL Commons

Transport functions for BEL Commons.

BEL Commons is a free, open-source platform for hosting BEL content. Because it was originally developed and published in an academic capacity at Fraunhofer SCAI, a public instance can be found at <https://bel-commons-dev.scai.fraunhofer.de>. However, this instance is only supported out of posterity and will not be updated. If you would like to host your own instance of BEL Commons, there are instructions on its GitHub page.

`pybel.from_bel_commons(network_id, host=None)`

Retrieve a public network from BEL Commons.

In the future, this function may be extended to support authentication.

Parameters

- **network_id** (int) – The BEL Commons network identifier
- **host** (Optional[str]) – The location of the BEL Commons server. Alternatively, looks up in PyBEL config with PYBEL_REMOTE_HOST or the environment as PYBEL_REMOTE_HOST Defaults to `pybel.constants.DEFAULT_SERVICE_URL`

Return type BELGraph

`pybel.to_bel_commons(graph, host=None, user=None, password=None, public=False)`

Send a graph to the receiver service and returns the `requests` response object.

Parameters

- **graph** (BELGraph) – A BEL graph
- **host** (Optional[str]) – The location of the BEL Commons server. Alternatively, looks up in PyBEL config with PYBEL_REMOTE_HOST or the environment as PYBEL_REMOTE_HOST Defaults to `pybel.constants.DEFAULT_SERVICE_URL`
- **user** (Optional[str]) – Username for BEL Commons. Alternatively, looks up in PyBEL config with PYBEL_REMOTE_USER or the environment as PYBEL_REMOTE_USER
- **password** (Optional[str]) – Password for BEL Commons. Alternatively, looks up in PyBEL config with PYBEL_REMOTE_PASSWORD or the environment as PYBEL_REMOTE_PASSWORD

Return type Response

Returns The response object from `requests`

11.5.2 BioDati

Transport functions for BioDati.

BioDati is a paid, closed-source platform for hosting BEL content. However, they do have a demo instance running at <https://studio.demo.biодати.com> with which the examples in this module will be described.

As noted in the transport functions for BioDati, you should change the URLs to point to your own instance of BioDati. If you’re looking for an open source storage system for hosting your own BEL content, you may consider [BEL Commons](#), with the caveat that it is currently maintained in an academic capacity. Disclosure: BEL Commons is developed by the developers of PyBEL.

```
pybel.to_biodati(graph, *, username='demo@biодати.com', password='demo',
                  base_url='https://nanopubstore.demo.biодати.com',
                  chunksize=None,
                  use_tqdm=True, collections=None, overwrite=False, validate=True, email=False)
```

Post this graph to a BioDati server.

Parameters

- **graph** (BELGraph) – A BEL graph
- **username** (str) – The email address to log in to BioDati. Defaults to “`demo@biодати.com`” for the demo server
- **password** (str) – The password to log in to BioDati. Defaults to “`demo`” for the demo server
- **base_url** (str) – The BioDati nanopub store base url. Defaults to “<https://nanopubstore.demo.biодати.com>” for the demo server’s nanopub store
- **chunksize** (Optional[int]) – The number of nanopubs to post at a time. By default, does all.
- **use_tqdm** (bool) – Should tqdm be used when iterating?
- **collections** (Optional[Iterable[str]]) – Tags to add to the nanopubs for lookup on BioDati
- **overwrite** (bool) – Set the BioDati upload “`overwrite`” setting
- **validate** (bool) – Set the BioDati upload “`validate`” setting
- **email** (Union[bool, str]) – Who should get emailed with results about the upload? If true, emails to user used for login. If string, emails to that user. If false, no email.

Return type Response

Returns The response from the BioDati server (last response if using chunking)

Warning: BioDati does not support large uploads (yet?).

Warning: The default public BioDati server has been put here. You should switch it to yours. It will look like `https://nanopubstore.<YOUR NAME>.biodati.com`.

```
pybel.from_biodati(network_id, username='demo@biodati.com', password='demo',  
                   base_url='https://networkstore.demo.biodati.com')
```

Get a graph from a BioDati network store based on its network identifier.

Parameters

- **network_id** (`str`) – The internal identifier of the network you want to download.
- **username** (`str`) – The email address to log in to BioDati. Defaults to “`demo@biodati.com`” for the demo server
- **password** (`str`) – The password to log in to BioDati. Defaults to “`demo`” for the demo server
- **base_url** (`str`) – The BioDati network store base url. Defaults to “`https://networkstore.demo.biodati.com`” for the demo server’s network store

Example usage:

```
from pybel import from_biodati  
network_id = '01E46GDFQAGK5W8EFS9S9WMH12' # COVID-19 graph example from Wendy  
    ↪ Zimmermann  
graph = from_biodati(  
    network_id=network_id,  
    username='demo@biodati.com',  
    password='demo',  
    base_url='https://networkstore.demo.biodati.com',  
)  
graph.summarize()
```

Warning: The default public BioDati server has been put here. You should switch it to yours. It will look like `https://networkstore.<YOUR NAME>.biodati.com`.

Return type `BELGraph`

11.5.3 Fraunhofer OrientDB

Transport functions for Franhofer’s OrientDB.

Fraunhofer hosts an instance of `OrientDB` that contains BEL in a schema similar to `pybel.io.umbrella_nodelink`. However, they include custom relations that do not come from a controlled vocabulary, and have not made the schema, ETL scripts, or documentation available.

Unlike BioDati and BEL Commons, the Fraunhofer OrientDB does not allow for uploads, so only a single function `pybel.from_fraunhofer_orientdb()` is provided by PyBEL.

```
pybel.from_fraunhofer_orientdb(database='covid',      user='covid_user',      password='covid',
                                query=None)
```

Get a BEL graph from the Fraunhofer OrientDB.

Parameters

- **database** (`str`) – The OrientDB database to connect to
- **user** (`str`) – The user to connect to OrientDB
- **password** (`str`) – The password to connect to OrientDB
- **query** (`Optional[str]`) – The query to run. Defaults to the URL encoded version of `select from E`, where E is all edges in the OrientDB edge database. Likely does not need to be changed, except in the case of selecting specific subsets of edges. Make sure you URL encode it properly, because OrientDB’s RESTful API puts it in the URL’s path.

By default, this function connects to the `covid` database, that corresponds to the COVID-19 Knowledge Graph⁰. If other databases in the Fraunhofer OrientDB are published and demo username/password combinations are given, the following table will be updated.

Database	Username	Password
covid	covid_user	covid

The `covid` database can be downloaded and converted to a BEL graph like this:

```
import pybel
graph = pybel.from_fraunhofer_orientdb(
    database='covid',
    user='covid_user',
    password='covid',
)
graph.summarize()
```

However, because the source BEL scripts for the COVID-19 Knowledge Graph are available on [GitHub](#) and the authors pre-enabled it for PyBEL, it can be downloaded with `pip install git+https://github.com/covid19kg/covid19kg.git` and used with the following python code:

```
import covid19kg
graph = covid19kg.get_graph()
graph.summarize()
```

Warning: It was initially planned to handle some of the non-standard relationships listed in the Fraunhofer OrientDB’s `schema` in their OrientDB Studio instance, but none of them actually appear in the only network that is accessible. If this changes, please leave an issue at <https://github.com/pybel/pybel/issues> so it can be addressed.

Return type `BELGraph`

⁰ Domingo-Fernández, D., et al. (2020). COVID-19 Knowledge Graph: a computable, multi-modal, cause-and-effect knowledge model of COVID-19 pathophysiology. *bioRxiv* 2020.04.14.040667.

11.6 Databases

11.6.1 SQL Databases

Conversion functions for BEL graphs with a SQL database.

`pybel.from_database(name, version=None, manager=None)`

Load a BEL graph from a database.

If `name` and `version` are given, finds it exactly with `pybel.manager.Manager.get_network_by_name_version()`. If just the name is given, finds most recent with `pybel.manager.Manager.get_network_by_name()`

Parameters

- `name` (`str`) – The name of the graph
- `version` (`Optional[str]`) – The version string of the graph. If not specified, loads most recent graph added with this name

Returns A BEL graph loaded from the database

Return type `Optional[BELGraph]`

`pybel.to_database(graph, manager=None, use_tqdm=False)`

Store a graph in a database.

Parameters `graph` (`BELGraph`) – A BEL graph

Returns If successful, returns the network object from the database.

Return type `Optional[Network]`

11.6.2 Neo4j

Output functions for BEL graphs to Neo4j.

`pybel.to_neo4j(graph, neo_connection, use_tqdm=False)`

Upload a BEL graph to a Neo4j graph database using py2neo.

Parameters

- `graph` (`pybel.BELGraph`) – A BEL Graph
- `neo_connection` (`str or py2neo.Graph`) – A py2neo connection object. Refer to the [py2neo documentation](#) for how to build this object.

Example Usage:

```
>>> import py2neo
>>> import pybel
>>> from pybel.examples import sialic_acid_graph
>>> neo_graph = py2neo.Graph("http://localhost:7474/db/data/") # use your own
   ↵connection settings
>>> pybel.to_neo4j(sialic_acid_graph, neo_graph)
```

11.7 Lossy Export

11.7.1 Umbrella Node-Link JSON

The Umbrella Node-Link JSON format is similar to node-link but uses full BEL terms as nodes.

Given a BEL statement describing that X phosphorylates Y like $\text{act}(\text{p}(X)) \rightarrow \text{p}(Y, \text{pmod}(\text{Ph}))$, PyBEL usually stores the `act()` information about X as part of the relationship. In Umbrella mode, this stays as part of the node.

Note that this generates additional nodes in the network for each of the “modified” versions of the node. For example, $\text{act}(\text{p}(X))$ will be represented as individual node instead of $\text{p}(X)$, as in the standard node-link JSON exporter.

A user might want to use this exporter in the following scenarios:

- Represent transitivity in activities like $\text{in}(\text{p}(X, \text{pmod}(\text{Ph})) \rightarrow \text{act}(\text{p}(X)) \rightarrow \text{p}(Y, \text{pmod}(\text{Ph})) \rightarrow \text{act}(\text{p}(Y))$ with four nodes that are more ammenable to simulations (e.g., boolean networks, petri nets).
- Visualizing networks that in similar way to the legacy BEL [Cytoscape plugin](#) from the BEL Framework (warning: now defunct) using tools like Cytoscape.

`pybel.to_umbrella_nodelink(graph)`
Convert this graph to an umbrella node-link JSON object.

Parameters `graph` (BELGraph) – A BEL graph

Return type Mapping[str, Any]

`pybel.to_umbrella_nodelink_file(graph, path, **kwargs)`
Write this graph as an umbrella node-link JSON to a file.

Parameters

- `graph` (BELGraph) – A BEL graph
- `path` (Union[str, TextIO]) – A path or file-like

Return type None

`pybel.to_umbrella_nodelink_gz(graph, path, **kwargs)`
Write this graph as an umbrella node-link JSON to a gzipped file.

Return type None

11.7.2 GraphML

Conversion functions for BEL graphs with [GraphML](#).

`pybel.to_graphml(graph, path, schema=None)`
Write a graph to a GraphML XML file using `networkx.write_graphml()`.

Parameters

- `graph` (BELGraph) – BEL Graph
- `path` (Union[str, BinaryIO]) – Path to the new exported file
- `schema` (Optional[str]) – Type of export. Currently supported: “simple” and “umbrella”.

The .graphml file extension is suggested so Cytoscape can recognize it. By default, this function exports using the PyBEL schema of including modifier information into the edges. As an alternative, this function can also distinguish between

Return type None

11.7.3 Miscellaneous

This module contains IO functions for outputting BEL graphs to lossy formats, such as GraphML and CSV.

`pybel.to_csv(graph, path, sep=None)`

Write the graph as a tab-separated edge list.

The resulting file will contain the following columns:

1. Source BEL term
2. Relation
3. Target BEL term
4. Edge data dictionary

See the Data Models section of the documentation for which data are stored in the edge data dictionary, such as queryable information about transforms on the subject and object and their associated metadata.

Return type None

`pybel.to_sif(graph, path, sep=None)`

Write the graph as a tab-separated SIF file.

The resulting file will contain the following columns:

1. Source BEL term
2. Relation
3. Target BEL term

This format is simple and can be used readily with many applications, but is lossy in that it does not include relation metadata.

Return type None

`pybel.to_gsea(graph, path)`

Write the genes/gene products to a GRP file for use with GSEA gene set enrichment analysis.

See also:

- [GRP format specification](#)
- [GSEA publication](#)

Return type None

12.1 Manager API

The BaseManager takes care of building and maintaining the connection to the database via SQLAlchemy.

class `pybel.manager.BaseManager(engine, session)`

A wrapper around a SQLAlchemy engine and session.

Instantiate a manager from an engine and session.

base

alias of `sqlalchemy.ext.declarative.api.Base`

create_all (`checkfirst=True`)

Create the PyBEL cache's database and tables.

Parameters `checkfirst (bool)` – Check if the database exists before trying to re-make it

Return type `None`

drop_all (`checkfirst=True`)

Drop all data, tables, and databases for the PyBEL cache.

Parameters `checkfirst (bool)` – Check if the database exists before trying to drop it

Return type `None`

bind()

Bind the metadata to the engine and session.

Return type `None`

The Manager collates multiple groups of functions for interacting with the database. For sake of code clarity, they are separated across multiple classes that are documented below.

class `pybel.manager.Manager(connection=None, engine=None, session=None, **kwargs)`

Bases: `pybel.manager.cache_manager._Manager`

A manager for the PyBEL database.

Create a connection to database and a persistent session using SQLAlchemy.

A custom default can be set as an environment variable with the name `pybel.constants.PYBEL_CONNECTION`, using an [RFC-1738](#) string. For example, a MySQL string can be given with the following form:

```
mysql+pymysql://<username>:<password>@<host>/<dbname>?  
charset=utf8[&<options>]
```

A SQLite connection string can be given in the form:

```
sqlite:///~/Desktop/cache.db
```

Further options and examples can be found on the SQLAlchemy documentation on [engine configuration](#).

Parameters

- **connection** (*Optional[str]*) – An RFC-1738 database connection string. If `None`, tries to load from the environment variable `PYBEL_CONNECTION` then from the config file `~/.config/pybel/config.json` whose value for `PYBEL_CONNECTION` defaults to `pybel.constants.DEFAULT_CACHE_LOCATION`.
- **engine** – Optional engine to use. Must be specified with a session and no connection.
- **session** – Optional session to use. Must be specified with an engine and no connection.
- **echo** (*bool*) – Turn on echoing sql
- **autoflush** (*Optional[bool]*) – Defaults to `True` if not specified in kwargs or configuration.
- **autocommit** (*Optional[bool]*) – Defaults to `False` if not specified in kwargs or configuration.
- **expire_on_commit** (*Optional[bool]*) – Defaults to `False` if not specified in kwargs or configuration.
- **scopefunc** – Scoped function to pass to `sqlalchemy.orm.scoped_session()`

From the Flask-SQLAlchemy documentation:

An extra key '`scopefunc`' can be set on the options dict to specify a custom scope function. If it's not provided, Flask's app context stack identity is used. This will ensure that sessions are created and removed with the request/response cycle, and should be fine in most cases.

Allowed Usages:

Instantiation with connection string as positional argument

```
>>> my_connection = 'sqlite:///~/Desktop/cache.db'  
>>> manager = Manager(my_connection)
```

Instantiation with connection string as positional argument with keyword arguments

```
>>> my_connection = 'sqlite:///~/Desktop/cache.db'  
>>> manager = Manager(my_connection, echo=True)
```

Instantiation with connection string as keyword argument

```
>>> my_connection = 'sqlite:///~/Desktop/cache.db'  
>>> manager = Manager(connection=my_connection)
```

Instantiation with connection string as keyword argument with keyword arguments

```
>>> my_connection = 'sqlite:///~/Desktop/cache.db'  
>>> manager = Manager(connection=my_connection, echo=True)
```

Instantiation with user-supplied engine and session objects as keyword arguments

```
>>> my_engine, my_session = ... # magical creation! See SQLAlchemy documentation  
>>> manager = Manager(engine=my_engine, session=my_session)
```

12.2 Manager Components

```
class pybel.manager.NetworkManager(engine, session)
    Groups functions for inserting and querying networks in the database's network store.

    Instantiate a manager from an engine and session.

count_networks()
    Count the networks in the database.

    Return type int

list_networks()
    List all networks in the database.

    Return type List[Network]

list_recent_networks()
    List the most recently created version of each network (by name).

    Return type List[Network]

has_name_version(name, version)
    Check if there exists a network with the name/version combination in the database.

    Return type bool

drop_networks()
    Drop all networks.

    Return type None

drop_network_by_id(network_id)
    Drop a network by its database identifier.

    Return type None

drop_network(network)
    Drop a network, while also cleaning up any edges that are no longer part of any network.

    Return type None

query_singleton_edges_from_network(network)
    Return a query selecting all edge ids that only belong to the given network.

    Return type Query

get_network_versions(name)
    Return all of the versions of a network with the given name.

    Return type Set[str]

get_network_by_name_version(name, version)
    Load the network with the given name and version if it exists.

    Return type Optional[Network]

get_graph_by_name_version(name, version)
    Load the BEL graph with the given name, or allows for specification of version.

    Return type Optional[BELGraph]

get_networks_by_name(name)
    Get all networks with the given name. Useful for getting all versions of a given network.

    Return type List[Network]
```

```
get_most_recent_network_by_name(name)
    Get the most recently created network with the given name.

    Return type Optional[Network]

get_graph_by_most_recent(name)
    Get the most recently created network with the given name as a pybel.BELGraph.

    Return type Optional[BELGraph]

get_network_by_id(network_id)
    Get a network from the database by its identifier.

    Return type Network

get_graph_by_id(network_id)
    Get a network from the database by its identifier and converts it to a BEL graph.

    Return type BELGraph

get_networks_by_ids(network_ids)
    Get a list of networks with the given identifiers.

    Note: order is not necessarily preserved.

    Return type List[Network]

get_graphs_by_ids(network_ids)
    Get a list of networks with the given identifiers and converts to BEL graphs.

    Return type List[BELGraph]

get_graph_by_ids(network_ids)
    Get a combine BEL Graph from a list of network identifiers.

    Return type BELGraph

class pybel.manager.QueryManager(engine, session)
    An extension to the Manager to make queries over the database.

    Instantiate a manager from an engine and session.

    count_nodes()
        Count the number of nodes in the database.

        Return type int

    query_nodes(bel=None, type=None, namespace=None, name=None)
        Query nodes in the database.

        Parameters
            • bel (Optional[str]) – BEL term that describes the biological entity. e.g. p(HGNC:APP)
            • type (Optional[str]) – Type of the biological entity. e.g. Protein
            • namespace (Optional[str]) – Namespace keyword that is used in BEL. e.g. HGNC
            • name (Optional[str]) – Name of the biological entity. e.g. APP

        Return type List[Node]

    count_edges()
        Count the number of edges in the database.

        Return type int
```

get_edges_with_citation(*citation*)

Get the edges with the given citation.

Return type `List[Edge]`

get_edges_with_citations(*citations*)

Get edges with one of the given citations.

Return type `List[Edge]`

search_edges_with_evidence(*evidence*)

Search edges with the given evidence.

Parameters `evidence` (`str`) – A string to search evidences. Can use wildcard percent symbol (%).

Return type `List[Edge]`

search_edges_with_bel(*bel*)

Search edges with given BEL.

Parameters `bel` (`str`) – A BEL string to use as a search

Return type `List[Edge]`

get_edges_with_annotation(*annotation*, *value*)

Search edges with the given annotation/value pair.

Return type `List[Edge]`

query_edges(*bel=None*, *source_function=None*, *source=None*, *target_function=None*, *target=None*, *relation=None*)

Return a query over the edges in the database.

Usually this means that you should call `list()` or `.all()` on this result.

Parameters

- `bel` (`Optional[str]`) – BEL statement that represents the desired edge.
- `source_function` (`Optional[str]`) – Filter source nodes with the given BEL function
- `source` (`Union[None, str, Node]`) – BEL term of source node e.g. `p (HGNC:APP)` or `Node` object.
- `target_function` (`Optional[str]`) – Filter target nodes with the given BEL function
- `target` (`Union[None, str, Node]`) – BEL term of target node e.g. `p (HGNC:APP)` or `Node` object.
- `relation` (`Optional[str]`) – The relation that should be present between source and target node.

query_citations(*db=None*, *db_id=None*, *name=None*, *author=None*, *date=None*, *evidence_text=None*)

Query citations in the database.

Parameters

- `db` (`Optional[str]`) – Type of the citation. e.g. PubMed
- `db_id` (`Optional[str]`) – The identifier used for the citation. e.g. PubMed_ID
- `name` (`Optional[str]`) – Title of the citation.

- **author** (`Union[None, str, List[str]]`) – The name or a list of names of authors participated in the citation.
- **date** (`Union[None, str, date]`) – Publishing date of the citation.
- **evidence_text** (`Optional[str]`) –

Return type `List[Citation]`

query_edges_by_pubmed_identifiers (`pubmed_identifiers`)

Get all edges annotated to the documents identified by the given PubMed identifiers.

Return type `List[Edge]`

query_induction (`nodes`)

Get all edges between any of the given nodes (minimum length of 2).

Return type `List[Edge]`

query_neighbors (`nodes`)

Get all edges incident to any of the given nodes.

