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# **NWB Specification Language**

*Release v1.2.0a*

**Aug 15, 2017**



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## NWB Specification Language

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Version: v1.2.0a Aug 15, 2017<sup>1</sup>

### Introduction

In order to support the formal and verifiable specification of neurodata file formats, NWB-N defines and uses the NWB specification language. The specification language is defined in YAML (or optionally JSON) and defines formal structures for describing the organization of complex data using basic concepts, e.g., Groups, Datasets, Attributes, and Links. A specification typically consists of a declaration of a namespace and a set of schema specifications. Data publishers can use the specification language to extend the format in order to store types of data not supported by the NWB core format ([Section 1.2](#)).

### Extensions

As mentioned, extensions to the core format are specified via custom user namespaces. Each namespace must have a unique name (i.e, must be different from NWB). The schema of new neurodata\_types (groups, datasets etc.) are then specified in seperate schema specification files. While it is possible to define multiple namespaces in the same file, most commonly, each new namespace will be defined in a separate file with corresponding schema specifications being stored in one ore more additional YAML (or JSON) files. One or more namespaces can be used simultaneously, so that multiple extensions can be used at the same time while avoiding potential name and type collisions between extensions (as well as extensions and the NWB core spec).

The specification of namespaces is described in detail next in [Section 1.3](#) and the specification of schema specifications is described in [Section 1.4](#) and subsequent sections.

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**Tip:** The `form` package as part of the PyNWB Python API provides dedicated data structures and utilities that support programmatic generation of extensions via Python programs, compared to writing YAML (or JSON) extension documents by hand. One main advantage of using PyNWB is that it is easier to use and maintain. E.g., using PyNWB

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<sup>1</sup> The version number given here is for the specification language and is independent of the version number for the NWB format. The date after the version number is the last modification date of this document.

helps ensure compliance of the generated specification files with the current specification language and the Python programs can often easily be just rerun to generate updated versions of extension files (with little to no changes to the program itself).

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**Tip:** The `nwb-schema` repo includes tools to generate Sphinx documentation from format specifications. In particular the tool `utils/init_sphinx_extension_doc.py` provides functionality to setup documentation for a format or extension defined by a namespace (similar to the documentation for NWB core namespace at <http://nwb-schema.readthedocs.io/en/latest/>). Use `python init_sphinx_extension_doc.py --help` to view the list of options for generating the docs. The repo also includes the tool `utils/generate_format_docs.py` which is used for generating actual reStructuredText files and figures from YAML/JSON specification sources.

---

## Namespaces

Namespaces are used to define a collections of specifications, to enable users to develop extensions in their own namespace and, hence, to avoid name/type collisions. Namespaces are defined in separate YAML files. The specification of a namespace looks as follows:

```
namespaces:
- doc: NWB namespace
  name: NWB
  full_name: NWB core
  version: 1.2.0
  date: 2017-04-25 18:05:00
  author:
  - Keith Godfrey
  - Jeff Teeters
  - Oliver Ruebel
  - Andrew Tritt
  contact:
  - keithg@alleninstitute.org
  - jteeters@berkeley.edu
  - oruebel@lbl.gov
  - ajtritt@lbl.gov
  schema:
  - source: nwb.base.yaml
    neurodata_types: null
  - ...
```

The top-level key must be `namespaces`. The value of `namespaces` is a list with the specification of one (or more) namespaces.

### Namespace declaration keys

#### **doc**

Text description of the namespace.

#### **name**

Unique name used to refer to the namespace

**full\_name**

Optional string with extended full name for the namespace.

**version**

Version string for the namespace

**date**

Date the namespace has been last modified or released. Formatting is %Y-%m-%d %H:%M:%S, e.g, 2017-04-25 17:14:13

**author**

List of strings with the names of the authors of the namespace.

**contact**

List of strings with the contact information for the authors. Ordering of the contacts should match the ordering of the authors.

**schema**

List of the schema to be included in this namespace. The specification looks as follows:

```
- source: .base.yaml
- source: nwb.ephys.yaml
  neurodata_types: ElectricalSeries
- namespace: core
  neurodata_types: Interface
```

- `source` describes the relative path to the YAML (or JSON) files with the schema specifications
- `namespace` describes a named reference to another namespace. In contrast to `source`, this is a reference by name to a known namespace (i.e., the namespace is resolved during the build and must point to an already existing namespace). This mechanism is used to allow, e.g., extension of a core namespace (here the NWB core namespace) without requiring hard paths to the files describing the core namespace.
- `neurodata_types` then is an optional list of strings indicating which `neurodata_types` should be included from the given specification source or namespace. The default is `neurodata_types: null` indicating that all `neurodata_types` should be included.