Return type `List[Edge]`

CHAPTER
THIRTEEN

MODELS

This module contains the SQLAlchemy database models that support the definition cache and graph cache.

class `pybel.manager.models.Base` (`**kwargs`)

The most base type

A simple constructor that allows initialization from kwargs.

Sets attributes on the constructed instance using the names and values in kwargs.

Only keys that are present as attributes of the instance's class are allowed. These could be, for example, any mapped columns or relationships.

class `pybel.manager.models.Namespace` (`**kwargs`)

Represents a BEL Namespace.

A simple constructor that allows initialization from kwargs.

Sets attributes on the constructed instance using the names and values in kwargs.

Only keys that are present as attributes of the instance's class are allowed. These could be, for example, any mapped columns or relationships.

uploaded

The date of upload

keyword

Keyword that is used in a BEL file to identify a specific namespace

pattern

Contains regex pattern for value identification.

miriam_id

MIRIAM resource identifier matching the regular expression ^MIR:001\d{5}\$

version

Version of the namespace

url

BELNS Resource location as URL

name

Name of the given namespace

domain

Domain for which this namespace is valid

species

Taxonomy identifiers for which this namespace is valid

description

Optional short description of the namespace

created

DateTime of the creation of the namespace definition file

query_url

URL that can be used to query the namespace (externally from PyBEL)

author

The author of the namespace

license

License information

contact

Contact information

get_term_to_encodings()

Return the term (db, id, name) to encodings from this namespace.

Return type `Mapping[Tuple[Optional[str], str], str]`

to_json (*include_id=False*)

Return the most useful entries as a dictionary.

Parameters `include_id (bool)` – If true, includes the model identifier

Return type `Mapping[str, str]`

class `pybel.manager.models.NamespaceEntry (**kwargs)`

Represents a name within a BEL namespace.

A simple constructor that allows initialization from kwargs.

Sets attributes on the constructed instance using the names and values in kwargs.

Only keys that are present as attributes of the instance's class are allowed. These could be, for example, any mapped columns or relationships.

name

Name that is defined in the corresponding namespace definition file

identifier

The database accession number

encoding

The biological entity types for which this name is valid

to_json (*include_id=False*)

Describe the namespaceEntry as dictionary of Namespace-Keyword and Name.

Parameters `include_id (bool)` – If true, includes the model identifier

Return type `Mapping[str, str]`

classmethod `name_contains (name_query)`

Make a filter if the name contains a certain substring.

class `pybel.manager.models.Network (**kwargs)`

Represents a collection of edges, specified by a BEL Script.

A simple constructor that allows initialization from kwargs.

Sets attributes on the constructed instance using the names and values in kwargs.

Only keys that are present as attributes of the instance's class are allowed. These could be, for example, any mapped columns or relationships.

name

Name of the given Network (from the BEL file)

version

Release version of the given Network (from the BEL file)

authors

Authors of the underlying BEL file

contact

Contact email from the underlying BEL file

description

Descriptive text from the underlying BEL file

copyright

Copyright information

disclaimer

Disclaimer information

licenses

License information

blob

A pickled version of this network

to_json (include_id=False)

Return this network as JSON.

Parameters `include_id (bool)` – If true, includes the model identifier

Return type `Mapping[str, Any]`

classmethod name_contains (name_query)

Build a filter for networks whose names contain the query.

classmethod description_contains (description_query)

Build a filter for networks whose descriptions contain the query.

classmethod id_in (network_ids)

Build a filter for networks whose identifiers appear in the given sequence.

as_bel ()

Get this network and loads it into a BELGraph.

Return type `pybel.BELGraph`

store_bel (graph)

Insert a BEL graph.

Parameters `graph (pybel.BELGraph)` – A BEL Graph

class pybel.manager.models.Modification (kwargs)**

The modifications that are present in the network are stored in this table.

A simple constructor that allows initialization from kwargs.

Sets attributes on the constructed instance using the names and values in kwargs.

Only keys that are present as attributes of the instance's class are allowed. These could be, for example, any mapped columns or relationships.

type
Type of the stored modification e.g. Fusion, gmod, pmod, etc

variantString
HGVS string if sequence modification

residue
Three letter amino acid code if PMOD

position
Position of PMOD or GMOD

to_json()
Recreate a is_variant dictionary for BELGraph.

Returns Dictionary that describes a variant or a fusion.

Return type *Variant* or *FusionBase*

class `pybel.manager.models.Node(**kwargs)`
Represents a BEL Term.

A simple constructor that allows initialization from kwargs.

Sets attributes on the constructed instance using the names and values in kwargs.

Only keys that are present as attributes of the instance's class are allowed. These could be, for example, any mapped columns or relationships.

type
The type of the represented biological entity e.g. Protein or Gene

is_variant
Identifies whether or not the given node is a variant

has_fusion
Identifies whether or not the given node is a fusion

bel
Canonical BEL term that represents the given node

data
PyBEL BaseEntity as JSON

classmethod bel_contains(bel_query)
Build a filter for nodes whose BEL contain the query.

as_bel()
Serialize this node as a PyBEL DSL object.

Return type *pybel.dsl.BaseEntity*

to_json()
Serialize this node as a JSON object using as_bel().

class `pybel.manager.models.Author(**kwargs)`
Contains all author names.

A simple constructor that allows initialization from kwargs.

Sets attributes on the constructed instance using the names and values in kwargs.

Only keys that are present as attributes of the instance's class are allowed. These could be, for example, any mapped columns or relationships.

```
classmethod name_contains(name_query)
    Build a filter for authors whose names contain the given query.

classmethod has_name_in(names)
    Build a filter if the author has any of the given names.

class pybel.manager.models.Citation(**kwargs)
    The information about the citations that are used to prove a specific relation are stored in this table.

    A simple constructor that allows initialization from kwargs.

    Sets attributes on the constructed instance using the names and values in kwargs.

    Only keys that are present as attributes of the instance's class are allowed. These could be, for example, any mapped columns or relationships.

    db
        Type of the stored publication e.g. PubMed

    db_id
        Reference identifier of the publication e.g. PubMed_ID

    title
        Title of the publication

    journal
        Journal name

    volume
        Volume of the journal

    issue
        Issue within the volume

    pages
        Pages of the publication

    date
        Publication date

    first_id
        First author

    last_id
        Last author

    property is_pubmed
        Return if this is a PubMed citation.

        Return type bool

    property is_enriched
        Return if this citation has been enriched for name, title, and other metadata.

        Return type bool

    to_json(include_id=False)
        Create a citation dictionary that is used to recreate the edge data dictionary of a BELGraph.

        Parameters include_id(bool) – If true, includes the model identifier

        Return type Mapping[str, Any]

        Returns Citation dictionary for the recreation of a BELGraph.
```

class `pybel.manager.models.Evidence(**kwargs)`

This table contains the evidence text that proves a specific relationship and refers the source that is cited.

A simple constructor that allows initialization from kwargs.

Sets attributes on the constructed instance using the names and values in kwargs.

Only keys that are present as attributes of the instance's class are allowed. These could be, for example, any mapped columns or relationships.

text

Supporting text from a given publication

to_json(include_id=False)

Create a dictionary that is used to recreate the edge data dictionary for a BELGraph.

Parameters `include_id(bool)` – If true, includes the model identifier

Returns Dictionary containing citation and evidence for a BELGraph edge.

Return type `dict`

class `pybel.manager.models.Property(**kwargs)`

The property table contains additional information that is used to describe the context of a relation.

A simple constructor that allows initialization from kwargs.

Sets attributes on the constructed instance using the names and values in kwargs.

Only keys that are present as attributes of the instance's class are allowed. These could be, for example, any mapped columns or relationships.

is_subject

Identifies which participant of the edge is affected by the given property

modifier

The modifier: one of activity, degradation, location, or translocation

relative_key

Relative key of effect e.g. to_tloc or from_tloc

property side

Return either `pybel.constants.SUBJECT` or `pybel.constants.OBJECT`.

Return type `str`

to_json()

Create a property dict that is used to recreate an edge dictionary for a BELGraph.

Returns Property dictionary of an edge that is participant (sub/obj) related.

Return type `dict`

class `pybel.manager.models.Edge(**kwargs)`

Relationships between BEL nodes and their properties, annotations, and provenance.

A simple constructor that allows initialization from kwargs.

Sets attributes on the constructed instance using the names and values in kwargs.

Only keys that are present as attributes of the instance's class are allowed. These could be, for example, any mapped columns or relationships.

bel

Valid BEL statement that represents the given edge

md5

The hash of the source, target, and associated metadata

data

The stringified JSON representing this edge

get_annotations_json()

Format the annotations properly.

Return type Optional[`dict[str, dict[str, bool]]`]

to_json(*include_id=False*)

Create a dictionary of one BEL Edge that can be used to create an edge in a BELGraph.

Parameters `include_id` (`bool`) – Include the database identifier?

Return type `Mapping[str, Any]`

Returns Dictionary that contains information about an edge of a BELGraph. Including participants and edge data information.

insert_into_graph(*graph*)

Insert this edge into a BEL graph.

Parameters `graph` (`pybel.BELGraph`) – A BEL graph

CHAPTER
FOURTEEN

COOKBOOK

An extensive set of examples can be found on the [PyBEL Notebooks](#) repository on GitHub. These notebooks contain basic usage and also make numerous references to the analytical package [PyBEL Tools](#)

14.1 Configuration

The default connection string can be set as an environment variable in your `~/.bashrc`. If you're using MySQL or MariaDB, it could look like this:

```
$ export PYBEL_CONNECTION="mysql+pymysql://user:password@server_name/database_name?  
↪charset=utf8"
```

14.1.1 Prepare a Cytoscape Network

Load, compile, and export to Cytoscape format:

```
$ pybel convert --path ~/Desktop/example.bel --graphml ~/Desktop/example.graphml
```

In Cytoscape, open with Import > Network > From File.

COMMAND LINE INTERFACE

Note: The command line wrapper might not work on Windows. Use `python3 -m pybel` if it has issues.

PyBEL automatically installs the command `pybel`. This command can be used to easily compile BEL documents and convert to other formats. See `pybel --help` for usage details. This command makes logs of all conversions and warnings to the directory `~/ .pybel/`.

15.1 `pybel`

PyBEL CLI on `/home/docs/checkouts/readthedocs.org/user_builds/pybel/envs/v0.14.9/bin/python`

```
pybel [OPTIONS] COMMAND [ARGS] ...
```

Options

`--version`

Show the version and exit.

`-c, --connection <connection>`

Database connection string. [default: `sqlite:///home/docs/.pybel/pybel_0.14.0_cache.db`]

15.1.1 `compile`

Compile a BEL script to a graph.

```
pybel compile [OPTIONS] PATH
```

Options

`--allow-naked-names`

Enable lenient parsing for naked names

`--disallow-nested`

Disable lenient parsing for nested statements

`--disallow-unqualified-translocations`

Disallow unqualified translocations

```
--no-identifier-validation
    Turn off identifier validation

--no-citation-clearing
    Turn off citation clearing

-r, --required-annotations <required_annotations>
    Specify multiple required annotations

--upgrade-urls

--skip-tqdm

-v, --verbose
```

Arguments

PATH

Required argument

15.1.2 insert

Insert a graph to the database.

```
pybel insert [OPTIONS] path
```

Arguments

path

Required argument

15.1.3 machine

Get content from the INDRA machine and upload to BEL Commons.

```
pybel machine [OPTIONS] [AGENTS] ...
```

Options

```
--local
    Upload to local database.

--host <host>
    URL of BEL Commons. Defaults to https://bel-commons.scai.fraunhofer.de
```

Arguments

AGENTS

Optional argument(s)

15.1.4 manage

Manage the database.

```
pybel manage [OPTIONS] COMMAND [ARGS] ...
```

drop

Drop the database.

```
pybel manage drop [OPTIONS]
```

Options

--yes

Confirm the action without prompting.

edges

Manage edges.

```
pybel manage edges [OPTIONS] COMMAND [ARGS] ...
```

ls

List edges.

```
pybel manage edges ls [OPTIONS]
```

Options

--offset <offset>

--limit <limit>

examples

Load examples to the database.

```
pybel manage examples [OPTIONS]
```

Options

-v, --debug

namespaces

Manage namespaces.

```
pybel manage namespaces [OPTIONS] COMMAND [ARGS] ...
```

drop

Drop a namespace by URL.

```
pybel manage namespaces drop [OPTIONS] URL
```

Arguments

URL

Required argument

insert

Add a namespace by URL.

```
pybel manage namespaces insert [OPTIONS] URL
```

Arguments

URL

Required argument

ls

List cached namespaces.

```
pybel manage namespaces ls [OPTIONS]
```

Options

- u, --url <url>**
Specific resource URL to list
- i, --namespace-id <namespace_id>**
Specific resource URL to list

networks

Manage networks.

```
pybel manage networks [OPTIONS] COMMAND [ARGS] ...
```

drop

Drop a network by its identifier or drop all networks.

```
pybel manage networks drop [OPTIONS]
```

Options

- n, --network-id <network_id>**
Identifier of network to drop
- y, --yes**
Drop all networks without confirmation if no identifier is given

ls

List network names, versions, and optionally, descriptions.

```
pybel manage networks ls [OPTIONS]
```

nodes

Manage nodes.

```
pybel manage nodes [OPTIONS] COMMAND [ARGS] ...
```

prune

Prune nodes not belonging to any edges.

```
pybel manage nodes prune [OPTIONS]
```

summarize

Summarize the contents of the database.

```
pybel manage summarize [OPTIONS]
```

15.1.5 neo

Upload to neo4j.

```
pybel neo [OPTIONS] path
```

Options

--connection <connection>
Connection string for neo4j upload.

--password <password>

Arguments

path
Required argument

15.1.6 post

Upload a graph to BEL Commons.

```
pybel post [OPTIONS] path
```

Options

--host <host>
URL of BEL Commons. Defaults to <https://bel-commons.scai.fraunhofer.de>

Arguments

path
Required argument

15.1.7 serialize

Serialize a graph to various formats.

```
pybel serialize [OPTIONS] path
```

Options

--tsv <tsv>
Path to output a TSV file.

--edgelist <edgelist>
Path to output a edgelist file.

--sif <sif>
Path to output an SIF file.

--gsea <gsea>
Path to output a GRP file for gene set enrichment analysis.

--graphml <graphml>
Path to output a GraphML file. Use .graphml for Cytoscape.

--nodelink <nodelink>
Path to output a node-link JSON file.

--bel <bel>
Output canonical BEL.

Arguments

path
Required argument

15.1.8 summarize

Summarize a graph.

```
pybel summarize [OPTIONS] path
```

Arguments

path
Required argument

15.1.9 warnings

List warnings from a graph.

```
pybel warnings [OPTIONS] path
```

Arguments

path

Required argument

CONSTANTS

Constants for PyBEL.

This module maintains the strings used throughout the PyBEL codebase to promote consistency.

`pybel.constants.get_cache_connection()`

Get the preferred RFC-1738 database connection string.

1. Check the environment variable PYBEL_CONNECTION
2. Check the PYBEL_CONNECTION key in the config file `~/.config/pybel/config.json`. Optionally, this config file might be in a different place if the environment variable PYBEL_CONFIG_DIRECTORY has been set.
3. Return a default connection string using a SQLite database in the `~/.pybel`. Optionally, this directory might be in a different place if the environment variable PYBEL_RESOURCE_DIRECTORY has been set.

Return type `str`

`pybel.constants.BEL_DEFAULT_NAMESPACE = 'bel'`

The default namespace given to entities in the BEL language

`pybel.constants.CITATION_TYPES = {'Book': None, 'DOI': 'doi', 'Journal': None, 'Online Re`

The valid citation types .. seealso:: <https://wiki.openbel.org/display/BELNA/Citation>

`pybel.constants.NAMESPACE_DOMAIN_TYPES = {'BiologicalProcess', 'Chemical', 'Gene and Gene P`

The valid namespace types .. seealso:: <https://wiki.openbel.org/display/BELNA/Custom+Namespaces>

`pybel.constants.CITATION_DB = 'db'`

Represents the key for the citation type in a citation dictionary

`pybel.constants.CITATION_IDENTIFIER = 'db_id'`

Represents the key for the citation reference in a citation dictionary

`pybel.constants.CITATION_DB_NAME = 'db_name'`

Represents the key for the optional PyBEL citation title entry in a citation dictionary

`pybel.constants.CITATION_DATE = 'date'`

Represents the key for the citation date in a citation dictionary

`pybel.constants.CITATION_AUTHORS = 'authors'`

Represents the key for the citation authors in a citation dictionary

`pybel.constants.CITATION_JOURNAL = 'db_name'`

Represents the key for the citation comment in a citation dictionary

`pybel.constants.CITATION_VOLUME = 'volume'`

Represents the key for the optional PyBEL citation volume entry in a citation dictionary

```
pybel.constants.CITATION_ISSUE = 'issue'
    Represents the key for the optional PyBEL citation issue entry in a citation dictionary

pybel.constants.CITATION_PAGES = 'pages'
    Represents the key for the optional PyBEL citation pages entry in a citation dictionary

pybel.constants.CITATION_FIRST_AUTHOR = 'first'
    Represents the key for the optional PyBEL citation first author entry in a citation dictionary

pybel.constants.CITATION_LAST_AUTHOR = 'last'
    Represents the key for the optional PyBEL citation last author entry in a citation dictionary

pybel.constants.FUNCTION = 'function'
    The node data key specifying the node's function (e.g. GENE, MIRNA, BIOPROCESS, etc.)

pybel.constants.CONCEPT = 'concept'
    The key specifying a concept

pybel.constants.NAMESPACE = 'namespace'
    The key specifying an identifier dictionary's namespace. Used for nodes, activities, and transformations.

pybel.constants.NAME = 'name'
    The key specifying an identifier dictionary's name. Used for nodes, activities, and transformations.

pybel.constants.IDENTIFIER = 'identifier'
    The key specifying an identifier dictionary

pybel.constants.LABEL = 'label'
    The key specifying an optional label for the node

pybel.constants.DESCRIPTION = 'description'
    The key specifying an optional description for the node

pybel.constants.XREFS = 'xref'
    The key specifying xrefs

pybel.constants.MEMBERS = 'members'
    They key representing the nodes that are a member of a composite or complex

pybel.constants.REACTANTS = 'reactants'
    The key representing the nodes appearing in the reactant side of a biochemical reaction

pybel.constants.PRODUCTS = 'products'
    The key representing the nodes appearing in the product side of a biochemical reaction

pybel.constants.PARTNER_3P = 'partner_3p'
    The key specifying the identifier dictionary of the fusion's 3-Prime partner

pybel.constants.PARTNER_5P = 'partner_5p'
    The key specifying the identifier dictionary of the fusion's 5-Prime partner

pybel.constants.RANGE_3P = 'range_3p'
    The key specifying the range dictionary of the fusion's 3-Prime partner

pybel.constants.RANGE_5P = 'range_5p'
    The key specifying the range dictionary of the fusion's 5-Prime partner

pybel.constants.VARIANTS = 'variants'
    The key specifying the node has a list of associated variants

pybel.constants.KIND = 'kind'
    The key representing what kind of variation is being represented
```

```
pybel.constants.HGVS = 'hgvs'
    The value for KIND for an HGVS variant

pybel.constants.PMOD = 'pmod'
    The value for KIND for a protein modification

pybel.constants.GMOD = 'gmod'
    The value for KIND for a gene modification

pybel.constants.FRAGMENT = 'frag'
    The value for KIND for a fragment

pybel.constants.PYBEL_VARIANT_KINDS = {'frag', 'gmod', 'hgvs', 'pmod'}
    The allowed values for KIND

pybel.constants.PYBEL_NODE_DATA_KEYS = {'function', 'fusion', 'identifier', 'members', 'name'}
    The group of all BEL-provided keys for node data dictionaries, used for hashing.

pybel.constants.DIRTY = 'dirty'
    Used as a namespace when none is given when lenient parsing mode is turned on. Not recommended!

pybel.constants.ABUNDANCE = 'Abundance'
    Represents the BEL abundance, abundance()

pybel.constants.GENE = 'Gene'
    Represents the BEL abundance, geneAbundance() .. seealso:: http://openbel.org/language/version\_2.0/bel\_specification\_version\_2.0.html#Xabundancea

pybel.constants.RNA = 'RNA'
    Represents the BEL abundance, rnaAbundance()

pybel.constants.MIRNA = 'miRNA'
    Represents the BEL abundance, microRNAAbundance()

pybel.constants.PROTEIN = 'Protein'
    Represents the BEL abundance, proteinAbundance()

pybel.constants.BIOPROCESS = 'BiologicalProcess'
    Represents the BEL function, biologicalProcess()

pybel.constants.PATHOLOGY = 'Pathology'
    Represents the BEL function, pathology()

pybel.constants.POPULATION = 'Population'
    Represents the BEL function, populationAbundance()

pybel.constants.COMPOSITE = 'Composite'
    Represents the BEL abundance, compositeAbundance()

pybel.constants.COMPLEX = 'Complex'
    Represents the BEL abundance, complexAbundance()

pybel.constants.REACTION = 'Reaction'
    Represents the BEL transformation, reaction()

pybel.constants.PYBEL_NODE_FUNCTIONS = {'Abundance', 'BiologicalProcess', 'Complex', 'Compound', 'Disease', 'Entity', 'Function', 'Gene', 'Location', 'Pathway', 'Protein', 'Reaction', 'Region', 'Species', 'Strain', 'Substance', 'Variant'}
    A set of all of the valid PyBEL node functions

pybel.constants.rev_abundance_labels = {'Abundance': 'a', 'BiologicalProcess': 'bp', 'Complex': 'c', 'Compound': 'comp', 'Disease': 'd', 'Entity': 'e', 'Function': 'f', 'Gene': 'g', 'Location': 'l', 'Pathway': 'p', 'Protein': 'protein', 'Reaction': 'r', 'Region': 'region', 'Species': 's', 'Strain': 'strain', 'Substance': 'substance', 'Variant': 'v'}
    The mapping from PyBEL node functions to BEL strings

pybel.constants.RELATION = 'relation'
    The key for an internal edge data dictionary for the relation string
```

```
pybel.constants.CITATION = 'citation'
    The key for an internal edge data dictionary for the citation dictionary

pybel.constants.EVIDENCE = 'evidence'
    The key for an internal edge data dictionary for the evidence string

pybel.constants.ANNOTATIONS = 'annotations'
    The key for an internal edge data dictionary for the annotations dictionary

pybel.constants.SUBJECT = 'subject'
    The key for an internal edge data dictionary for the subject modifier dictionary

pybel.constants.OBJECT = 'object'
    The key for an internal edge data dictionary for the object modifier dictionary

pybel.constants.LINE = 'line'
    The key or an internal edge data dictionary for the line number

pybel.constants.HASH = 'hash'
    The key representing the hash of the other

pybel.constants.PYBEL_EDGE_DATA_KEYS = {'annotations', 'citation', 'evidence', 'object', 'subject'}
    The group of all BEL-provided keys for edge data dictionaries, used for hashing.

pybel.constants.PYBEL_EDGE_METADATA_KEYS = {'hash', 'line'}
    The group of all PyBEL-specific keys for edge data dictionaries, not used for hashing.

pybel.constants.PYBEL_EDGE_ALL_KEYS = {'annotations', 'citation', 'evidence', 'hash', 'line', 'subject', 'object'}
    The group of all PyBEL annotated keys for edge data dictionaries

pybel.constants.HAS_REACTANT = 'hasReactant'
    A BEL relationship

pybel.constants.HAS_PRODUCT = 'hasProduct'
    A BEL relationship

pybel.constants.HAS_VARIANT = 'hasVariant'
    A BEL relationship

pybel.constants.TRANSCRIBED_TO = 'transcribedTo'
    A BEL relationship GENE to RNA is called transcription

pybel.constants.TRANSLATED_TO = 'translatedTo'
    A BEL relationship RNA to PROTEIN is called translation

pybel.constants.INCREASES = 'increases'
    A BEL relationship

pybel.constants.DIRECTLY_INCREASES = 'directlyIncreases'
    A BEL relationship

pybel.constants.DECREASES = 'decreases'
    A BEL relationship

pybel.constants.DIRECTLY_DECREASES = 'directlyDecreases'
    A BEL relationship

pybel.constants.CAUSES_NO_CHANGE = 'causesNoChange'
    A BEL relationship

pybel.constants.REGULATES = 'regulates'
    A BEL relationship
```

```
pybel.constants.BINDS = 'binds'
    A BEL relationship

pybel.constants.CORRELATION = 'correlation'
    A BEL relationship

pybel.constants.NO_CORRELATION = 'noCorrelation'
    A BEL relationship

pybel.constants.NEGATIVE_CORRELATION = 'negativeCorrelation'
    A BEL relationship

pybel.constants.POSITIVE_CORRELATION = 'positiveCorrelation'
    A BEL relationship

pybel.constants.ASSOCIATION = 'association'
    A BEL relationship

pybel.constants.ORTHOLOGOUS = 'orthologous'
    A BEL relationship

pybel.constants.ANALOGOUS_TO = 'analogousTo'
    A BEL relationship

pybel.constants.IS_A = 'isA'
    A BEL relationship

pybel.constants.RATE_LIMITING_STEP_OF = 'rateLimitingStepOf'
    A BEL relationship

pybel.constants.SUBPROCESS_OF = 'subProcessOf'
    A BEL relationship

pybel.constants.BIOMARKER_FOR = 'biomarkerFor'
    A BEL relationship

pybel.constants.PROGNOSTIC BIOMARKER_FOR = 'prognosticBiomarkerFor'
    A BEL relationship

pybel.constants.EQUIVALENT_TO = 'equivalentTo'
    A BEL relationship, added by PyBEL

pybel.constants.PART_OF = 'partOf'
    A BEL relationship, added by PyBEL

pybel.constants.CAUSAL_INCREASE_RELATIONS = {'directlyIncreases', 'increases'}
    A set of all causal relationships that have an increasing effect

pybel.constants.CAUSAL_DECREASE_RELATIONS = {'decreases', 'directlyDecreases'}
    A set of all causal relationships that have a decreasing effect

pybel.constants.DIRECT_CAUSAL_RELATIONS = {'directlyDecreases', 'directlyIncreases'}
    A set of direct causal relations

pybel.constants.INDIRECT_CAUSAL_RELATIONS = {'decreases', 'increases', 'regulates'}
    A set of direct causal relations

pybel.constants.CAUSAL_POLAR_RELATIONS = {'decreases', 'directlyDecreases', 'directlyIncreases'}
    A set of causal relationships that are polar

pybel.constants.CAUSAL_RELATIONS = {'decreases', 'directlyDecreases', 'directlyIncreases', 'increases'}
    A set of all causal relationships
```

```
pybel.constants.CORRELATIVE_RELATIONS = {'correlation', 'negativeCorrelation', 'noCorrelation'}
A set of all correlative relationships

pybel.constants.POLAR_RELATIONS = {'decreases', 'directlyDecreases', 'directlyIncreases', 'increases', 'directlyIncreases'}
A set of polar relations

pybel.constants.TWO_WAY_RELATIONS = {'analogousTo', 'association', 'binds', 'correlation', 'equivalentTo', 'hasProduct', 'hasReactant', 'hasVariant'}
A set of all relationships that are inherently directionless, and are therefore added to the graph twice

pybel.constants.UNQUALIFIED_EDGES = {'equivalentTo', 'hasProduct', 'hasReactant', 'hasVariant'}
A list of relationship types that don't require annotations or evidence

pybel.constants.GRAPH_METADATA = 'document_metadata'
The key for the document metadata dictionary. Can be accessed by graph.graph[GRAPH_METADATA], or by using the property built in to the pybel.BELGraph, pybel.BELGraph.document()

pybel.constants.METADATA_NAME = 'name'
The key for the document name. Can be accessed by graph.document[METADATA_NAME] or by using the property built into the pybel.BELGraph class, pybel.BELGraph.name()

pybel.constants.METADATA_VERSION = 'version'
The key for the document version. Can be accessed by graph.document[METADATA_VERSION]

pybel.constants.METADATA_DESCRIPTION = 'description'
The key for the document description. Can be accessed by graph.document[METADATA_DESCRIPTION]

pybel.constants.METADATA_AUTHORS = 'authors'
The key for the document authors. Can be accessed by graph.document[METADATA_NAME]

pybel.constants.METADATA_CONTACT = 'contact'
The key for the document contact email. Can be accessed by graph.document[METADATA_CONTACT]

pybel.constants.METADATA_LICENSES = 'licenses'
The key for the document licenses. Can be accessed by graph.document[METADATA_LICENSES]

pybel.constants.METADATA_COPYRIGHT = 'copyright'
The key for the document copyright information. Can be accessed by graph.document[METADATA_COPYRIGHT]

pybel.constants.METADATA_DISCLAIMER = 'disclaimer'
The key for the document disclaimer. Can be accessed by graph.document[METADATA_DISCLAIMER]

pybel.constants.METADATA_PROJECT = 'project'
The key for the document project. Can be accessed by graph.document[METADATA_PROJECT]

pybel.constants.DOCUMENT_KEYS = {'Authors': 'authors', 'ContactInfo': 'contact', 'Copyright': 'copyright'}
Provides a mapping from BEL language keywords to internal PyBEL strings

pybel.constants.METADATA_INSERT_KEYS = {'authors', 'contact', 'copyright', 'description'}
The keys to use when inserting a graph to the cache

pybel.constants.INVERSE_DOCUMENT_KEYS = {'authors': 'Authors', 'contact': 'ContactInfo'}
Provides a mapping from internal PyBEL strings to BEL language keywords. Is the inverse of DOCUMENT_KEYS

pybel.constants.REQUIRED_METADATA = {'authors', 'contact', 'description', 'name', 'version'}
A set representing the required metadata during BEL document parsing

pybel.constants.FRAGMENT_START = 'start'
The key for the starting position of a fragment range
```

```

pybel.constants.FRAGMENT_STOP = 'stop'
    The key for the stopping position of a fragment range

pybel.constants.FRAGMENT_MISSING = 'missing'
    The key signifying that there is neither a start nor stop position defined

pybel.constants.FRAGMENT_DESCRIPTION = 'description'
    The key for any additional descriptive data about a fragment

pybel.constants.GMOD_ORDER = ['kind', 'identifier']
    The order for serializing gene modification data

pybel.constants.GSUB_REFERENCE = 'reference'
    The key for the reference nucleotide in a gene substitution. Only used during parsing since this is converted to HGVS.

pybel.constants.GSUB_POSITION = 'position'
    The key for the position of a gene substitution. Only used during parsing since this is converted to HGVS

pybel.constants.GSUB_VARIANT = 'variant'
    The key for the effect of a gene substitution. Only used during parsing since this is converted to HGVS

pybel.constants.PMOD_CODE = 'code'
    The key for the protein modification code.

pybel.constants.PMOD_POSITION = 'pos'
    The key for the protein modification position.

pybel.constants.PMOD_ORDER = ['kind', 'identifier', 'code', 'pos']
    The order for serializing information about a protein modification

pybel.constants.PSUB_REFERENCE = 'reference'
    The key for the reference amino acid in a protein substitution. Only used during parsing since this is converted to HGVS

pybel.constants.PSUB_POSITION = 'position'
    The key for the position of a protein substitution. Only used during parsing since this is converted to HGVS.

pybel.constants.PSUB_VARIANT = 'variant'
    The key for the variant of a protein substitution. Only used during parsing since this is converted to HGVS.

pybel.constants.TRUNCATION_POSITION = 'position'
    The key for the position at which a protein is truncated

pybel.constants.belns_encodings = {'A': {'Abundance', 'Complex', 'Gene', 'Protein', 'RNA'},
    The mapping from BEL namespace codes to PyBEL internal abundance constants ..seealso:: https://wiki.openbel.org/display/BELNA/Assignment+of+Encoding+%28Allowed+Functions%29+for+BEL+Namespaces

pybel.constants.DEFAULT_SERVICE_URL = 'https://bel-commons.scai.fraunhofer.de'
    The default location of PyBEL Web

```

Language constants for BEL.

This module contains mappings between PyBEL's internal constants and BEL language keywords.

```
class pybel.language.Entity(*, namespace, name=None, identifier=None)
    Represents a named entity with a namespace and name/identifier.
```

Create a dictionary representing a reference to an entity.

Parameters

- **namespace** (`str`) – The namespace to which the entity belongs
- **name** (`Optional[str]`) – The name of the entity

- **identifier** (Optional[str]) – The identifier of the entity in the namespace

property namespace

The entity's namespace.

Return type str

property name

The entity's name or label.

Return type str

property identifier

The entity's identifier.

Return type str

property curie

Return this entity as a CURIE.

Return type str

property obo

Return this entity as an OBO-style CURIE.

Return type str

```
pybel.language.activity_labels = {'cat': 'cat', 'catalyticActivity': 'cat', 'chap': 'ch'
A dictionary of activity labels used in the ma() function in activity(p(X), ma(Y))
```

```
pybel.language.activity_mapping = {'cat': {'identifier': '0003824', 'name': 'catalytic a
Maps the default BEL molecular activities to Gene Ontology Molecular Functions
```

```
pybel.language.compartment_mapping = {'cell surface': {'identifier': '0009986', 'name':
Maps the default BEL cellular components to Gene Ontology Cellular Components
```

```
pybel.language.abundance_labels = {'a': 'Abundance', 'abundance': 'Abundance', 'biologica
Provides a mapping from BEL terms to PyBEL internal constants
```

```
pybel.language.abundance_sbo_mapping = {'BiologicalProcess': {'identifier': '0000375', 'name': 'r
Maps the BEL abundance types to the Systems Biology Ontology
```

```
pybel.language.pmod_namespace = {'ADP-ribosylation': 'ADPRib', 'ADPRib': 'ADPRib', 'Ac': 'Ac
A dictionary of default protein modifications to their preferred value
```

```
pybel.language.pmod_mappings = {'ADPRib': {'synonyms': ['ADPRib', 'ADP-ribosylation', 'ADP-ribosylation
Use Gene Ontology children of go_0006464: "cellular protein modification process"
```

```
pybel.language.pmod_legacy_labels = {'A': 'Ac', 'F': 'Farn', 'G': 'Glyco', 'H': 'Hy', 'M': 'Me
A dictionary of legacy (BEL 1.0) default namespace protein modifications to their BEL 2.0 preferred value
```

```
pybel.language.gmod_namespace = {'ADPRib': 'ADPRib', 'M': 'Me', 'Me': 'Me', 'methylation': 'methylat
A dictionary of default gene modifications. This is a PyBEL variant to the BEL specification.
```

```
pybel.language.gmod_mappings = {'ADPRib': {'synonyms': ['ADPRib'], 'xrefs': [{"namespace": "GO", "id": "GO:0006304"}]
Use Gene Ontology children of go:0006304 ! "DNA modification"
```

CHAPTER
SEVENTEEN

PARSERS

This page is for users who want to squeeze the most bizarre possibilities out of PyBEL. Most users will not need this reference.

PyBEL makes extensive use of the PyParsing module. The code is organized to different modules to reflect the different faces of the BEL language. These parsers support BEL 2.0 and have some backwards compatibility for rewriting BEL v1.0 statements as BEL v2.0. The biologist and bioinformatician using this software will likely never need to read this page, but a developer seeking to extend the language will be interested to see the inner workings of these parsers.

See: https://github.com/OpenBEL/language/blob/master/version_2.0/MIGRATE_BEL1_BEL2.md

17.1 BEL Parser

```
class pybel.parser.parse_bel.BELParser(graph, namespace_to_term_to_encoding=None,
                                         namespace_to_pattern=None, annotation_to_term=None, annotation_to_pattern=None, annotation_to_local=None,
                                         allow_naked_names=False, disallow_nested=False, disallow_unqualified_translocations=False, citation_clearing=True, skip_validation=False,
                                         autostreamline=True, required_annotations=None)
```

Build a parser backed by a given dictionary of namespaces.

Build a BEL parser.

Parameters

- **graph** (`pybel.BELGraph`) – The BEL Graph to use to store the network
- **namespace_to_term_to_encoding** (`Optional[Mapping[str, Mapping[Tuple[Optional[str], str], str]]]`) – A dictionary of {namespace: {name: encoding}}. Delegated to `pybel.parser.parse_identifier.IdentifierParser`
- **namespace_to_pattern** (`Optional[Mapping[str, Pattern]]`) – A dictionary of {namespace: regular expression strings}. Delegated to `pybel.parser.parse_identifier.IdentifierParser`
- **annotation_to_term** (`Optional[Mapping[str, Set[str]]]`) – A dictionary of {annotation: set of values}. Delegated to `pybel.parser.ControlParser`
- **annotation_to_pattern** (`Optional[Mapping[str, Pattern]]`) – A dictionary of {annotation: regular expression strings}. Delegated to `pybel.parser.ControlParser`

- **annotation_to_local** (`Optional[Mapping[str, Set[str]]]`) – A dictionary of {annotation: set of values}. Delegated to `pybel.parser.ControlParser`
- **allow_naked_names** (`bool`) – If true, turn off naked namespace failures. Delegated to `pybel.parser.parse_identifier.IdentifierParser`
- **disallow_nested** (`bool`) – If true, turn on nested statement failures. Delegated to `pybel.parser.parse_identifier.IdentifierParser`
- **disallow_unqualified_translocations** (`bool`) – If true, allow translocations without TO and FROM clauses.
- **citation_clearing** (`bool`) – Should SET Citation statements clear evidence and all annotations? Delegated to `pybel.parser.ControlParser`
- **autostreamline** (`bool`) – Should the parser be streamlined on instantiation?
- **required_annotations** (`Optional[List[str]]`) – Optional list of required annotations

pmod = None

2.2.1

location = None

2.2.4

gmod = None

PyBEL BEL Specification variant

fusion = None

2.6.1

general_abundance = None

2.1.1

gene = None

2.1.4

mirna = None

2.1.5

protein = None

2.1.6

rna = None

2.1.7

complex_singleton = None

2.1.2

composite_abundance = None

2.1.3

molecular_activity = None

2.4.1

biological_process = None

2.3.1

pathology = None

2.3.2

activity = None

2.3.3

```
translocation = None
    2.5.1
```

```
degradation = None
    2.5.2
```

```
reactants = None
    2.5.3
```

```
rate_limit = None
    3.1.5
```

```
subprocess_of = None
    3.4.6
```

```
transcribed = None
    3.3.2
```

```
translated = None
    3.3.3
```

```
has_member = None
    3.4.1
```

```
abundance_list = None
    3.4.2
```

```
get_annotations()
    Get the current annotations in this parser.
```

Return type `Dict`

```
clear()
```

Clear the graph and all control parser data (current citation, annotations, and statement group).

```
handle_nested_relation(line, position, tokens)
```

Handle nested statements.

If `self.disallow_nested` is True, raises a `NestedRelationWarning`.

Raises `NestedRelationWarning`

```
check_function_semantics(line, position, tokens)
```

Raise an exception if the function used on the tokens is wrong.

Raises `InvalidFunctionSemantic`

Return type `ParseResults`

```
handle_term(_, __, tokens)
```

Handle BEL terms (the subject and object of BEL relations).

Return type `ParseResults`

```
handle_has_members(_, __, tokens)
```

Handle list relations like `p(X) hasMembers list(p(Y), p(Z), ...)`.

Return type `ParseResults`

```
handle_has_components(_, __, tokens)
```

Handle list relations like `p(X) hasComponents list(p(Y), p(Z), ...)`.

Return type `ParseResults`

```
handle_unqualified_relation(_, __, tokens)
```

Handle unqualified relations.

Return type ParseResults

handle_inverse_unqualified_relation(*_*, *__*, *tokens*)
Handle unqualified relations that should go reverse.

Return type ParseResults

ensure_node(*tokens*)
Turn parsed tokens into canonical node name and makes sure its in the graph.

Return type BaseEntity

handle_translocation_illegal(*line*, *position*, *tokens*)
Handle a malformed translocation.

Return type None

```
pybel.io.line_utils.parse_lines(graph, lines, manager=None, disallow_nested=False,
                                 citation_clearing=True, use_tqdm=False,
                                 tqdm_kwargs=None, no_identifier_validation=False,
                                 disallow_unqualified_translocations=False, allow_redefinition=False,
                                 allow_naked_names=False, required_annotations=None,
                                 upgrade_urls=False)
```

Parse an iterable of lines into this graph.

Delegates to `parse_document()`, `parse_definitions()`, and `parse_statements()`.

Parameters

- **graph** (BELGraph) – A BEL graph
- **lines** (Iterable[str]) – An iterable over lines of BEL script
- **manager** (Optional[Manager]) – A PyBEL database manager
- **disallow_nested** (bool) – If true, turns on nested statement failures
- **citation_clearing** (bool) – Should SET Citation statements clear evidence and all annotations? Delegated to `pybel.parser.ControlParser`
- **use_tqdm** (bool) – Use tqdm to show a progress bar?
- **tqdm_kwargs** (Optional[Mapping[str, Any]]) – Keywords to pass to tqdm
- **disallow_unqualified_translocations** (bool) – If true, allow translocations without TO and FROM clauses.
- **required_annotations** (Optional[List[str]]) – Annotations that are required for all statements
- **upgrade_urls** (bool) – Automatically upgrade old namespace URLs. Defaults to false.

Warning: These options allow concessions for parsing BEL that is either **WRONG** or **UNSCIENTIFIC**. Use them at risk to reproducibility and validity of your results.

Parameters

- **no_identifier_validation** (bool) – If true, turns off namespace validation
- **allow_naked_names** (bool) – If true, turns off naked namespace failures
- **allow_redefinition** (bool) – If true, doesn't fail on second definition of same name or annotation

- **allow_definition_failures** (`bool`) – If true, allows parsing to continue if a terminology file download/parse fails

Return type `None`

17.2 Metadata Parser

```
class pybel.parser.parse_metadata.MetadataParser(manager, namespace_to_term_to_encoding=None, namespace_to_pattern=None, annotation_to_term=None, annotation_to_pattern=None, annotation_to_local=None, default_namespace=None, allow_redefinition=False, skip_validation=False, uprade_urls=False)
```

A parser for the document and definitions section of a BEL document.

See also:

BEL 1.0 Specification for the `DEFINE` keyword

Build a metadata parser.