**Attention:** As with any language, we can only use what is defined. This means that similar to include or import statements in programming languages, e.g., Python, the `source` and `namespace` keys must be in order of use. E.g., `nwb.ephys.yaml` defines `ElectricalSeries` which inherits from `Timeseries` that is defined in `nwb.base.yaml`. This means that we have to list `nwb.base.yaml` before `nwb.ephys.yaml` since otherwise `Timeseries` would not be defined when `nwb.ephys.yaml` is trying to use it.

## Schema specification

The schema specification defines the groups, datasets and relationship that make up the format. Schema specifications are stored in dict `spec` and consist of a list of Group specifications. Schemas may be distributed across multiple YAML files to improve readability and to support logical organization of types. This is the main part of the format specification. It is described in the following sections.

```
specs:  
- ...
```

---

**Note:** Schema specifications are agnostic to namespaces, i.e., a schema (or type) becomes part of a namespace by including it in the namespace as part of the `schema` description of the namespace. Hence, the same schema can be reused across namespaces.

---

## Groups

Groups are specified as part of the top-level list or via lists stored in the key `groups`. The specification of a group is described in YAML as follows:

```
# Group specification  
- name: Optional fixed name for the group. A group must either have a unique_  
  ↳ neurodata_type or a unique, fixed name.  
  default_name: Default name for the group  
  doc: Required description of the group  
  neurodata_type_def: Optional new neurodata_type for the group  
  neurodata_type_inc: Optional neurodata_type the group should inherit from  
  quantity: Optional quantity identifier for the group (default=1).  
  linkable: Boolean indicating whether the group is linkable (default=True)  
  attributes: Optional list of attribute specifications describing the attributes_  
  ↳ of the group  
  datasets: Optional list of dataset specifications describing the datasets_  
  ↳ contained in the group  
  links: Optional list of link specification describing the links contained in the_  
  ↳ group  
  groups: Optional list of group specifications describing the sub-groups contained_  
  ↳ in the group
```

The key/value pairs that make up a group specification are described in more detail next in [Section 1.5.1](#).

### Group specification keys

#### **name**

String with the optional fixed name for the group.

---

**Note:** Every group must have either a unique fixed name or a unique `neurodata_type` determined by (`neurodata_type_def` and `neurodata_type_inc`) to enable the unique identification of groups when stored on disk.

---



**default\_name**

Default name of the group.

---

**Note:** Only one of either `name` or `default_name` (or neither) should be specified as the fixed name given by `name` would always overwrite the behavior of `default_name`.

---

**doc**

The value of the group specification `doc` key is a string describing the group. The `doc` key is required.

---

**Note:** In earlier versions (before version 1.2a) this key was called `description`

---

**neurodata\_type\_inc and neurodata\_type\_def**

The concept of a `neurodata_type` is similar to the concept of `Class` in object-oriented programming. A `neurodata_type` is a unique identifier for a specific type of group (or dataset) in a specification. By assigning a `neurodata_type` to a group (or dataset) enables others to reuse that type by inclusion or inheritance (*Note:* only groups (or datasets) with a specified type can be reused).

- ``neurodata_type_def``: This key is used to define (i.e, create) a new `neurodata_type` and to assign that type to the current group (or dataset).
- ``neurodata_type_inc``: The value of the `neurodata_type_inc` key describes the base type of a group (or dataset). The value must be an existing type.

Both ``neurodata_type_def`` and ``neurodata_type_inc`` are optional keys. To enable the unique identification, every group (and dataset) must either have a fixed name and/or a unique `neurodata_type`. This means, any group (or dataset) with a variable name must have a unique `neurodata_type`.

The `neurodata_type` is determined by the value of the `neurodata_type_def` key or if no new type is defined then the value of `neurodata_type_inc` is used to determine type. Or in other words, the `neurodata_type` is determined by the last type in the ancestry (i.e, inheritance hierarchy) of an object.