Parameters

- **manager** – A cache manager
- **namespace_to_term_to_encoding** (`Optional[Mapping[str, Mapping[Tuple[Optional[str], str], str]]]`) – An enumerated namespace mapping from {namespace keyword: {(identifier, name): encoding}}
- **namespace_to_pattern** (`Optional[Mapping[str, Pattern]]`) – A regular expression namespace mapping from {namespace keyword: regex string}
- **annotation_to_term** (`Optional[Mapping[str, Set[str]]]`) – Enumerated annotation mapping from {annotation keyword: set of valid values}
- **annotation_to_pattern** (`Optional[Mapping[str, Pattern]]`) – Regular expression annotation mapping from {annotation keyword: regex string}
- **default_namespace** (`Optional[Set[str]]`) – A set of strings that can be used without a namespace
- **skip_validation** (`bool`) – If true, don't download and cache namespaces/annotations

manager = None

This metadata parser's internal definition cache manager

namespace_to_term_to_encoding = None

A dictionary of cached {namespace keyword: {(identifier, name): encoding}}

uncachable_namespaces = None

A set of namespaces's URLs that can't be cached

namespace_to_pattern = None

A dictionary of {namespace keyword: regular expression string}

default_namespace = None

A set of names that can be used without a namespace

```
annotation_to_term = None
    A dictionary of cached {annotation keyword: set of values}

annotation_to_pattern = None
    A dictionary of {annotation keyword: regular expression string}

annotation_to_local = None
    A dictionary of cached {annotation keyword: set of values}

document_metadata = None
    A dictionary containing the document metadata

namespace_url_dict = None
    A dictionary from {namespace keyword: BEL namespace URL}

annotation_url_dict = None
    A dictionary from {annotation keyword: BEL annotation URL}

handle_document(line, position, tokens)
    Handle statements like SET DOCUMENT X = "Y".
        Raises InvalidMetadataException
        Raises VersionFormatWarning
        Return type ParseResults

raise_for_redefined_namespace(line, position, namespace)
    Raise an exception if a namespace is already defined.
        Raises RedefinedNamespaceError
        Return type None

handle_namespace_url(line, position, tokens)
    Handle statements like DEFINE NAMESPACE X AS URL "Y".
        Raises RedefinedNamespaceError
        Raises pybel.resources.exc.ResourceError
        Return type ParseResults

handle_namespace_pattern(line, position, tokens)
    Handle statements like DEFINE NAMESPACE X AS PATTERN "Y".
        Raises RedefinedNamespaceError
        Return type ParseResults

raise_for_redefined_annotation(line, position, annotation)
    Raise an exception if the given annotation is already defined.
        Raises RedefinedAnnotationError
        Return type None

handle_annotations_url(line, position, tokens)
    Handle statements like DEFINE ANNOTATION X AS URL "Y".
        Raises RedefinedAnnotationError
        Return type ParseResults

handle_annotation_list(line, position, tokens)
    Handle statements like DEFINE ANNOTATION X AS LIST {"Y", "Z", ...}.
```

Raises RedefinedAnnotationError

Return type ParseResults

handle_annotation_pattern (*line, position, tokens*)
Handle statements like DEFINE ANNOTATION X AS PATTERN "Y".

Raises RedefinedAnnotationError

Return type ParseResults

has_enumerated_annotation (*annotation*)
Check if this annotation is defined by an enumeration.

Return type bool

has_regex_annotation (*annotation*)
Check if this annotation is defined by a regular expression.

Return type bool

has_local_annotation (*annotation*)
Check if this annotation is defined by an locally.

Return type bool

has_annotation (*annotation*)
Check if this annotation is defined.

Return type bool

has_enumerated_namespace (*namespace*)
Check if this namespace is defined by an enumeration.

Return type bool

has_regex_namespace (*namespace*)
Check if this namespace is defined by a regular expression.

Return type bool

has_namespace (*namespace*)
Check if this namespace is defined.

Return type bool

raise_for_version (*line, position, version*)
Check that a version string is valid for BEL documents.

This means it's either in the YYYYMMDD or semantic version format.

Parameters

- **line** (str) – The line being parsed
- **position** (int) – The position in the line being parsed
- **version** (str) – A version string

Raises VersionFormatWarning

Return type None

17.3 Control Parser

```
class pybel.parser.parse_control.ControlParser(annotation_to_term=None,           an-
                                                notation_to_pattern=None,      an-
                                                notation_to_local=None,       ci-
                                                citation_clearing=True,      re-
                                                required_annotations=None)
```

A parser for BEL control statements.

See also:

BEL 1.0 specification on [control records](#)

Initialize the control statement parser.

Parameters

- **annotation_to_term** (`Optional[Mapping[str, Set[str]]]`) – A dictionary of {annotation: set of valid values} defined with URL for parsing
- **annotation_to_pattern** (`Optional[Mapping[str, Pattern]]`) – A dictionary of {annotation: regular expression string}
- **annotation_to_local** (`Optional[Mapping[str, Set[str]]]`) – A dictionary of {annotation: set of valid values} for parsing defined with LIST
- **citation_clearing** (`bool`) – Should SET Citation statements clear evidence and all annotations?
- **required_annotations** (`Optional[List[str]]`) – Annotations that are required

property citation_is_set

Check if the citation is set.

Return type `bool`

has_enumerated_annotation(annotation)

Check if the annotation is defined as an enumeration.

Return type `bool`

has_regex_annotation(annotation)

Check if the annotation is defined as a regular expression.

Return type `bool`

has_local_annotation(annotation)

Check if the annotation is defined locally.

Return type `bool`

has_annotation(annotation)

Check if the annotation is defined.

Return type `bool`

raise_for_undefined_annotation(line, position, annotation)

Raise an exception if the annotation is not defined.

Raises `UndefinedAnnotationWarning`

Return type `None`

raise_for_invalid_annotation_value(line, position, key, value)

Raise an exception if the annotation is not defined.

Raises IllegalAnnotationValueWarning or MissingAnnotationRegexWarning

Return type None

raise_for_missing_citation(*line, position*)

Raise an exception if there is no citation present in the parser.

Raises MissingCitationException

Return type None

handle_annotation_key(*line, position, tokens*)

Handle an annotation key before parsing to validate that it's either enumerated or as a regex.

Raise MissingCitationException or UndefinedAnnotationWarning

Return type ParseResults

handle_set_statement_group(*_, __, tokens*)

Handle a SET STATEMENT_GROUP = "X" statement.

Return type ParseResults

handle_set_citation(*line, position, tokens*)

Handle a SET Citation = {"X", "Y", "Z", ...} statement.

Return type ParseResults

handle_set_evidence(*_, __, tokens*)

Handle a SET Evidence = "" statement.

Return type ParseResults

handle_set_command(*line, position, tokens*)

Handle a SET X = "Y" statement.

Return type ParseResults

handle_set_command_list(*line, position, tokens*)

Handle a SET X = {"Y", "Z", ...} statement.

Return type ParseResults

handle_unset_statement_group(*line, position, tokens*)

Unset the statement group, or raises an exception if it is not set.

Raises MissingAnnotationKeyWarning

Return type ParseResults

handle_unset_citation(*line, position, tokens*)

Unset the citation, or raise an exception if it is not set.

Raises MissingAnnotationKeyWarning

Return type ParseResults

handle_unset_evidence(*line, position, tokens*)

Unset the evidence, or throws an exception if it is not already set.

The value for `tokens` [EVIDENCE] corresponds to which alternate of SupportingText or Evidence was used in the BEL script.

Raises MissingAnnotationKeyWarning

Return type ParseResults

```
validate_unset_command(line, position, annotation)
    Raise an exception when trying to UNSET X if X is not already set.

    Raises MissingAnnotationKeyWarning

    Return type None

handle_unset_command(line, position, tokens)
    Handle an UNSET X statement or raises an exception if it is not already set.

    Raises MissingAnnotationKeyWarning

    Return type ParseResults

handle_unset_list(line, position, tokens)
    Handle UNSET {A, B, ...} or raises an exception of any of them are not present.

    Consider that all unsets are in peril if just one of them is wrong!

    Raises MissingAnnotationKeyWarning

    Return type ParseResults

handle_unset_all(_, __, tokens)
    Handle an UNSET_ALL statement.

    Return type ParseResults

get_annotations()
    Get the current annotations.

    Return type Dict

get_citationReturn type CitationDict

get_missing_required_annotations()
    Return missing required annotations.

    Return type List[str]

clear_citation()
    Clear the citation and if citation clearing is enabled, clear the evidence and annotations.

    Return type None

clear()
    Clear the statement_group, citation, evidence, and annotations.

    Return type None
```

17.4 Concept Parser

```
class pybel.parser.parse_concept.ConceptParser(namespace_to_term_to_encoding=None,
                                                namespace_to_pattern=None,
                                                default_namespace=None,           al-
                                                low_naked_names=False,
                                                skip_validation=False)
```

A parser for concepts in the form of namespace:name or namespace:identifier!name.

Can be made more lenient when given a default namespace or enabling the use of naked names.

Initialize the concept parser.

Parameters

- **namespace_to_term_to_encoding** (`Optional[Mapping[str, Mapping[Tuple[Optional[str], str], str]]]`) – A dictionary of {namespace: {(identifier, name): encoding}}
- **namespace_to_pattern** (`Optional[Mapping[str, Pattern]]`) – A dictionary of {namespace: regular expression string} to compile
- **default_namespace** (`Optional[Set[str]]`) – A set of strings that can be used without a namespace
- **allow_naked_names** (`bool`) – If true, turn off naked namespace failures

`has_enumerated_namespace(namespace)`

Check that the namespace has been defined by an enumeration.

Return type `bool`

`has_regex_namespace(namespace)`

Check that the namespace has been defined by a regular expression.

Return type `bool`

`raise_for_missing_namespace(line, position, namespace, name)`

Raise an exception if the namespace is not defined.

Return type `None`

`raise_for_missing_name(line, position, namespace, name)`

Raise an exception if the namespace is not defined or if it does not validate the given name.

Return type `None`

`handle_identifier_fqualified(line, position, tokens)`

Handle parsing a qualified OBO-style identifier.

Return type `ParseResults`

`handle_identifier_qualified(line, position, tokens)`

Handle parsing a qualified identifier.

Return type `ParseResults`

`handle_namespace_default(line, position, tokens)`

Handle parsing an identifier for the default namespace.

Return type `ParseResults`

`static handle_namespace_lenient(line, position, tokens)`

Handle parsing an identifier for names missing a namespace that are outside the default namespace.

Return type `ParseResults`

`handle_namespace_invalid(line, position, tokens)`

Raise an exception when parsing a name missing a namespace.

Return type `None`

17.5 Sub-Parsers

Parsers for modifications to abundances.

INTERNAL DOMAIN SPECIFIC LANGUAGE

PyBEL implements an internal domain-specific language (DSL).

This enables you to write BEL using Python scripts. Even better, you can programmatically generate BEL using Python. See the Bio2BEL [paper](#) and [repository](#) for many examples.

Internally, the BEL parser converts BEL script into the BEL DSL then adds it to a BEL graph object. When you iterate through the `pybel.BELGraph`, the nodes are instances of subclasses of `pybel.dsl.BaseEntity`.

18.1 Primitives

```
class pybel.dsl.Entity(*, namespace, name=None, identifier=None)
```

Represents a named entity with a namespace and name/identifier.

Create a dictionary representing a reference to an entity.

Parameters

- `namespace (str)` – The namespace to which the entity belongs
- `name (Optional[str])` – The name of the entity
- `identifier (Optional[str])` – The identifier of the entity in the namespace

```
class pybel.dsl BaseEntity
```

This is the superclass for all BEL terms.

A BEL term has three properties:

1. It has a type. Subclasses of this function should set the class variable `function`.
2. It can be converted to BEL. Note, this is an abstract class, so all sub-classes must implement this functionality in `as_bel()`.
3. It can be hashed, based on the BEL conversion

```
class pybel.dsl.BaseAbundance(namespace, name=None, identifier=None, xrefs=None)
```

The superclass for all named BEL terms.

A named BEL term has:

1. A type (taken care of by being a subclass of `BaseEntity`)
2. A named `Entity`. Though this doesn't directly inherit from `Entity`, it creates one internally using the namespace, identifier, and name. Ideally, both the identifier and name are given. If one is missing, it can be looked up with `pybel.grounding.ground()`
3. An optional list of xrefs, corresponding to the whole entity, not just the namespace/name. For example, the BEL term `p(HGNC:APP, frag(672_713)` could xref CHEBI:64647.

Build an abundance from a function, namespace, and a name and/or identifier.

Parameters

- **namespace** (`str`) – The name of the namespace
- **name** (`Optional[str]`) – The name of this abundance
- **identifier** (`Optional[str]`) – The database identifier for this abundance
- **xrefs** (`Optional[List[Entity]]`) – Alternate identifiers for the entity

```
class pybel.dsl.ListAbundance(members)
```

The superclass for all BEL terms defined by lists, as opposed to by names like in `BaseAbundance`.

Build a list abundance node.

Parameters `members` (`Union[BaseAbundance, Iterable[BaseAbundance]]`) – A list of PyBEL node data dictionaries

18.2 Named Entities

```
class pybel.dsl.Abundance(namespace, name=None, identifier=None, xrefs=None)
```

Builds an abundance node.

```
>>> from pybel.dsl import Abundance
>>> Abundance(namespace='CHEBI', name='water')
```

Build an abundance from a function, namespace, and a name and/or identifier.

Parameters

- **namespace** (`str`) – The name of the namespace
- **name** (`Optional[str]`) – The name of this abundance
- **identifier** (`Optional[str]`) – The database identifier for this abundance
- **xrefs** (`Optional[List[Entity]]`) – Alternate identifiers for the entity

```
class pybel.dsl.BiologicalProcess(namespace, name=None, identifier=None, xrefs=None)
```

Builds a biological process node.

```
>>> from pybel.dsl import BiologicalProcess
>>> BiologicalProcess(namespace='GO', name='apoptosis')
```

Build an abundance from a function, namespace, and a name and/or identifier.

Parameters

- **namespace** (`str`) – The name of the namespace
- **name** (`Optional[str]`) – The name of this abundance
- **identifier** (`Optional[str]`) – The database identifier for this abundance
- **xrefs** (`Optional[List[Entity]]`) – Alternate identifiers for the entity

```
class pybel.dsl.Pathology(namespace, name=None, identifier=None, xrefs=None)
```

Build a pathology node.

```
>>> from pybel.dsl import Pathology
>>> Pathology(namespace='DO', name='Alzheimer Disease')
```

Build an abundance from a function, namespace, and a name and/or identifier.

Parameters

- **namespace** (`str`) – The name of the namespace
- **name** (`Optional[str]`) – The name of this abundance
- **identifier** (`Optional[str]`) – The database identifier for this abundance
- **xrefs** (`Optional[List[Entity]]`) – Alternate identifiers for the entity

```
class pybel.dsl.Population(namespace, name=None, identifier=None, xrefs=None)
```

Builds a population node.

```
>>> from pybel.dsl import Population
>>> Population(namespace='uberon', name='blood')
```

Build an abundance from a function, namespace, and a name and/or identifier.

Parameters

- **namespace** (`str`) – The name of the namespace
- **name** (`Optional[str]`) – The name of this abundance
- **identifier** (`Optional[str]`) – The database identifier for this abundance
- **xrefs** (`Optional[List[Entity]]`) – Alternate identifiers for the entity

18.3 Central Dogma

```
class pybel.dsl.CentralDogma(namespace, name=None, identifier=None, xrefs=None, variants=None)
```

The base class for “central dogma” abundances (i.e., genes, miRNAs, RNAs, and proteins).

Build a node for a gene, RNA, miRNA, or protein.

Parameters

- **namespace** (`str`) – The name of the database used to identify this entity
- **name** (`Optional[str]`) – The database’s preferred name or label for this entity
- **identifier** (`Optional[str]`) – The database’s identifier for this entity
- **xrefs** (`Optional[List[Entity]]`) – Alternative database cross references
- **variants** (`Union[None, Variant, Iterable[Variant]]`) – An optional variant or list of variants

```
class pybel.dsl.Gene(namespace, name=None, identifier=None, xrefs=None, variants=None)
```

Builds a gene node.

Build a node for a gene, RNA, miRNA, or protein.

Parameters

- **namespace** (`str`) – The name of the database used to identify this entity
- **name** (`Optional[str]`) – The database’s preferred name or label for this entity
- **identifier** (`Optional[str]`) – The database’s identifier for this entity
- **xrefs** (`Optional[List[Entity]]`) – Alternative database cross references

- **variants** (`Union[None, Variant, Iterable[Variant]]`) – An optional variant or list of variants

```
class pybel.dsl.Transcribable(namespace, name=None, identifier=None, xrefs=None, variants=None)
```

A base class for RNA and micro-RNA to share getting of their corresponding genes.

Build a node for a gene, RNA, miRNA, or protein.

Parameters

- **namespace** (`str`) – The name of the database used to identify this entity
- **name** (`Optional[str]`) – The database's preferred name or label for this entity
- **identifier** (`Optional[str]`) – The database's identifier for this entity
- **xrefs** (`Optional[List[Entity]]`) – Alternative database cross references
- **variants** (`Union[None, Variant, Iterable[Variant]]`) – An optional variant or list of variants

```
class pybel.dsl.Rna(namespace, name=None, identifier=None, xrefs=None, variants=None)
```

Builds an RNA node.

Example: AKT1 protein coding gene's RNA:

```
>>> from pybel.dsl import Rna
>>> Rna(namespace='HGNC', name='AKT1', identifier='391')
```

Non-coding RNAs can also be encoded such as U85:

```
>>> from pybel.dsl import Rna
>>> Rna(namespace='SNORNABASE', identifier='SR0000073')
```

Build a node for a gene, RNA, miRNA, or protein.

Parameters

- **namespace** (`str`) – The name of the database used to identify this entity
- **name** (`Optional[str]`) – The database's preferred name or label for this entity
- **identifier** (`Optional[str]`) – The database's identifier for this entity
- **xrefs** (`Optional[List[Entity]]`) – Alternative database cross references
- **variants** (`Union[None, Variant, Iterable[Variant]]`) – An optional variant or list of variants

```
class pybel.dsl.MicroRna(namespace, name=None, identifier=None, xrefs=None, variants=None)
```

Represents an micro-RNA.

Human miRNA's are listed on HUGO's MicroRNAs (MIR) gene family.

MIR1-1 from HGNC:

```
>>> from pybel.dsl import MicroRna
>>> MicroRna(namespace='HGNC', name='MIR1-1', identifier='31499')
```

MIR1-1 from miRBase:

```
>>> from pybel.dsl import MicroRna
>>> MicroRna(namespace='MIRBASE', identifier='MI0000651')
```

MIR1-1 from Entrez Gene

```
>>> from pybel.dsl import MicroRna
>>> MicroRna(namespace='ENTREZ', identifier='406904')
```

Build a node for a gene, RNA, miRNA, or protein.

Parameters

- **namespace** (str) – The name of the database used to identify this entity
- **name** (Optional[str]) – The database’s preferred name or label for this entity
- **identifier** (Optional[str]) – The database’s identifier for this entity
- **xrefs** (Optional[List[Entity]]) – Alternative database cross references
- **variants** (Union[None, Variant, Iterable[Variant]]) – An optional variant or list of variants

class pybel.dsl.Protein(namespace, name=None, identifier=None, xrefs=None, variants=None)

Builds a protein node.

Example: AKT

```
>>> from pybel.dsl import Protein
>>> Protein(namespace='HGNC', name='AKT1')
```

Example: AKT with optionally included HGNC database identifier

```
>>> from pybel.dsl import Protein
>>> Protein(namespace='HGNC', name='AKT1', identifier='391')
```

Example: AKT with phosphorylation

```
>>> from pybel.dsl import Protein, ProteinModification
>>> Protein(namespace='HGNC', name='AKT', variants=[ProteinModification('Ph', ↴code='Thr', position=308)])
```

Build a node for a gene, RNA, miRNA, or protein.

Parameters

- **namespace** (str) – The name of the database used to identify this entity
- **name** (Optional[str]) – The database’s preferred name or label for this entity
- **identifier** (Optional[str]) – The database’s identifier for this entity
- **xrefs** (Optional[List[Entity]]) – Alternative database cross references
- **variants** (Union[None, Variant, Iterable[Variant]]) – An optional variant or list of variants

18.3.1 Variants

```
class pybel.dsl.Variant(kind)
```

The superclass for variant dictionaries.

Build the variant data dictionary.

Parameters `kind` (`str`) – The kind of variant

```
class pybel.dsl.ProteinModification(name, code=None, position=None, namespace=None,
                                      identifier=None, xrefs=None)
```

Build a protein modification variant dictionary.

Build a protein modification variant data dictionary.

Parameters

- `name` (`str`) – The name of the modification
- `code` (`Optional[str]`) – The three letter amino acid code for the affected residue. Capital first letter.
- `position` (`Optional[int]`) – The position of the affected residue
- `namespace` (`Optional[str]`) – The namespace to which the name of this modification belongs
- `identifier` (`Optional[str]`) – The identifier of the name of the modification
- `xrefs` (`Optional[List[Entity]]`) – Alternative database xrefs

Either the name or the identifier must be used. If the namespace is omitted, it is assumed that a name is specified from the BEL default namespace.

Example from BEL default namespace:

```
>>> from pybel.dsl import ProteinModification
>>> ProteinModification('Ph', code='Thr', position=308)
```

Example from custom namespace:

```
>>> from pybel.dsl import ProteinModification
>>> ProteinModification(name='protein phosphorylation', namespace='GO', code='Thr',
   ↴', position=308)
```

Example from custom namespace additionally qualified with identifier:

```
>>> from pybel.dsl import ProteinModification
>>> ProteinModification(name='protein phosphorylation', namespace='GO',
   >>>                         identifier='0006468', code='Thr', position=308)
```

```
class pybel.dsl.GeneModification(name, namespace=None, identifier=None, xrefs=None)
```

Build a gene modification variant dictionary.

Either the name or the identifier must be used. If the namespace is omitted, it is assumed that a name is specified from the BEL default namespace.

Example from BEL default namespace:

```
>>> from pybel.dsl import GeneModification
>>> GeneModification(name='Me')
```

Example from custom namespace:

```
>>> from pybel.dsl import GeneModification
>>> GeneModification(name='DNA methylation', namespace='GO', identifier='0006306',
    ↵)
```

Build a variant that has a reference.

Parameters

- **name** (`str`) – The name of the modification
- **namespace** (`Optional[str]`) – The namespace to which the name of this modification belongs
- **identifier** (`Optional[str]`) – The identifier of the name of the modification
- **xrefs** (`Optional[List[Entity]]`) – Alternative database xrefs

Either the name or the identifier must be used. If the namespace is omitted, it is assumed that a name is specified from the BEL default namespace.

class `pybel.dsl.Hgvs` (`variant`)

Builds a HGVS variant dictionary.

Build an HGVS variant data dictionary.

Parameters **variant** (`str`) – The HGVS variant string

```
>>> from pybel.dsl import Protein, Hgvs
>>> Protein(namespace='HGNC', name='AKT1', variants=[Hgvs('P.Alal27Tyr')])
```

class `pybel.dsl.HgvsReference`

Represents the “reference” variant in HGVS.

Build an HGVS variant data dictionary.

Parameters **variant** – The HGVS variant string

```
>>> from pybel.dsl import Protein, Hgvs
>>> Protein(namespace='HGNC', name='AKT1', variants=[Hgvs('P.Alal27Tyr')])
```

class `pybel.dsl.HgvsUnspecified`

Represents an unspecified variant in HGVS.

Build an HGVS variant data dictionary.

Parameters **variant** – The HGVS variant string

```
>>> from pybel.dsl import Protein, Hgvs
>>> Protein(namespace='HGNC', name='AKT1', variants=[Hgvs('P.Alal27Tyr')])
```

class `pybel.dsl.ProteinSubstitution` (`from_aa, position, to_aa`)

A protein substitution variant.

Build an HGVS variant data dictionary for the given protein substitution.

Parameters

- **from_aa** (`str`) – The 3-letter amino acid code of the original residue
- **position** (`int`) – The position of the residue
- **to_aa** (`str`) – The 3-letter amino acid code of the new residue

```
>>> from pybel.dsl import Protein, ProteinSubstitution
>>> Protein(namespace='HGNC', name='AKT1', variants=[ProteinSubstitution('Ala', ↵127, 'Tyr')])
```

class pybel.dsl.Fragment (*start=None, stop=None, description=None*)

Represent the information about a protein fragment.

Build a protein fragment data dictionary.

Parameters

- **start** (Union[None, int, str]) – The starting position
- **stop** (Union[None, int, str]) – The stopping position
- **description** (Optional[str]) – An optional description

Example of specified fragment:

```
>>> from pybel.dsl import Protein, Fragment
>>> Protein(name='APP', namespace='HGNC', variants=[Fragment(start=672, ↵stop=713)])
```

Example of unspecified fragment:

```
>>> from pybel.dsl import Protein, Fragment
>>> Protein(name='APP', namespace='HGNC', variants=[Fragment()])
```

18.4 Fusions

class pybel.dsl.FusionBase (*partner_5p, partner_3p, range_5p=None, range_3p=None*)

The superclass for building fusion node data dictionaries.

Build a fusion node.

Parameters

- **partner_5p** (CentralDogma) – A PyBEL node for the 5-prime partner
- **partner_3p** (CentralDogma) – A PyBEL node for the 3-prime partner
- **range_5p** (Optional[FusionRangeBase]) – A fusion range for the 5-prime partner
- **range_3p** (Optional[FusionRangeBase]) – A fusion range for the 3-prime partner

class pybel.dsl.GeneFusion (*partner_5p, partner_3p, range_5p=None, range_3p=None*)

Builds a gene fusion node.

Example, using fusion ranges with the ‘c’ qualifier

```
>>> from pybel.dsl import GeneFusion, Gene
>>> GeneFusion(
>>> ... partner_5p=Gene(namespace='HGNC', name='TMPRSS2'),
>>> ... range_5p=EnumeratedFusionRange('c', 1, 79),
>>> ... partner_3p=Gene(namespace='HGNC', name='ERG'),
>>> ... range_3p=EnumeratedFusionRange('c', 312, 5034)
>>> )
```

Example with missing fusion ranges:

```
>>> from pybel.dsl import GeneFusion, Gene
>>> GeneFusion(
>>> ... partner_5p=Gene(namespace='HGNC', name='TMPRSS2'),
>>> ... partner_3p=Gene(namespace='HGNC', name='ERG'),
>>> )
```

Build a fusion node.

Parameters

- **partner_5p** (CentralDogma) – A PyBEL node for the 5-prime partner
- **partner_3p** (CentralDogma) – A PyBEL node for the 3-prime partner
- **range_5p** (Optional[FusionRangeBase]) – A fusion range for the 5-prime partner
- **range_3p** (Optional[FusionRangeBase]) – A fusion range for the 3-prime partner

```
class pybel.dsl.RnaFusion(partner_5p, partner_3p, range_5p=None, range_3p=None)
```

Builds an RNA fusion node.

Example, with fusion ranges using the ‘r’ qualifier:

```
>>> from pybel.dsl import RnaFusion, Rna
>>> RnaFusion(
>>> ... partner_5p=Rna(namespace='HGNC', name='TMPRSS2'),
>>> ... range_5p=EnumeratedFusionRange('r', 1, 79),
>>> ... partner_3p=Rna(namespace='HGNC', name='ERG'),
>>> ... range_3p=EnumeratedFusionRange('r', 312, 5034)
>>> )
```

Example with missing fusion ranges:

```
>>> from pybel.dsl import RnaFusion, Rna
>>> RnaFusion(
>>> ... partner_5p=Rna(namespace='HGNC', name='TMPRSS2'),
>>> ... partner_3p=Rna(namespace='HGNC', name='ERG'),
>>> )
```

Build a fusion node.

Parameters

- **partner_5p** (CentralDogma) – A PyBEL node for the 5-prime partner
- **partner_3p** (CentralDogma) – A PyBEL node for the 3-prime partner
- **range_5p** (Optional[FusionRangeBase]) – A fusion range for the 5-prime partner
- **range_3p** (Optional[FusionRangeBase]) – A fusion range for the 3-prime partner

```
class pybel.dsl.ProteinFusion(partner_5p, partner_3p, range_5p=None, range_3p=None)
```

Builds a protein fusion node.

Build a fusion node.

Parameters

- **partner_5p** (CentralDogma) – A PyBEL node for the 5-prime partner
- **partner_3p** (CentralDogma) – A PyBEL node for the 3-prime partner
- **range_5p** (Optional[FusionRangeBase]) – A fusion range for the 5-prime partner
- **range_3p** (Optional[FusionRangeBase]) – A fusion range for the 3-prime partner

18.4.1 Fusion Ranges

```
class pybel.dsl.FusionRangeBase
    The superclass for fusion range data dictionaries.
```

```
class pybel.dsl.EnumeratedFusionRange (reference, start, stop)
    Represents an enumerated fusion range.
```

Build an enumerated fusion range.

Parameters

- **reference** (`str`) – The reference code
- **or str start** (`int`) – The start position, either specified by its integer position, or ‘?’
- **or str stop** (`int`) – The stop position, either specified by its integer position, ‘?’, or ‘*’

Example fully specified RNA fusion range:

```
>>> EnumeratedFusionRange ('r', 1, 79)
```

```
class pybel.dsl.MissingFusionRange
```

Represents a fusion range with no defined start or end.

Build a missing fusion range.

18.4.2 List Abundances

```
class pybel.dsl.ComplexAbundance (members, namespace=None, name=None, identifier=None,
                                    xrefs=None)
```

Build a complex abundance node with the optional ability to specify a name.

Build a complex list node.

Parameters

- **members** (`Iterable[BaseAbundance]`) – A list of PyBEL node data dictionaries
- **namespace** (`Optional[str]`) – The namespace from which the name originates
- **name** (`Optional[str]`) – The name of the complex
- **identifier** (`Optional[str]`) – The identifier in the namespace in which the name originates
- **xrefs** (`Optional[List[Entity]]`) – Alternate identifiers for the entity if it is named

```
class pybel.dsl.CompositeAbundance (members)
```

Build a composite abundance node.

This node is effectively the “AND” inside BEL, which can help represent when two things need to be true at the same time. For example, in COVID 19, if both the NF-KB and IL6-STAT complex are present, then acute respiratory distress syndrome happens.

```
>>> from pybel.dsl import CompositeAbundance, ComplexAbundance, Protein, ...
>>> CompositeAbundance([
...     NamedComplexAbundance('fplx', 'nfkb'),
...     ComplexAbundance([
...         Protein('hgnc', identifier='6018', name='IL6'),
...         Protein('hgnc', identifier='11364', name='STAT3'),
```

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```
...    ]),  
... ])
```

Build a list abundance node.

Parameters `members` (`Union[BaseAbundance, Iterable[BaseAbundance]]`) – A list of PyBEL node data dictionaries

```
class pybel.dsl.Reaction(reactants, products)
```

Build a reaction node.

Build a reaction node.

Parameters

- `reactants` (`Union[BaseAbundance, Iterable[BaseAbundance]]`) – A list of PyBEL node data dictionaries representing the reactants
- `products` (`Union[BaseAbundance, Iterable[BaseAbundance]]`) – A list of PyBEL node data dictionaries representing the products

```
>>> from pybel.dsl import Reaction, Protein, Abundance  
>>> Reaction([Protein(namespace='HGNC', name='KNG1')], [Abundance(namespace='CHEBI  
<br>', name='bradykinin')])
```

18.5 Utilities

The following functions are useful to build DSL objects from dictionaries:

```
pybel.tokens.parse_result_to_dsl(tokens)
```

Convert a ParseResult to a PyBEL DSL object.

Return type `BaseEntity`

LOGGING MESSAGES

19.1 Errors

This module contains base exceptions that are shared through the package.

`exception pybel.exceptions.PyBELWarning`

The base class for warnings during compilation from which PyBEL can recover.

19.2 Parse Exceptions

Exceptions for the BEL parser.

A message for “General Parser Failure” is displayed when a problem was caused due to an unforeseen error. The line number and original statement are printed for the user to debug.

`exception pybel.parser.exc.BELParserWarning (line_number, line, position, *args)`

The base PyBEL parser exception, which holds the line and position where a parsing problem occurred.

Initialize the BEL parser warning.

Parameters

- `line_number (int)` – The line number on which this warning occurred
- `line (str)` – The content of the line
- `position (int)` – The position within the line where the warning occurred

`exception pybel.parser.exc.BELSyntaxError (line_number, line, position, *args)`

For general syntax errors.

Initialize the BEL parser warning.

Parameters

- `line_number (int)` – The line number on which this warning occurred
- `line (str)` – The content of the line
- `position (int)` – The position within the line where the warning occurred

`exception pybel.parser.exc.InconsistentDefinitionError (line_number, line, position, definition)`

Base PyBEL error for redefinition.

Initialize the BEL parser warning.

Parameters

- **line_number** (`int`) – The line number on which this warning occurred
- **line** (`str`) – The content of the line
- **position** (`int`) – The position within the line where the warning occurred

exception `pybel.parser.exc.RedefinedNamespaceError` (`line_number, line, position, definition`)

Raised when a namespace is redefined.

Initialize the BEL parser warning.

Parameters

- **line_number** (`int`) – The line number on which this warning occurred
- **line** (`str`) – The content of the line
- **position** (`int`) – The position within the line where the warning occurred

exception `pybel.parser.exc.RedefinedAnnotationError` (`line_number, line, position, definition`)

Raised when an annotation is redefined.

Initialize the BEL parser warning.

Parameters

- **line_number** (`int`) – The line number on which this warning occurred
- **line** (`str`) – The content of the line
- **position** (`int`) – The position within the line where the warning occurred

exception `pybel.parser.exc.NameWarning` (`line_number, line, position, name, *args`)

The base class for errors related to nomenclature.

Build a warning wrapping a given name.

exception `pybel.parser.exc.NakedNameWarning` (`line_number, line, position, name, *args`)

Raised when there is an identifier without a namespace. Enable lenient mode to suppress.

Build a warning wrapping a given name.

exception `pybel.parser.exc.MissingDefaultNameWarning` (`line_number, line, position, name, *args`)

Raised if reference to value not in default namespace.

Build a warning wrapping a given name.

exception `pybel.parser.exc.NamespaceIdentifierWarning` (`line_number, line, position, namespace, name`)

The base class for warnings related to namespace:name identifiers.

Initialize the namespace identifier warning.

Parameters

- **line_number** (`int`) – The line number of the line that caused the exception
- **line** (`str`) – The line that caused the exception
- **position** (`int`) – The line's position of the exception
- **namespace** (`str`) – The namespace of the identifier
- **name** (`str`) – The name of the identifier

```
exception pybel.parser.exc.UndefinedNamespaceWarning(line_number, line, position,
                                                     namespace, name)
```

Raised if reference made to undefined namespace.

Initialize the namespace identifier warning.

Parameters

- **line_number** (`int`) – The line number of the line that caused the exception
- **line** (`str`) – The line that caused the exception
- **position** (`int`) – The line's position of the exception
- **namespace** (`str`) – The namespace of the identifier
- **name** (`str`) – The name of the identifier

```
exception pybel.parser.exc.MissingNamespaceNameWarning(line_number, line, position,
                                                       namespace, name)
```

Raised if reference to value not in namespace.

Initialize the namespace identifier warning.

Parameters

- **line_number** (`int`) – The line number of the line that caused the exception
- **line** (`str`) – The line that caused the exception
- **position** (`int`) – The line's position of the exception
- **namespace** (`str`) – The namespace of the identifier
- **name** (`str`) – The name of the identifier

```
exception pybel.parser.exc.MissingNamespaceRegexWarning(line_number, line, position,
                                                       namespace, name)
```

Raised if reference not matching regex.

Initialize the namespace identifier warning.

Parameters

- **line_number** (`int`) – The line number of the line that caused the exception
- **line** (`str`) – The line that caused the exception
- **position** (`int`) – The line's position of the exception
- **namespace** (`str`) – The namespace of the identifier
- **name** (`str`) – The name of the identifier

```
exception pybel.parser.exc.AnnotationWarning(line_number, line, position, annotation,
                                             *args)
```

Base exception for annotation warnings.

Build an AnnotationWarning.

Parameters

- **line_number** (`int`) – The line number on which the warning occurred
- **line** (`str`) – The line on which the warning occurred
- **position** (`int`) – The position in the line that caused the warning
- **annotation** (`str`) – The annotation name that caused the warning

```
exception pybel.parser.exc.UndefinedAnnotationWarning(line_number, line, position,
                                                      annotation, *args)
```

Raised when an undefined annotation is used.

Build an AnnotationWarning.

Parameters

- **line_number** (*int*) – The line number on which the warning occurred
- **line** (*str*) – The line on which the warning occurred
- **position** (*int*) – The position in the line that caused the warning
- **annotation** (*str*) – The annotation name that caused the warning

```
exception pybel.parser.exc.MissingAnnotationKeyWarning(line_number, line, position,
                                                       annotation, *args)
```

Raised when trying to unset an annotation that is not set.

Build an AnnotationWarning.

Parameters

- **line_number** (*int*) – The line number on which the warning occurred
- **line** (*str*) – The line on which the warning occurred
- **position** (*int*) – The position in the line that caused the warning
- **annotation** (*str*) – The annotation name that caused the warning

```
exception pybel.parser.exc.AnnotationIdentifierWarning(line_number, line, position,
                                                       annotation, value)
```

Base exception for annotation:value pairs.

Build an AnnotationWarning.

Parameters

- **line_number** (*int*) – The line number on which the warning occurred
- **line** (*str*) – The line on which the warning occurred
- **position** (*int*) – The position in the line that caused the warning
- **annotation** (*str*) – The annotation name that caused the warning

```
exception pybel.parser.exc.IllegalAnnotationValueWarning(line_number, line, position,
                                                       annotation, value)
```

Raised when an annotation has a value that does not belong to the original set of valid annotation values.

Build an AnnotationWarning.

Parameters

- **line_number** (*int*) – The line number on which the warning occurred
- **line** (*str*) – The line on which the warning occurred
- **position** (*int*) – The position in the line that caused the warning
- **annotation** (*str*) – The annotation name that caused the warning

```
exception pybel.parser.exc.MissingAnnotationRegexWarning(line_number, line, position,
                                                       annotation, value)
```

Raised if annotation doesn't match regex.

Build an AnnotationWarning.

Parameters

- **line_number** (`int`) – The line number on which the warning occurred
- **line** (`str`) – The line on which the warning occurred
- **position** (`int`) – The position in the line that caused the warning
- **annotation** (`str`) – The annotation name that caused the warning

```
exception pybel.parser.exc.VersionFormatWarning(line_number, line, position, version_string)
```

Raised if the version string doesn't adhere to semantic versioning or YYYYMMDD format.

Initialize the BEL parser warning.

Parameters

- **line_number** – The line number on which this warning occurred
- **line** – The content of the line
- **position** – The position within the line where the warning occurred

```
exception pybel.parser.exc.MetadataException(line_number, line, position, *args)
```

Base exception for issues with document metadata.

Initialize the BEL parser warning.

Parameters

- **line_number** (`int`) – The line number on which this warning occurred
- **line** (`str`) – The content of the line
- **position** (`int`) – The position within the line where the warning occurred

```
exception pybel.parser.exc.MalformedMetadataException(line_number, line, position, *args)
```

Raised when an invalid metadata line is encountered.

Initialize the BEL parser warning.

Parameters

- **line_number** (`int`) – The line number on which this warning occurred
- **line** (`str`) – The content of the line
- **position** (`int`) – The position within the line where the warning occurred

```
exception pybel.parser.exc.InvalidMetadataException(line_number, line, position, key, value)
```

Raised when an incorrect document metadata key is used.

Hint: Valid document metadata keys are:

- Authors
- ContactInfo
- Copyright
- Description
- Disclaimer
- Licenses

- Name
 - Version
-

See also:

BEL specification on the [properties section](#)

Initialize the BEL parser warning.

Parameters

- **line_number** – The line number on which this warning occurred
- **line** – The content of the line
- **position** – The position within the line where the warning occurred

exception `pybel.parser.exc.MissingMetadataException(line_number, line, position, key)`

Raised when a BEL Script is missing critical metadata.

Initialize the BEL parser warning.

Parameters

- **line_number** – The line number on which this warning occurred
- **line** – The content of the line
- **position** – The position within the line where the warning occurred

static make(key)

Build an instance of this class with auto-filled dummy values.

Unlike normal classes, polymorphism on `__init__` can't be used for exceptions when pickling/unpickling.

exception `pybel.parser.exc.InvalidCitationLengthException(line_number, line, position, *args)`

Base exception raised when the format for a citation is wrong.

Initialize the BEL parser warning.

Parameters

- **line_number (int)** – The line number on which this warning occurred
- **line (str)** – The content of the line
- **position (int)** – The position within the line where the warning occurred

exception `pybel.parser.exc.CitationTooShortException(line_number, line, position, *args)`

Raised when a citation does not have the minimum of {type, name, reference}.

Initialize the BEL parser warning.

Parameters

- **line_number (int)** – The line number on which this warning occurred
- **line (str)** – The content of the line
- **position (int)** – The position within the line where the warning occurred

exception `pybel.parser.exc.CitationTooLongException(line_number, line, position, *args)`

Raised when a citation has more than the allowed entries, {type, name, reference, date, authors, comments}.

Initialize the BEL parser warning.

Parameters

- **line_number** (`int`) – The line number on which this warning occurred
- **line** (`str`) – The content of the line
- **position** (`int`) – The position within the line where the warning occurred

```
exception pybel.parser.exc.MissingCitationException(line_number, line, position,  
                                                 *args)
```

Raised when trying to parse a BEL statement, but no citation is currently set.

This might be due to a previous error in the formatting of a citation.

Though it's not a best practice, some BEL curators set other annotations before the citation. If this is the case in your BEL document, and you're *absolutely* sure that all UNSET statements are correctly written, you can use `citation_clearing=True` as a keyword argument in any of the IO functions in `pybel.from_lines()`, `pybel.from_url()`, or `pybel.from_path()`.

Initialize the BEL parser warning.

Parameters

- **line_number** (`int`) – The line number on which this warning occurred
- **line** (`str`) – The content of the line
- **position** (`int`) – The position within the line where the warning occurred

```
exception pybel.parser.exc.MissingSupportWarning(line_number, line, position, *args)
```

Raised when trying to parse a BEL statement, but no evidence is currently set.

All BEL statements must be qualified with evidence.

If your data is serialized from a database and provenance information is not readily accessible, consider referencing the publication for the database, or a url pointing to the data from either a programmatically or human-readable endpoint.

Initialize the BEL parser warning.

Parameters

- **line_number** (`int`) – The line number on which this warning occurred
- **line** (`str`) – The content of the line
- **position** (`int`) – The position within the line where the warning occurred

```
exception pybel.parser.exc.MissingAnnotationWarning(line_number, line, position, required_annotations)
```

Raised when trying to parse a BEL statement and a required annotation is not present.

Initialize the BEL parser warning.

Parameters

- **line_number** – The line number on which this warning occurred
- **line** – The content of the line
- **position** – The position within the line where the warning occurred

```
exception pybel.parser.exc.InvalidCitationType(line_number, line, position, citation_type)
```

Raised when a citation is set with an incorrect type.

Hint: Valid citation types include:

- Book
 - PubMed
 - Journal
 - Online Resource
 - URL
 - DOI
 - Other
-

See also:

OpenBEL wiki on [citations](#)

Initialize the BEL parser warning.

Parameters

- **line_number** – The line number on which this warning occurred
- **line** – The content of the line
- **position** – The position within the line where the warning occurred

exception `pybel.parser.exc.InvalidPubMedIdentifierWarning(line_number, line, position, reference)`

Raised when a citation is set whose type is PubMed but whose database identifier is not a valid integer.

Initialize the BEL parser warning.

Parameters

- **line_number** – The line number on which this warning occurred
- **line** – The content of the line
- **position** – The position within the line where the warning occurred

exception `pybel.parser.exc.MalformedTranslocationWarning(line_number, line, position, tokens)`

Raised when there is a translocation statement without location information.

Initialize the BEL parser warning.

Parameters

- **line_number** – The line number on which this warning occurred
- **line** – The content of the line
- **position** – The position within the line where the warning occurred

exception `pybel.parser.exc.PlaceholderAminoAcidWarning(line_number, line, position, code)`

Raised when an invalid amino acid code is given.

One example might be the usage of X, which is a colloquial signifier for a truncation in a given position. Text mining efforts for knowledge extraction make this mistake often. X might also signify a placeholder amino acid.

Initialize the BEL parser warning.

Parameters

- **line_number** – The line number on which this warning occurred
- **line** – The content of the line
- **position** – The position within the line where the warning occurred

exception `pybel.parser.exc.NestedRelationWarning`(*line_number*, *line*, *position*, **args*)

Raised when encountering a nested statement.

See our the docs for an explanation of why we explicitly do not support nested statements.

Initialize the BEL parser warning.

Parameters

- **line_number** (`int`) – The line number on which this warning occurred
- **line** (`str`) – The content of the line
- **position** (`int`) – The position within the line where the warning occurred

exception `pybel.parser.exc.InvalidEntity`(*line_number*, *line*, *position*, *namespace*, *name*)

Raised when using a non-entity name for a name.

Initialize the BEL parser warning.

Parameters

- **line_number** – The line number on which this warning occurred
- **line** – The content of the line
- **position** – The position within the line where the warning occurred

exception `pybel.parser.exc.InvalidFunctionSemantic`(*line_number*, *line*, *position*, *func*, *namespace*, *name*, *allowed_functions*)

Raised when an invalid function is used for a given node.

For example, an HGNC symbol for a protein-coding gene YFG cannot be referenced as an miRNA with `m(HGNC:YFG)`

Initialize the BEL parser warning.

Parameters

- **line_number** – The line number on which this warning occurred
- **line** – The content of the line
- **position** – The position within the line where the warning occurred

REFERENCES

If you find PyBEL useful for your work, please consider citing¹:

20.1 Related Publications

We have used PyBEL in several other projects and publications. Below is a sample:

- Domingo-Fernández, D., Mubeen, S., Marin-Llao, J., Hoyt, C. T., & Hofmann-Apitius, M. (2018). PathMe: Merging and exploring mechanistic pathway knowledge. bioRxiv, 451625.
- Domingo-Fernández, D., Hoyt, C. T., Alvarez, C. B., Marin-Llao, J., Hofmann-Apitius, M. (2018). ComPath: an ecosystem for exploring, analyzing, and curating mappings across pathway databases. Npj Systems Biology and Applications, 5(1), 3. <https://doi.org/10.1038/s41540-018-0078-8>
- Hoyt, C. T., et al. (2018). A systematic approach for identifying shared mechanisms in epilepsy and its comorbidities. Database, 2018(1). <https://doi.org/10.1093/database/bay050>
- Hoyt, C. T., et al. (2019). Re-curation and Rational Enrichment of Knowledge Graphs in Biological Expression Language. BioRxiv, 536409. <https://doi.org/10.1101/536409>
- Hoyt, C. T., Domingo-Fernández, D., & Hofmann-Apitius, M. (2018). BEL Commons: an environment for exploration and analysis of networks encoded in Biological Expression Language. Database, 2018(3), 1–11. <https://doi.org/10.1093/database/bay126>
- Ali, M., et al. (2019). BioKEEN: A library for learning and evaluating biological knowledge graph embeddings. Bioinformatics, btz117. <https://doi.org/10.1093/bioinformatics/btz117>

20.2 Software using PyBEL

- <https://github.com/cthoyt/bel-repository>
- <https://github.com/bio2bel>
- <https://github.com/sorgerlab/indra>
- <https://github.com/smartDataAnalytics/biokeen>

¹ Hoyt, C. T., et al. (2017). PyBEL: a Computational Framework for Biological Expression Language. Bioinformatics, 34(December), 1–2.

20.3 BEL Content

- <https://github.com/neurommsig/neurommsig-knowledge>
- <https://github.com/pharmacome/knowledge>

CHAPTER
TWENTYONE

ROADMAP

This project road map documents not only the PyBEL repository, but the PyBEL Tools and BEL Commons repositories as well as the Bio2BEL project.

21.1 PyBEL

- **Performance improvements**
 - Parallelization of parsing
 - On-the-fly validation with OLS or MIRIAM

21.2 Bio2BEL

- **Generation of new namespaces, equivalencies, and hierarchical knowledge (isA and partOf relations)**
 - FlyBase
 - InterPro (Done)
 - UniProt (Done)
 - Human Phenotype Ontology
 - Uber Anatomy Ontology
 - HGNC Gene Families (Done)
 - Enzyme Classification (Done)
- **Integration of knowledge sources**
 - ChEMBL
 - Comparative Toxicogenomics Database (Done)
 - BRENDA
 - MetaCyc
 - Protein complex definitions

21.3 Data2BEL

Integration of analytical pipelines to convert data to BEL

- LD Block Analysis
- Gene Co-expression Analysis
- Differential Gene Expression Analysis

21.4 PyBEL Tools

- **Biological Grammar**
 - Network motif identification
 - Stability analysis
 - **Prior knowledge comparision**
 - * Molecular activity annotation
 - * SNP Impact
- **Implementation of standard BEL Algorithms**
 - RCR
 - NPA
 - SST
- **Development of new algorithms**
 - Heat diffusion algorithms
 - Cart Before the Horse
 - Metapath analysis
 - Reasoning and inference rules
 - Subgraph Expansion application in NeuroMMSigDB
 - Chemical Enrichment in NeuroMMSigDB

21.5 BEL Commons

- Integration with BELIEF
- Integration with NeuroMMSigDB (Done)
- Import and export from NDEx

CHAPTER
TWENTYTWO

CURRENT ISSUES

22.1 Speed

Speed is still an issue, because documents above 100K lines still take a couple minutes to run. This issue is exacerbated by (optionally) logging output to the console, which can make it more than 3x or 4x as slow.

22.2 Namespaces

The default namespaces from OpenBEL do not follow a standard file format. They are similar to INI config files, but do not use consistent delimiters. Also, many of the namespaces don't respect that the delimiter should not be used in the namespace names. There are also lots of names with strange characters, which may have been caused by copying from a data source that had specific escape characters without proper care.

22.3 Testing

Testing was very difficult because the example documents on the OpenBEL website had many semantic errors, such as using names and annotation values that were not defined within their respective namespace and annotation definition files. They also contained syntax errors like naked names, which are not only syntactically incorrect, but lead to bad science; and improper usage of activities, like illegally nesting an activity within a composite statement.

CHAPTER
TWENTYTHREE

TECHNOLOGY

This page is meant to describe the development stack for PyBEL, and should be a useful introduction for contributors.

23.1 Versioning

PyBEL is versioned on GitHub so changes in its code can be tracked over time and to make use of the variety of software development plugins. Code is produced following the [Git Flow](#) philosophy, which means that new features are coded in branches off of the development branch and merged after they are triaged. Finally, develop is merged into master for releases. If there are bugs in releases that need to be fixed quickly, “hot fix” branches from master can be made, then merged back to master and develop after fixing the problem.

23.2 Testing in PyBEL

PyBEL is written with extensive unit testing and integration testing. Whenever possible, test- driven development is practiced. This means that new ideas for functions and features are encoded as blank classes/functions and directly writing tests for the desired output. After tests have been written that define how the code should work, the implementation can be written.

Test-driven development requires us to think about design before making quick and dirty implementations. This results in better code. Additionally, thorough testing suites make it possible to catch when changes break existing functionality.

Tests are written with the standard `unittest` library.

23.2.1 Unit Testing

Unit tests check that the functionality of the different parts of PyBEL work independently.

An example unit test can be found in `tests.test_parse_bel.TestAbundance.test_short_abundance`. It ensures that the parser is able to handle a given string describing the abundance of a chemical/other entity in BEL. It tests that the parser produces the correct output, that the BEL statement is converted to the correct internal representation. In this example, this is a tuple describing the abundance of oxygen atoms. Finally, it tests that this representation is added as a node in the underlying BEL graph with the appropriate attributes added.

23.2.2 Integration Testing

Integration tests are more high level, and ensure that the software accomplishes more complicated goals by using many components. An example integration test is found in `tests.test_import.TestImport.test_from_fileURL`. This test ensures that a BEL script can be read and results in a NetworkX object that contains all of the information described in the script

23.2.3 Tox

While IDEs like PyCharm provide excellent testing tools, they are not programmatic. `Tox` is python package that provides a CLI interface to run automated testing procedures (as well as other build functions, that aren't important to explain here). In PyBEL, it is used to run the unit tests in the `tests` folder with the `pytest` harness. It also runs `check-manifest`, builds the documentation with `sphinx`, and computes the code coverage of the tests. The entire procedure is defined in `tox.ini`. Tox also allows test to be done on many different versions of Python.

23.2.4 Continuous Integration

Continuous integration is a philosophy of automatically testing code as it changes. PyBEL makes use of the Travis CI server to perform testing because of its tight integration with GitHub. Travis automatically installs git hooks inside GitHub so it knows when a new commit is made. Upon each commit, Travis downloads the newest commit from GitHub and runs the tests configured in the `.travis.yml` file in the top level of the PyBEL repository. This file effectively instructs the Travis CI server to run Tox. It also allows for the modification of the environment variables. This is used in PyBEL to test many different versions of python.

23.2.5 Code Coverage

After building, Travis sends code coverage results to codecov.io. This site helps visualize untested code and track the improvement of testing coverage over time. It also integrates with GitHub to show which feature branches are inadequately tested. In development of PyBEL, inadequately tested code is not allowed to be merged into develop.

23.2.6 Versioning

PyBEL uses semantic versioning. In general, the project's version string will have a suffix `-dev` like in `0.3.4-dev` throughout the development cycle. After code is merged from feature branches to develop and it is time to deploy, this suffix is removed and develop branch is merged into master.

The version string appears in multiple places throughout the project, so `BumpVersion` is used to automate the updating of these version strings. See `.bumpversion.cfg` for more information.

23.3 Deployment

PyBEL is also distributed through PyPI (pronounced Py-Pee-Eye). Travis CI has a wonderful integration with PyPI, so any time a tag is made on the master branch (and also assuming the tests pass), a new distribution is packed and sent to PyPI. Refer to the "deploy" section at the bottom of the `.travis.yml` file for more information, or the Travis CI [PyPI deployment documentation](#). As a side note, Travis CI has an encryption tool so the password for the PyPI account can be displayed publicly on GitHub. Travis decrypts it before performing the upload to PyPI.

23.3.1 Steps

1. bumpversion release on development branch
2. Push to git
3. After tests pass, merge develop in to master
4. After tests pass, create a tag on GitHub with the same name as the version number (on master)
5. Travis will automatically deploy to PyPI after tests pass. After checking deployment has been successful, switch to develop and bumpversion patch

CHAPTER
TWENTYFOUR

INDICES AND TABLES

- genindex
- modindex
- search

BIBLIOGRAPHY

[Tarcă2009] Tarcă, A. L., *et al* (2009). A novel signaling pathway impact analysis. *Bioinformatics*, 25(1), 75–82.

PYTHON MODULE INDEX

p

pybel.constants, 103
pybel.dsl, 123
pybel.examples, 35
pybel.examples.braf_example, 36
pybel.examples.egf_example, 35
pybel.examples.sialic_acid_example, 36
pybel.examples.statin_example, 36
pybel.examples.tloc_example, 37
pybel.exceptions, 135
pybel.io, 57
pybel.io.bel_commons_client, 72
pybel.io.biobase_client, 73
pybel.io.cx, 62
pybel.io.extras, 78
pybel.io.fraunhofer_orientdb, 74
pybel.io.gpickle, 60
pybel.io.graphdati, 65
pybel.io.graphml, 77
pybel.io.hetionet, 59
pybel.io.hipathia, 69
pybel.io.indra, 66
pybel.io.jgif, 63
pybel.io.jupyter, 68
pybel.io.neo4j, 76
pybel.io.nodelink, 61
pybel.io.pynpa, 69
pybel.io.spia, 71
pybel.io.tsv, 71
pybel.io.umbrella_nodelink, 77
pybel.language, 109
pybel.manager.database_io, 76
pybel.manager.models, 85
pybel.parser.exc, 135
pybel.parser.modifiers, 122
pybel.parser.modifiers.fragment, 26
pybel.parser.modifiers.fusion, 27
pybel.parser.modifiers.gene_modification,
 23
pybel.parser.modifiers.gene_substitution,
 22
pybel.parser.modifiers.location, 32

pybel.parser.modifiers.protein_modification,
 24
pybel.parser.modifiers.protein_substitution,
 24
pybel.parser.modifiers.truncation, 25
pybel.parser.modifiers.variant, 21
pybel.struct, 11
pybel.struct.filters, 47
pybel.struct.grouping, 51
pybel.struct.mutation, 49
pybel.struct.pipeline.decorators, 55
pybel.struct.pipeline.exc, 56
pybel.struct.summary, 39

INDEX

Symbols

```
--add__() (pybel.BELGraph method), 12
__and__() (pybel.BELGraph method), 13
__iadd__() (pybel.BELGraph method), 13
__iand__() (pybel.BELGraph method), 13
--allow-naked-names
    pybel-compile command line option,
        95
--bel <bel>
    pybel-serialize command line
        option, 101
--connection <connection>
    pybel command line option, 95
    pybel-neo command line option, 100
--debug
    pybel-manage-examples command line
        option, 98
--disallow-nested
    pybel-compile command line option,
        95
--disallow-unqualified-translocations
    pybel-compile command line option,
        95
--edgelist <edgelist>
    pybel-serialize command line
        option, 101
--graphml <graphml>
    pybel-serialize command line
        option, 101
--gsea <gsea>
    pybel-serialize command line
        option, 101
--host <host>
    pybel-machine command line option,
        96
    pybel-post command line option, 100
--limit <limit>
    pybel-manage-edges-ls command line
        option, 97
--local
    pybel-machine command line option,
        96
--namespace-id <namespace_id>
    pybel-manage-namespaces-ls command
        line option, 99
--network-id <network_id>
    pybel-manage-networks-drop command
        line option, 99
--no-citation-clearing
    pybel-compile command line option,
        96
--no-identifier-validation
    pybel-compile command line option,
        95
--nodelink <nodelink>
    pybel-serialize command line
        option, 101
--offset <offset>
    pybel-manage-edges-ls command line
        option, 97
--password <password>
    pybel-neo command line option, 100
--required-annotations
    <required_annotations>
    pybel-compile command line option,
        96
--sif <sif>
    pybel-serialize command line
        option, 101
--skip-tqdm
    pybel-compile command line option,
        96
--tsv <tsv>
    pybel-serialize command line
        option, 101
--upgrade-urls
    pybel-compile command line option,
        96
--url <url>
    pybel-manage-namespaces-ls command
        line option, 99
--verbose
    pybel-compile command line option,
        96
```

```
--version
    pybel command line option, 95
--yes
    pybel-manage-drop command line
        option, 97
    pybel-manage-networks-drop command
        line option, 99
-c
    pybel command line option, 95
-i
    pybel-manage-namespaces-ls command
        line option, 99
-n
    pybel-manage-networks-drop command
        line option, 99
-r
    pybel-compile command line option,
        96
-u
    pybel-manage-namespaces-ls command
        line option, 99
-v
    pybel-compile command line option,
        96
    pybel-manage-examples command line
        option, 98
-y
    pybel-manage-networks-drop command
        line option, 99
```

A

Abundance (*class in pybel.dsl*), 124
ABUNDANCE (*in module pybel.constants*), 105
abundance_labels (*in module pybel.language*), 110
abundance_list (*pybel.parser.parse_bel.BELParser attribute*), 113
abundance_sbo_mapping (*in module pybel.language*), 110
activity (*pybel.parser.parse_bel.BELParser attribute*), 112
activity_labels (*in module pybel.language*), 110
activity_mapping (*in module pybel.language*), 110
add_activates() (*pybel.BELGraph method*), 19
add_association() (*pybel.BELGraph method*), 18
add_binds() (*pybel.BELGraph method*), 18
add_causes_no_change() (*pybel.BELGraph method*), 19
add_correlation() (*pybel.BELGraph method*), 18
add_decreases() (*pybel.BELGraph method*), 18
add_dephosphorylates() (*pybel.BELGraph method*), 19
add_directly_decreases() (*pybel.BELGraph method*), 18
add_directly_dephosphorylates() (*pybel.BELGraph method*), 19
add_directly_increases() (*pybel.BELGraph method*), 18
add_directly_phosphorylates() (*pybel.BELGraph method*), 19
add_equivalence() (*pybel.BELGraph method*), 17
add_has_product() (*pybel.BELGraph method*), 17
add_has_reactant() (*pybel.BELGraph method*), 17
add_has_variant() (*pybel.BELGraph method*), 17
add_increases() (*pybel.BELGraph method*), 18
add_inhibits() (*pybel.BELGraph method*), 19
add_is_a() (*pybel.BELGraph method*), 17
add_negative_correlation() (*pybel.BELGraph method*), 19
add_no_correlation() (*pybel.BELGraph method*), 18
add_node_from_data() (*pybel.BELGraph method*), 20
add_orthology() (*pybel.BELGraph method*), 17
add_part_of() (*pybel.BELGraph method*), 17
add_phosphorylates() (*pybel.BELGraph method*), 19
add_positive_correlation() (*pybel.BELGraph method*), 18
add_qualified_edge() (*pybel.BELGraph method*), 17
add_reaction() (*pybel.BELGraph method*), 20
add_regulates() (*pybel.BELGraph method*), 18
add_transcription() (*pybel.BELGraph method*), 16
add_translation() (*pybel.BELGraph method*), 17
add_unqualified_edge() (*pybel.BELGraph method*), 16
AGENTS
 pybel-machine command line option, 97
ANALOGOUS_TO (*in module pybel.constants*), 107
annotation_list() (*pybel.BELGraph property*), 15
annotation_pattern() (*pybel.BELGraph property*), 15
annotation_to_local (*pybel.parser.parse_metadata.MetadataParser attribute*), 116
annotation_to_pattern (*pybel.parser.parse_metadata.MetadataParser attribute*), 116
annotation_to_term (*pybel.parser.parse_metadata.MetadataParser attribute*), 115
annotation_url() (*pybel.BELGraph property*), 15
annotation_url_dict (*pybel.parser.parse_metadata.MetadataParser*

attribute), 116
AnnotationIdentifierWarning, 138
ANNOTATIONS (in module pybel.constants), 106
AnnotationWarning, 137
append() (pybel.Pipeline method), 54
as_bel() (pybel.manager.models.Network method), 87
as_bel() (pybel.manager.models.Node method), 88
ASSOCIATION (in module pybel.constants), 107
Author (class in pybel.manager.models), 88
author (pybel.manager.models.Namespace attribute), 86
authors (pybel.manager.models.Network attribute), 87
authors() (pybel.BELGraph property), 14

B

Base (class in pybel.manager.models), 85
base (pybel.manager.BaseManager attribute), 79
BaseAbundance (class in pybel.dsl), 123
BaseEntity (class in pybel.dsl), 123
BaseManager (class in pybel.manager), 79
bel (pybel.manager.models.Edge attribute), 90
bel (pybel.manager.models.Node attribute), 88
bel_contains() (pybel.manager.models.Node class method), 88
BEL_DEFAULT_NAMESPACE (in module pybel.constants), 103
BELGraph (class in pybel), 12
belns_encodings (in module pybel.constants), 109
BELParser (class in pybel.parser.parse_bel), 111
BELParserWarning, 135
BELSyntaxError, 135
bind() (pybel.manager.BaseManager method), 79
BINDS (in module pybel.constants), 106
biological_process (pybel.parser.parse_bel.BELParser attribute), 112
BiologicalProcess (class in pybel.dsl), 124
BIOMARKER_FOR (in module pybel.constants), 107
BIOPROCESS (in module pybel.constants), 105
blob (pybel.manager.models.Network attribute), 87

C

calculate_error_by_annotation() (in module pybel.struct.summary), 39
calculate_incorrect_name_dict() (in module pybel.struct.summary), 39
CAUSAL_DECREASE_RELATIONS (in module pybel.constants), 107
CAUSAL_INCREASE_RELATIONS (in module pybel.constants), 107
CAUSAL_POLAR_RELATIONS (in module pybel.constants), 107
CAUSAL_RELATIONS (in module pybel.constants), 107
CAUSES_NO_CHANGE (in module pybel.constants), 106
CentralDogma (class in pybel.dsl), 125
check_function_semantics() (pybel.parser.parse_bel.BELParser method), 113
Citation (class in pybel.manager.models), 89
CITATION (in module pybel.constants), 105
CITATION_AUTHORS (in module pybel.constants), 103
CITATION_DATE (in module pybel.constants), 103
CITATION_DB (in module pybel.constants), 103
CITATION_DB_NAME (in module pybel.constants), 103
CITATION_FIRST_AUTHOR (in module pybel.constants), 104
CITATION_IDENTIFIER (in module pybel.constants), 103
citation_is_set() (pybel.parser.parse_control.ControlParser property), 118
CITATION_ISSUE (in module pybel.constants), 103
CITATION_JOURNAL (in module pybel.constants), 103
CITATION_LAST_AUTHOR (in module pybel.constants), 104
CITATION_PAGES (in module pybel.constants), 104
CITATION_TYPES (in module pybel.constants), 103
CITATION_VOLUME (in module pybel.constants), 103
CitationTooLongException, 140
CitationTooShortException, 140
clear() (pybel.parser.parse_bel.BELParser method), 113
clear() (pybel.parser.parse_control.ControlParser method), 120
clear_citation() (pybel.parser.parse_control.ControlParser method), 120
compartment_mapping (in module pybel.language), 110
COMPLEX (in module pybel.constants), 105
complex_singleton (pybel.parser.parse_bel.BELParser attribute), 112
ComplexAbundance (class in pybel.dsl), 132
COMPOSITE (in module pybel.constants), 105
composite_abundance (pybel.parser.parse_bel.BELParser attribute), 112
CompositeAbundance (class in pybel.dsl), 132
CONCEPT (in module pybel.constants), 104
ConceptParser (class in pybel.parser.parse_concept), 120
contact (pybel.manager.models.Namespace attribute), 86
contact (pybel.manager.models.Network attribute), 87
contact() (pybel.BELGraph property), 14
ControlParser (class in pybel.parser.parse_control), 118

copyright (*pybel.manager.models.Network attribute*), 87
 copyright () (*pybel.BELGraph property*), 14
 CORRELATION (*in module pybel.constants*), 107
 CORRELATIVE_RELATIONS (*in module pybel.constants*), 107
 count_annotations() (*in module pybel.struct.summary*), 42
 count_edges() (*pybel.manager.QueryManager method*), 82
 count_error_types() (*in module pybel.struct.summary*), 39
 count_functions() (*in module pybel.struct.summary*), 39
 count_naked_names() (*in module pybel.struct.summary*), 39
 count_names_by_namespace() (*in module pybel.struct.summary*), 40
 count_namespaces() (*in module pybel.struct.summary*), 40
 count_networks() (*pybel.manager.NetworkManager method*), 81
 count_nodes() (*pybel.manager.QueryManager method*), 82
 count_pathologies() (*in module pybel.struct.summary*), 41
 count_relations() (*in module pybel.struct.summary*), 42
 count_variants() (*in module pybel.struct.summary*), 40
 create_all() (*pybel.manager.BaseManager method*), 79
 created (*pybel.manager.models.Namespace attribute*), 86
 curie() (*pybel.language.Entity property*), 110

D

data (*pybel.manager.models.Edge attribute*), 91
 data (*pybel.manager.models.Node attribute*), 88
 date (*pybel.manager.models.Citation attribute*), 89
 db (*pybel.manager.models.Citation attribute*), 89
 db_id (*pybel.manager.models.Citation attribute*), 89
 DECREASES (*in module pybel.constants*), 106
 default_namespace (*pybel.parser.parse_metadata.MetadataParser attribute*), 115
 DEFAULT_SERVICE_URL (*in module pybel.constants*), 109
 defined_annotation_keywords() (*pybel.BELGraph property*), 16
 defined_namespace_keywords() (*pybel.BELGraph property*), 15

degradation (*pybel.parser.parse_bel.BELParser attribute*), 113
 DeprecationMappingError, 56
 DESCRIPTION (*in module pybel.constants*), 104
 description (*pybel.manager.models.Namespace attribute*), 85
 description (*pybel.manager.models.Network attribute*), 87
 description() (*pybel.BELGraph property*), 14
 description_contains() (*pybel.manager.models.Network class method*), 87
 DIRECT_CAUSAL_RELATIONS (*in module pybel.constants*), 107
 DIRECTLY_DECREASES (*in module pybel.constants*), 106
 DIRECTLY_INCREASES (*in module pybel.constants*), 106
 DIRTY (*in module pybel.constants*), 105
 disclaimer (*pybel.manager.models.Network attribute*), 87
 disclaimer() (*pybel.BELGraph property*), 14
 DOCUMENT_KEYS (*in module pybel.constants*), 108
 document_metadata (*pybel.parser.parse_metadata.MetadataParser attribute*), 116
 domain (*pybel.manager.models.Namespace attribute*), 85
 drop_all() (*pybel.manager.BaseManager method*), 79
 drop_network() (*pybel.manager.NetworkManager method*), 81
 drop_network_by_id() (*pybel.manager.NetworkManager method*), 81
 drop_networks() (*pybel.manager.NetworkManager method*), 81
 dump() (*in module pybel*), 57
 dump() (*pybel.Pipeline method*), 54
 dumps() (*pybel.Pipeline method*), 54

E

Edge (*class in pybel.manager.models*), 90
 edge_to_bel() (*pybel.BELGraph static method*), 20
 encoding (*pybel.manager.models.NamespaceEntry attribute*), 86
 ensure_node() (*pybel.parser.parse_bel.BELParser method*), 114
 Entity (*class in pybel.dsl*), 123
 Entity (*class in pybel.language*), 109
 EnumeratedFusionRange (*class in pybel.dsl*), 132
 EQUIVALENT_TO (*in module pybel.constants*), 107
 Evidence (*class in pybel.manager.models*), 89
 EVIDENCE (*in module pybel.constants*), 106

`extend()` (*pybel.Pipeline method*), 54

F

`first_id` (*pybel.manager.models.Citation attribute*), 89

`Fragment` (*class in pybel.dsl*), 130

`FRAGMENT` (*in module pybel.constants*), 105

`FRAGMENT_DESCRIPTION` (*in module pybel.constants*), 109

`FRAGMENT_MISSING` (*in module pybel.constants*), 109

`FRAGMENT_START` (*in module pybel.constants*), 108

`FRAGMENT_STOP` (*in module pybel.constants*), 108

`from_bel_commons()` (*in module pybel*), 72

`from_bel_script()` (*in module pybel*), 59

`from_bel_script_url()` (*in module pybel*), 59

`from_biodati()` (*in module pybel*), 74

`from_biopax()` (*in module pybel*), 68

`from_bytes()` (*in module pybel*), 60

`from_cbn_jgif()` (*in module pybel*), 64

`from_cbn_jgif_file()` (*in module pybel*), 65

`from_cx()` (*in module pybel*), 62

`from_cx_file()` (*in module pybel*), 62

`from_cx_gz()` (*in module pybel*), 63

`from_cx_jsons()` (*in module pybel*), 62

`from_database()` (*in module pybel*), 76

`from_fraunhofer_orientdb()` (*in module pybel*), 74

`from_functions()` (*pybel.Pipeline static method*), 53

`from_graphdati()` (*in module pybel*), 65

`from_graphdati_file()` (*in module pybel*), 66

`from_graphdati_gz()` (*in module pybel*), 66

`from_graphdati_jsons()` (*in module pybel*), 66

`from_hetionet_file()` (*in module pybel*), 59

`from_hetionet_gz()` (*in module pybel*), 59

`from_hetionet_json()` (*in module pybel*), 59

`from_hipathia_dfs()` (*in module pybel*), 71

`from_hipathia_paths()` (*in module pybel*), 71

`from_indra_statements()` (*in module pybel*), 67

`from_indra_statements_json()` (*in module pybel*), 67

`from_indra_statements_json_file()` (*in module pybel*), 67

`from_jgif()` (*in module pybel*), 63

`from_jgif_file()` (*in module pybel*), 64

`from_jgif_gz()` (*in module pybel*), 64

`from_jgif_jsons()` (*in module pybel*), 63

`from_json()` (*pybel.Pipeline static method*), 54

`from_nodelink()` (*in module pybel*), 61

`from_nodelink_file()` (*in module pybel*), 61

`from_nodelink_gz()` (*in module pybel*), 61

`from_nodelink_jsons()` (*in module pybel*), 61

`from_pickle()` (*in module pybel*), 60

`FUNCTION` (*in module pybel.constants*), 104

`fusion` (*pybel.parser.parse_bel.BELParser attribute*), 112

`FusionBase` (*class in pybel.dsl*), 130

`FusionRangeBase` (*class in pybel.dsl*), 132

G

`Gene` (*class in pybel.dsl*), 125

`GENE` (*in module pybel.constants*), 105

`gene` (*pybel.parser.parse_bel.BELParser attribute*), 112

`GeneFusion` (*class in pybel.dsl*), 130

`GeneModification` (*class in pybel.dsl*), 128

`general_abundance` (*pybel.parser.parse_bel.BELParser attribute*), 112

`get_annotation_values()` (*in module pybel.struct.summary*), 42

`get_annotation_values_by_annotation()` (*in module pybel.struct.summary*), 41

`get_annotations()` (*in module pybel.struct.summary*), 42

`get_annotations()` (*pybel.parser.parse_bel.BELParser method*), 113

`get_annotations()` (*pybel.parser.parse_control.ControlParser method*), 120

`get_annotations_json()` (*pybel.manager.models.Edge method*), 91

`get_authors()` (*pybel.BELGraph method*), 16

`get_cache_connection()` (*in module pybel.constants*), 103

`get_citation()` (*pybel.parser.parse_control.ControlParser method*), 120

`get_edge_annotations()` (*pybel.BELGraph method*), 20

`get_edge_citation()` (*pybel.BELGraph method*), 20

`get_edge_evidence()` (*pybel.BELGraph method*), 20

`get_edges_with_annotation()` (*pybel.manager.QueryManager method*), 83

`get_edges_with_citation()` (*pybel.manager.QueryManager method*), 82

`get_edges_with_citations()` (*pybel.manager.QueryManager method*), 83

`get_equivalent_nodes()` (*pybel.BELGraph method*), 20

`get_functions()` (*in module pybel.struct.summary*), 39

`get_graph_by_id()` (*pybel.manager.NetworkManager method*), 82

get_graph_by_ids()
 `bel.manager.NetworkManager`
 82

get_graph_by_most_recent()
 `bel.manager.NetworkManager`
 82

get_graph_by_name_version()
 `bel.manager.NetworkManager`
 81

get_graphs_by_ids()
 `bel.manager.NetworkManager`
 82

get_hetionet() (*in module pybel*), 59

get_metaedge_to_key() (*in module* `pybel.struct.summary`), 42

get_missing_required_annotations()
 (`pybel.parser.parse_control.ControlParser`
 method), 120

get_most_recent_network_by_name()
 (`pybel.manager.NetworkManager` method), 81

get_naked_names() (*in module* `pybel.struct.summary`), 39

get_names() (*in module* `pybel.struct.summary`), 40

get_names_by_namespace() (*in module* `pybel.struct.summary`), 40

get_namespaces() (*in module* `pybel.struct.summary`), 40

get_network_by_id()
 `bel.manager.NetworkManager`
 82

get_network_by_name_version()
 `bel.manager.NetworkManager`
 81

get_network_versions()
 `bel.manager.NetworkManager`
 81

get_networks_by_ids()
 `bel.manager.NetworkManager`
 82

get_networks_by_name()
 `bel.manager.NetworkManager`
 81

get_pubmed_identifiers() (*in module* `pybel.struct.summary`), 41

get_subgraphs_by_annotation() (*in module* `pybel.struct.grouping`), 51

get_subgraphs_by_citation() (*in module* `pybel.struct.grouping`), 51

get_syntax_errors() (*in module* `pybel.struct.summary`), 39

get_term_to_encodings()
 `bel.manager.models.Namespace`
 86

get_top_hubs() (*in module* `pybel.struct.summary`), 41

get_top_pathologies() (*in module* `bel.struct.summary`), 41

get_transformation() (*in module* `bel.struct.pipeline.decorators`), 55

get_unused_annotations() (*in module* `pybel.struct.summary`), 42

get_unused_list_annotation_values() (*in module* `pybel.struct.summary`), 42

get_unused_namespaces() (*in module* `pybel.struct.summary`), 40

GMOD (*in module* `pybel.constants`), 105

gmod (`pybel.parser.parse_bel.BELParser` attribute), 112

gmod_mappings (*in module* `pybel.language`), 110

gmod_namespace (*in module* `pybel.language`), 110

GMOD_ORDER (*in module* `pybel.constants`), 109

GRAPH_METADATA (*in module* `pybel.constants`), 108

GSUB_POSITION (*in module* `pybel.constants`), 109

GSUB_REFERENCE (*in module* `pybel.constants`), 109

GSUB_VARIANT (*in module* `pybel.constants`), 109

H

handle_annotation_key()
 (`bel.parser.parse_control.ControlParser`
 method), 119

handle_annotation_list()
 (`bel.parser.parse_metadata.MetadataParser`
 method), 116

handle_annotation_pattern()
 (`bel.parser.parse_metadata.MetadataParser`
 method), 117

handle_annotations_url()
 (`bel.parser.parse_metadata.MetadataParser`
 method), 116

handle_document()
 (`bel.parser.parse_metadata.MetadataParser`
 method), 116

handle_has_components()
 (`bel.parser.parse_bel.BELParser` method), 113

handle_has_members()
 (`bel.parser.parse_bel.BELParser` method), 113

handle_identifier_fqualified()
 (`bel.parser.parse_concept.ConceptParser`
 method), 121

handle_identifier_qualified()
 (`bel.parser.parse_concept.ConceptParser`
 method), 121

handle_inverse_unqualified_relation()
 (`pybel.parser.parse_bel.BELParser` method), 114

handle_namespace_default()
 (`pybel.parser.parse_concept.ConceptParser`

method), 121		
handle_namespace_invalid() (py- bel.parser.parse_concept.ConceptParser method), 121		handle_unset_statement_group() (py- bel.parser.parse_control.ControlParser method), 119
handle_namespace_lenient() (py- bel.parser.parse_concept.ConceptParser static method), 121		has_annotation() (py- bel.parser.parse_control.ControlParser method), 118
handle_namespace_pattern() (py- bel.parser.parse_metadata.MetadataParser method), 116		has_annotation() (py- bel.parser.parse_metadata.MetadataParser method), 117
handle_namespace_url() (py- bel.parser.parse_metadata.MetadataParser method), 116		has_edge_citation() (pybel.BELGraph method), 20
handle_nested_relation() (py- bel.parser.parse_bel.BELParser method), 113		has_edge_evidence() (pybel.BELGraph method), 20
handle_set_citation() (py- bel.parser.parse_control.ControlParser method), 119		has_enumerated_annotation() (py- bel.parser.parse_control.ControlParser method), 118
handle_set_command() (py- bel.parser.parse_control.ControlParser method), 119		has_enumerated_annotation() (py- bel.parser.parse_metadata.MetadataParser method), 117
handle_set_command_list() (py- bel.parser.parse_control.ControlParser method), 119		has_enumerated_namespace() (py- bel.parser.parse_concept.ConceptParser method), 121
handle_set_evidence() (py- bel.parser.parse_control.ControlParser method), 119		has_enumerated_namespace() (py- bel.parser.parse_metadata.MetadataParser method), 117
handle_set_statement_group() (py- bel.parser.parse_control.ControlParser method), 119		has_fusion (pybel.manager.models.Node attribute), 88
handle_term() (pybel.parser.parse_bel.BELParser method), 113		has_local_annotation() (py- bel.parser.parse_control.ControlParser method), 118
handle_translocation_illegal() (py- bel.parser.parse_bel.BELParser method), 114		has_local_annotation() (py- bel.parser.parse_metadata.MetadataParser method), 117
handle_unqualified_relation() (py- bel.parser.parse_bel.BELParser method), 113		has_member (pybel.parser.parse_bel.BELParser attribute), 113
handle_unset_all() (py- bel.parser.parse_control.ControlParser method), 120		has_name_in() (pybel.manager.models.Author class method), 89
handle_unset_citation() (py- bel.parser.parse_control.ControlParser method), 119		has_name_version() (py- bel.manager.NetworkManager method), 81
handle_unset_command() (py- bel.parser.parse_control.ControlParser method), 120		has_namespace() (py- bel.parser.parse_metadata.MetadataParser method), 117
handle_unset_evidence() (py- bel.parser.parse_control.ControlParser method), 119		HAS_PRODUCT (in module pybel.constants), 106
handle_unset_list() (py- bel.parser.parse_control.ControlParser method), 120		HAS_REACTANT (in module pybel.constants), 106
		has_regex_annotation() (py- bel.parser.parse_control.ControlParser method), 118
		has_regex_annotation() (py- bel.parser.parse_metadata.MetadataParser method), 117
		has_regex_namespace() (py- bel.parser.parse_concept.ConceptParser method), 121

```

has_regex_namespace() (pybel.parser.parse_metadata.MetadataParser
method), 117
HAS_VARIANT (in module pybel.constants), 106
HASH (in module pybel.constants), 106
Hgvs (class in pybel.dsl), 129
HGVS (in module pybel.constants), 104
HgvsReference (class in pybel.dsl), 129
HgvsUnspecified (class in pybel.dsl), 129

I
id_in() (pybel.manager.models.Network class
method), 87
IDENTIFIER (in module pybel.constants), 104
identifier (pybel.manager.models.NamespaceEntry
attribute), 86
identifier () (pybel.language.Entity property), 110
IllegalAnnotationValueWarning, 138
in_place_transformation() (in module pybel.struct.pipeline.decorators), 55
InconsistentDefinitionError, 135
INCREASES (in module pybel.constants), 106
INDIRECT_CAUSAL_RELATIONS (in module pybel.constants), 107
insert_into_graph() (pybel.manager.models.Edge method), 91
intersection() (pybel.Pipeline static method), 55
InvalidCitationLengthException, 140
InvalidCitationType, 141
InvalidEntity, 143
InvalidFunctionSemantic, 143
InvalidMetadataException, 139
InvalidPubMedIdentifierWarning, 142
INVERSE_DOCUMENT_KEYS (in module pybel.constants), 108
IS_A (in module pybel.constants), 107
is_enriched() (pybel.manager.models.Citation
property), 89
is_pubmed() (pybel.manager.models.Citation
property), 89
is_subject (pybel.manager.models.Property
attribute), 90
is_variant (pybel.manager.models.Node attribute),
88
issue (pybel.manager.models.Citation attribute), 89
iter_annotation_value_pairs() (in module
pybel.struct.summary), 41
iter_annotation_values() (in module pybel.struct.summary), 41
iter_equivalent_nodes() (pybel.BELGraph
method), 20
iterate_pubmed_identifiers() (in module pybel.struct.summary), 41

```

J
journal (pybel.manager.models.Citation attribute), 89

K
keyword (pybel.manager.models.Namespace attribute), 85
KIND (in module pybel.constants), 104

L
LABEL (in module pybel.constants), 104
last_id (pybel.manager.models.Citation attribute), 89
left_full_join() (in module pybel.struct), 45
left_outer_join() (in module pybel.struct), 45
license (pybel.manager.models.Namespace attribute), 86
license () (pybel.BELGraph property), 14
licenses (pybel.manager.models.Network attribute), 87
LINE (in module pybel.constants), 106
list_networks() (pybel.manager.NetworkManager
method), 81
list_recent_networks() (pybel.manager.NetworkManager
method), 81
ListAbundance (class in pybel.dsl), 124
load() (in module pybel), 57
load() (pybel.Pipeline static method), 55
loads() (pybel.Pipeline static method), 55
location (pybel.parser.parse_bel.BELParser
attribute), 112

M
make () (pybel.parser.exc.MissingMetadataException
static method), 140
MalformedMetadataException, 139
MalformedTranslocationWarning, 142
Manager (class in pybel.manager), 79
manager (pybel.parser.parse_metadata.MetadataParser
attribute), 115
md5 (pybel.manager.models.Edge attribute), 90
MEMBERS (in module pybel.constants), 104
METADATA_AUTHORS (in module pybel.constants), 108
METADATA_CONTACT (in module pybel.constants), 108
METADATA_COPYRIGHT (in module pybel.constants),
108
METADATA_DESCRIPTION (in module pybel.constants), 108
METADATA_DISCLAIMER (in module pybel.constants),
108
METADATA_INSERT_KEYS (in module pybel.constants), 108
METADATA_LICENSES (in module pybel.constants),
108

METADATA_NAME (*in module pybel.constants*), 108
 METADATA_PROJECT (*in module pybel.constants*), 108
 METADATA_VERSION (*in module pybel.constants*), 108
 MetadataException, 139
 MetadataParser (class in *pybel.parser.parse_metadata*), 115
 MetaValueError, 56
 MicroRna (class in *pybel.dsl*), 126
 miriam_id (*pybel.manager.models.Namespace* attribute), 85
 MIRNA (*in module pybel.constants*), 105
 mirna (*pybel.parser.parse_bel.BELParser* attribute), 112
 MissingAnnotationKeyWarning, 138
 MissingAnnotationRegexWarning, 138
 MissingAnnotationWarning, 141
 MissingCitationException, 141
 MissingDefaultNameWarning, 136
 MissingFusionRange (class in *pybel.dsl*), 132
 MissingMetadataException, 140
 MissingNamespaceNameWarning, 137
 MissingNamespaceRegexWarning, 137
 MissingPipelineFunctionError, 56
 MissingSupportWarning, 141
 MissingUniverseError, 56
 Modification (class in *pybel.manager.models*), 87
 modifier (*pybel.manager.models.Property* attribute), 90
 module
 pybel.constants, 103
 pybel.dsl, 123
 pybel.examples, 35
 pybel.examples.braf_example, 36
 pybel.examples.egf_example, 35
 pybel.examples.sialic_acid_example, 36
 pybel.examples.statin_example, 36
 pybel.examples.tloc_example, 37
 pybel.exceptions, 135
 pybel.io, 57
 pybel.io.bel_commons_client, 72
 pybel.io.bioclient, 73
 pybel.io.cx, 62
 pybel.io.extras, 78
 pybel.io.fraunhofer_orientdb, 74
 pybel.io.gpickle, 60
 pybel.io.graphdati, 65
 pybel.io.graphml, 77
 pybel.io.hetionet, 59
 pybel.io.hipathia, 69
 pybel.io.indra, 66
 pybel.io.jgif, 63
 pybel.io.jupyter, 68
 pybel.io.neo4j, 76
 pybel.io.nodelink, 61
 pybel.io.pynpa, 69
 pybel.io.spia, 71
 pybel.io.tsv, 71
 pybel.io.umbrella_nodelink, 77
 pybel.language, 109
 pybel.manager.database_io, 76
 pybel.manager.models, 85
 pybel.parser.exc, 135
 pybel.parser.modifiers, 122
 pybel.parser.modifiers.fragment, 26
 pybel.parser.modifiers.fusion, 27
 pybel.parser.modifiers.gene_modification, 23
 pybel.parser.modifiers.gene_substitution, 22
 pybel.parser.modifiers.location, 32
 pybel.parser.modifiers.protein_modification, 24
 pybel.parser.modifiers.protein_substitution, 24
 pybel.parser.modifiers.truncation, 25
 pybel.parser.modifiers.variant, 21
 pybel.struct, 11
 pybel.struct.filters, 47
 pybel.struct.grouping, 51
 pybel.struct.mutation, 49
 pybel.struct.pipeline.decorators, 55
 pybel.struct.pipeline.exc, 56
 pybel.struct.summary, 39
 molecular_activity (*pybel.parser.parse_bel.BELParser* attribute), 112

N

NakedNameWarning, 136
 NAME (*in module pybel.constants*), 104
 name (*pybel.manager.models.Namespace* attribute), 85
 name (*pybel.manager.models.NamespaceEntry* attribute), 86
 name (*pybel.manager.models.Network* attribute), 87
 name () (*pybel.BELGraph* property), 13
 name () (*pybel.language.Entity* property), 110
 name_contains () (*pybel.manager.models.Author* class method), 88
 name_contains () (*pybel.manager.models.NamespaceEntry* class method), 86
 name_contains () (*pybel.manager.models.Network* class method), 87
 Namespace (class in *pybel.manager.models*), 85
 NAMESPACE (*in module pybel.constants*), 104
 namespace () (*pybel.language.Entity* property), 110

NAMESPACE_DOMAIN_TYPES (in module `pybel.constants`), 103
namespace_pattern() (`pybel.BELGraph` property), 15
namespace_to_pattern (`pybel.parser.parse_metadata.MetadataParser` attribute), 115
namespace_to_term_to_encoding (`pybel.parser.parse_metadata.MetadataParser` attribute), 115
namespace_url() (`pybel.BELGraph` property), 15
namespace_url_dict (`pybel.parser.parse_metadata.MetadataParser` attribute), 116
NamespaceEntry (class in `pybel.manager.models`), 86
NamespaceIdentifierWarning, 136
NameWarning, 136
NEGATIVE_CORRELATION (in module `pybel.constants`), 107
NestedRelationWarning, 143
Network (class in `pybel.manager.models`), 86
NetworkManager (class in `pybel.manager`), 81
NO_CORRELATION (in module `pybel.constants`), 107
Node (class in `pybel.manager.models`), 88
node_has_namespace() (`pybel.BELGraph` method), 20
node_to_bel() (`pybel.BELGraph` static method), 20
number_of_authors() (`pybel.BELGraph` method), 16
number_of_citations() (`pybel.BELGraph` method), 16
number_of_warnings() (`pybel.BELGraph` method), 16

O

OBJECT (in module `pybel.constants`), 106
obo() (`pybel.language.Entity` property), 110
ORTHOLOGOUS (in module `pybel.constants`), 107

P

pages (`pybel.manager.models.Citation` attribute), 89
parse_lines() (in module `pybel.io.line_utils`), 114
parse_result_to_dsl() (in module `pybel.tokens`), 133
PART_OF (in module `pybel.constants`), 107
PARTNER_3P (in module `pybel.constants`), 104
PARTNER_5P (in module `pybel.constants`), 104
PATH
 pybel-compile command line option, 96
path
 pybel-insert command line option, 96
 pybel-neo command line option, 100
 pybel-post command line option, 100

pybel-serialize command line option, 101
pybel-summarize command line option, 101
pybel-warnings command line option, 102
path() (`pybel.BELGraph` property), 13
Pathology (class in `pybel.dsl`), 124
PATHOLOGY (in module `pybel.constants`), 105
pathology (`pybel.parser.parse_bel.BELParser` attribute), 112
pattern (`pybel.manager.models.Namespace` attribute), 85
Pipeline (class in `pybel`), 53
PipelineNameError, 56
PlaceholderAminoAcidWarning, 142
PMOD (in module `pybel.constants`), 105
pmod (`pybel.parser.parse_bel.BELParser` attribute), 112
PMOD_CODE (in module `pybel.constants`), 109
pmod_legacy_labels (in module `pybel.language`), 110
pmod_mappings (in module `pybel.language`), 110
pmod_namespace (in module `pybel.language`), 110
PMOD_ORDER (in module `pybel.constants`), 109
PMOD_POSITION (in module `pybel.constants`), 109
POLAR_RELATIONS (in module `pybel.constants`), 108
Population (class in `pybel.dsl`), 125
POPULATION (in module `pybel.constants`), 105
position (`pybel.manager.models.Modification` attribute), 88
POSITIVE_CORRELATION (in module `pybel.constants`), 107
post_jgif() (in module `pybel`), 64
PRODUCTS (in module `pybel.constants`), 104
PROGNOSTIC_BIOMARKER_FOR (in module `pybel.constants`), 107
Property (class in `pybel.manager.models`), 90
Protein (class in `pybel.dsl`), 127
PROTEIN (in module `pybel.constants`), 105
protein (`pybel.parser.parse_bel.BELParser` attribute), 112
ProteinFusion (class in `pybel.dsl`), 131
ProteinModification (class in `pybel.dsl`), 128
ProteinSubstitution (class in `pybel.dsl`), 129
PSUB_POSITION (in module `pybel.constants`), 109
PSUB_REFERENCE (in module `pybel.constants`), 109
PSUB_VARIANT (in module `pybel.constants`), 109
pybel command line option
 --connection <connection>, 95
 --version, 95
 -c, 95
pybel.constants
 module, 103
pybel.dsl

```

        module, 123
pybel.examples
    module, 35
pybel.examples.braf_example
    module, 36
pybel.examples.braf_example_graph (in
    module pybel.examples.braf_example), 36
pybel.examples.egf_example
    module, 35
pybel.examples.egf_graph (in module py-
    bel.examples.egf_example), 36
pybel.examples.ras_tloc_graph (in module
    pybel.examples.tloc_example), 37
pybel.examples.sialic_acid_example
    module, 36
pybel.examples.sialic_acid_graph (in mod-
    ule pybel.examples.sialic_acid_example), 36
pybel.examples.statin_example
    module, 36
pybel.examples.statin_graph (in module py-
    bel.examples.statin_example), 36
pybel.examples.tloc_example
    module, 37
pybel.exceptions
    module, 135
pybel.io
    module, 57
pybel.io.bel_commons_client
    module, 72
pybel.io.bioclient
    module, 73
pybel.io.cx
    module, 62
pybel.io.extras
    module, 78
pybel.io.fraunhofer_orientdb
    module, 74
pybel.io.gpickle
    module, 60
pybel.io.graphdati
    module, 65
pybel.io.graphml
    module, 77
pybel.io.hetionet
    module, 59
pybel.io.hipathia
    module, 69
pybel.io.indra
    module, 66
pybel.io.jgif
    module, 63
pybel.io.jupyter
    module, 68
pybel.io.neo4j
    module, 76
pybel.io.nodelink
    module, 61
pybel.io.pynpa
    module, 69
pybel.io.spia
    module, 71
pybel.io.tsv
    module, 71
pybel.io.umbrella_nodelink
    module, 77
pybel.language
    module, 109
pybel.manager.database_io
    module, 76
pybel.manager.models
    module, 85
pybel.parser.exc
    module, 135
pybel.parser.modifiers
    module, 122
pybel.parser.modifiers.fragment
    module, 26
pybel.parser.modifiers.fusion
    module, 27
pybel.parser.modifiers.gene_modification
    module, 23
pybel.parser.modifiers.gene_substitution
    module, 22
pybel.parser.modifiers.location
    module, 32
pybel.parser.modifiers.protein_modification
    module, 24
pybel.parser.modifiers.protein_substitution
    module, 24
pybel.parser.modifiers.truncation
    module, 25
pybel.parser.modifiers.variant
    module, 21
pybel.struct
    module, 11
pybel.struct.filters
    module, 47
pybel.struct.grouping
    module, 51
pybel.struct.mutation
    module, 49
pybel.struct.pipeline.decorators
    module, 55
pybel.struct.pipeline.exc
    module, 56
pybel.struct.summary
    module, 39

```

```

PYBEL_EDGE_ALL_KEYS (in module pybel.constants),           -i, 99
    106          -u, 99
PYBEL_EDGE_DATA_KEYS  (in module pybel.constants), 106      pybel-manage-networks-drop command
    bel.constants), 106          line option
PYBEL_EDGE_METADATA_KEYS  (in module pybel.constants), 106   --network-id <network_id>, 99
    bel.constants), 106          --yes, 99
PYBEL_NODE_DATA_KEYS  (in module pybel.constants), 105       -n, 99
    bel.constants), 105          -y, 99
PYBEL_NODE_FUNCTIONS  (in module pybel.constants), 105      pybel-neo command line option
    bel.constants), 105          --connection <connection>, 100
PYBEL_VARIANT_KINDS  (in module pybel.constants), 105        --password <password>, 100
    105          path, 100
pybel_version() (pybel.BELGraph property), 16      pybel-post command line option
pybel-compile command line option                         --host <host>, 100
    --allow-naked-names, 95      path, 100
    --disallow-nested, 95       pybel-serialize command line option
    --disallow-unqualified-translocations, 95            --bel <bel>, 101
    95          --edgelist <edgelist>, 101
    --no-citation-clearing, 96          --graphml <graphml>, 101
    --no-identifier-validation, 95          --gsea <gsea>, 101
    --required-annotations          --nodelink <nodelink>, 101
    <required_annotations>, 96          --sif <sif>, 101
    --skip-tqdm, 96          --tsv <tsv>, 101
    --upgrade-urls, 96          path, 101
    --verbose, 96       pybel-summarize command line option
    -r, 96          path, 101
    -v, 96          pybel-warnings command line option
    PATH, 96          path, 102
pybel-insert command line option                         PyBELWarning, 135
    path, 96
pybel-machine command line option                      Q
    --host <host>, 96
    --local, 96
    AGENTS, 97
pybel-manage-drop command line option                  query_citations() (pybel.manager.QueryManager
    --yes, 97          method), 83
pybel-manage-edges-ls command line option             query_edges()      (pybel.manager.QueryManager
    option          method), 83
    --limit <limit>, 97
    --offset <offset>, 97
pybel-manage-examples command line                   query_edges_by_pubmed_identifiers() (pybel.
    option          manager.QueryManager method), 84
    --debug, 98
    -v, 98
pybel-manage-namespaces-drop command                query_induction() (pybel.manager.QueryManager
    line option          method), 84
    URL, 98
pybel-manage-namespaces-insert command              query_neighbors() (pybel.manager.QueryManager
    line option          method), 84
    URL, 98
pybel-manage-namespaces-ls command                  query_nodes()      (pybel.manager.QueryManager
    line option          method), 82
    --namespace-id <namespace_id>, 99
    --url <url>, 99
query_singleton_edges_from_network() (pybel.
    manager.NetworkManager method), 81
query_url (pybel.manager.models.Namespace at-
    tribute), 86
QueryManager (class in pybel.manager), 82
R
raise_for_invalid_annotation_value() (pybel.parser.parse_control.ControlParser
    method), 118

```

raise_for_missing_citation() (*pybel.parser.parse_control.ControlParser method*), 119

raise_for_missing_name() (*pybel.parser.parse_concept.ConceptParser method*), 121

raise_for_missing_namespace() (*pybel.parser.parse_concept.ConceptParser method*), 121

raise_for_redefined_annotation() (*pybel.parser.parse_metadata.MetadataParser method*), 116

raise_for_redefined_namespace() (*pybel.parser.parse_metadata.MetadataParser method*), 116

raise_for_undefined_annotation() (*pybel.parser.parse_control.ControlParser method*), 118

raise_for_version() (*pybel.parser.parse_metadata.MetadataParser method*), 117

RANGE_3P (*in module pybel.constants*), 104

RANGE_5P (*in module pybel.constants*), 104

rate_limit (*pybel.parser.parse_bel.BELParser attribute*), 113

RATE_LIMITING_STEP_OF (*in module pybel.constants*), 107

REACTANTS (*in module pybel.constants*), 104

reactants (*pybel.parser.parse_bel.BELParser attribute*), 113

Reaction (*class in pybel.dsl*), 133

REACTION (*in module pybel.constants*), 105

RedefinedAnnotationError, 136

RedefinedNamespaceError, 136

REGULATES (*in module pybel.constants*), 106

RELATION (*in module pybel.constants*), 105

relative_key (*pybel.manager.models.Property attribute*), 90

REQUIRED_METADATA (*in module pybel.constants*), 108

residue (*pybel.manager.models.Modification attribute*), 88

rev_abundance_labels (*in module pybel.constants*), 105

Rna (*class in pybel.dsl*), 126

RNA (*in module pybel.constants*), 105

rna (*pybel.parser.parse_bel.BELParser attribute*), 112

RnaFusion (*class in pybel.dsl*), 131

run() (*pybel.Pipeline method*), 54

S

search_edges_with_bel() (*pybel.manager.QueryManager method*), 83

search_edges_with_evidence() (*pybel.manager.QueryManager method*), 83

side() (*pybel.manager.models.Property property*), 90

species (*pybel.manager.models.Namespace attribute*), 85

store_bel() (*pybel.manager.models.Network method*), 87

SUBJECT (*in module pybel.constants*), 106

SUBPROCESS_OF (*in module pybel.constants*), 107

subprocess_of (*pybel.parser.parse_bel.BELParser attribute*), 113

summarize() (*pybel.BELGraph method*), 21

summary_dict() (*pybel.BELGraph method*), 21

summary_str() (*pybel.BELGraph method*), 21

T

text (*pybel.manager.models.Evidence attribute*), 90

title (*pybel.manager.models.Citation attribute*), 89

to_bel_commons() (*in module pybel*), 72

to_bel_script() (*in module pybel*), 59

to_biodati() (*in module pybel*), 73

to_bytes() (*in module pybel*), 60

to_csv() (*in module pybel*), 78

to_cx() (*in module pybel*), 62

to_cx_file() (*in module pybel*), 62

to_cx_gz() (*in module pybel*), 63

to_cx_jsons() (*in module pybel*), 62

to_database() (*in module pybel*), 76

to_edgelist() (*in module pybel*), 72

to_graphdati() (*in module pybel*), 65

to_graphdati_file() (*in module pybel*), 65

to_graphdati_gz() (*in module pybel*), 66

to_graphdati_jsonl() (*in module pybel*), 66

to_graphdati_jsonl_gz() (*in module pybel*), 66

to_graphdati_jsons() (*in module pybel*), 66

to_graphml() (*in module pybel*), 77

to_gsea() (*in module pybel*), 78

to_hipathia() (*in module pybel*), 70

to_hipathia_dfs() (*in module pybel*), 70

to_indra_statements() (*in module pybel*), 68

to_indra_statements_json() (*in module pybel*), 68

to_indra_statements_json_file() (*in module pybel*), 68

to_jgif() (*in module pybel*), 63

to_jgif_file() (*in module pybel*), 64

to_jgif_gz() (*in module pybel*), 64

to_jgif_jsons() (*in module pybel*), 64

to_json() (*pybel.manager.models.Citation method*), 89

to_json() (*pybel.manager.models.Edge method*), 91

to_json() (*pybel.manager.models.Evidence method*), 90

to_json() (*pybel.manager.models.Modification method*), 88
 to_json() (*pybel.manager.models.Namespace method*), 86
 to_json() (*pybel.manager.models.NamespaceEntry method*), 86
 to_json() (*pybel.manager.models.Network method*), 87
 to_json() (*pybel.manager.models.Node method*), 88
 to_json() (*pybel.manager.models.Property method*), 90
 to_json() (*pybel.Pipeline method*), 54
 to_jupyter() (*in module pybel.io.jupyter*), 68
 to_neo4j() (*in module pybel*), 76
 to_nodelink() (*in module pybel*), 61
 to_nodelink_file() (*in module pybel*), 61
 to_nodelink_gz() (*in module pybel*), 61
 to_nodelink_jsons() (*in module pybel*), 61
 to_npa_dfs() (*in module pybel*), 69
 to_npa_directory() (*in module pybel*), 69
 to_pickle() (*in module pybel*), 60
 to_sif() (*in module pybel*), 78
 to_spia_dfs() (*in module pybel*), 71
 to_spia_excel() (*in module pybel*), 71
 to_spia_tsvs() (*in module pybel*), 71
 to_tsv() (*in module pybel*), 71
 to_umbrella_nodelink() (*in module pybel*), 77
 to_umbrella_nodelink_file() (*in module pybel*), 77
 to_umbrella_nodelink_gz() (*in module pybel*), 77
 Transcribable (*class in pybel.dsl*), 126
 transcribed (*pybel.parser.parse_bel.BELParser attribute*), 113
 TRANSCRIBED_TO (*in module pybel.constants*), 106
 transformation() (*in module pybel.struct.pipeline.decorators*), 55
 translated (*pybel.parser.parse_bel.BELParser attribute*), 113
 TRANSLATED_TO (*in module pybel.constants*), 106
 translocation (*pybel.parser.parse_bel.BELParser attribute*), 112
 TRUNCATION_POSITION (*in module pybel.constants*), 109
 TWO_WAY_RELATIONS (*in module pybel.constants*), 108
 type (*pybel.manager.models.Modification attribute*), 87
 type (*pybel.manager.models.Node attribute*), 88

U

uncachable_namespaces (*pybel.parser.parse_metadata.MetadataParser attribute*), 115

UndefinedAnnotationWarning, 137

UndefinedNamespaceWarning, 136
 uni_in_place_transformation() (*in module pybel.struct.pipeline.decorators*), 55
 uni_transformation() (*in module pybel.struct.pipeline.decorators*), 55
 union() (*in module pybel.struct*), 45
 union() (*pybel.Pipeline static method*), 55
 UNQUALIFIED_EDGES (*in module pybel.constants*), 108
 uploaded (*pybel.manager.models.Namespace attribute*), 85
 URL
 pybel-manage-namespaces-drop command line option, 98
 pybel-manage-namespaces-insert command line option, 98
 url (*pybel.manager.models.Namespace attribute*), 85

V

validate_unset_command() (*pybel.parser.parse_control.ControlParser method*), 119
 Variant (*class in pybel.dsl*), 128
 VARIANTS (*in module pybel.constants*), 104
 variantString (*pybel.manager.models.Modification attribute*), 88
 version (*pybel.manager.models.Namespace attribute*), 85
 version (*pybel.manager.models.Network attribute*), 87
 version() (*pybel.BELGraph property*), 13
 VersionFormatWarning, 139
 volume (*pybel.manager.models.Citation attribute*), 89

W

warnings() (*pybel.BELGraph property*), 16

X

XREFS (*in module pybel.constants*), 104