**Reusing existing neurodata\_types**

The combination of ``neurodata_type_inc`` and ``neurodata_type_def`` provides an easy-to-use mechanism for reuse of type specifications via inheritance (i.e., merge and extension of specifications) and inclusion (i.e, embedding of an existing type as a component, such as a subgroup, of a new specification). Here an overview of all relevant cases:

<code>neurodata_type_inc</code>	<code>neurodata_type_def</code>	Description
not set	not set	define a standard dataset or group without a type
not set	set	create a new <code>neurodata_type</code> from scratch
set	not set	include (reuse) <code>neurodata_type</code> without creating a new one (include)
set	set	merge/extend <code>neurodata_type</code> and create a new type (inheritance/merge)

**Example: Reuse by inheritance**

```
# Abbreviated YAML specification
- neurodata_type_def: Series
  datasets:
```

```

- name: A

- neurodata_type_def: MySeries
  neurodata_type_inc: Series
  datasets:
  - name: B
    
```

The result of this is that `MySeries` inherits dataset A from `Series` and adds its own dataset B, i.e., if we resolve the inheritance, then the above is equivalent to:

```

# Result:
- neurodata_type_def: MySeries
  datasets:
  - name: A
  - name: B
    
```

**Example: Reuse by inclusion**

```

# Abbreviated YAML specification
- neurodata_type_def: Series
  datasets:
  - name: A

- neurodata_type_def: MySeries
  groups:
  - neurodata_type_inc: Series
    
```

The result of this is that `MySeries` now includes a group of type `Series`, i.e., the above is equivalent to:

```

- neurodata_type_def: MySeries
  groups:
  - neurodata_type_inc: Series
    datasets:
    - name: A
    
```

---

**Note:** The keys ``neurodata_type_def`` and ``neurodata_type_inc`` were introduced in version 1.2a to simplify the concepts of inclusion and merging of specifications and replaced the keys ``include`` and ``merge`` (and ``merge+``).

---

**quantity**

The `quantity` describes how often the corresponding group (or dataset) can appear. The `quantity` indicates both minimum and maximum number of instances. Hence, if the minimum number of instances is 0 then the group (or dataset) is optional and otherwise it is required.

value	minimum quantity	maximum quantity	Comment
<code>`zero_or_more` or `*`</code>	0	unlimited	Zero or more instances
<code>`one_or_more` or `+`</code>	1	unlimited	One or more instances
<code>`zero_or_one` or `?`</code>	0	1	Zero or one instances
<code>`1`, `2`, `3`, ...</code>	n	n	Exactly n instances

---

**Note:** The `quantity` key was added in version 1.2a of the specification language as a replacement of the ``quantity_flag`` that was used to encode quantity information via a regular expression as part of the main

---

key of the group.

---

### **linkable**

Boolean describing whether the this group can be linked.

### **attributes**

List of attribute specifications describing the attributes of the group. See Section [\required](#) for details.

```
attributes:  
- ...
```

### **links**

List of link specifications describing all links to be stored as part of this group.

```
links:  
- doc: Link to target type  
  name: link name  
  target_type: type of target  
- ...
```

### **datasets**

List of dataset specifications describing all datasets to be stored as part of this group.

```
datasets:  
- name: data1  
  doc: My data 1  
  type: number  
  quantity: 'zero_or_one'  
- name: data2  
  doc: My data 2  
  type: text  
  attributes:  
    - ...  
- ...
```

### **groups**

List of group specifications describing all groups to be stored as part of this group

```
groups:  
- name: group1  
  quantity: 'zero_or_one'  
- ...
```

## `\_required`

**Attention:** TODO: The `\_required` key has been removed in version 1.2.x and later. An improved version will be added again in later version of the specification language.

## Attributes

Attributes are specified as part of lists stored in the key `attributes` as part of the specifications of groups and datasets. Attributes are typically used to further characterize or store metadata about the group, dataset, or link they are associated with. Similar to datasets, attributes can define arbitrary n-dimensional arrays, but are typically used to store smaller data. The specification of an attributes is described in YAML as follows:

```
...
attributes:
- name: Required string describing the name of the attribute
  doc: Required string with the description of the attribute
  dtype: Required string describing the data type of the attribute
  dims: Optional list describing the names of the dimensions of the data array stored_
  ↳by the attribute (default=None)
  shape: Optional list describing the allowed shape(s) of the data array stored by_
  ↳the attribute (default=None)
  required: Optional boolean indicating whether the attribute is required_
  ↳(default=True)
  value: Optional constant, fixed value for the attribute.
  default_value: Optional default value for variable-valued attributes. Only one of_
  ↳value or default_value should be set.
-
```

## Attribute specification keys

### `name`

String with the name for the attribute. The `name` key is required and must specify a unique attribute on the current parent object (e.g., group or dataset)

### `doc`

`doc` specifies the documentation string for the attribute and should describe the purpose and use of the attribute data. The `doc` key is required.

### `dtype`

String specifying the data type of the attribute. Allowable values are:

- `float` – indicates a floating point number
- `int` – indicates an integer
- `uint` – unsigned integer
- `number` – indicates either a floating point or an integer

- `text` – a text string

For all of the above types (except number and text), a default size (in bits) can be specified by appending the size to the type, e.g., `int32`. If “!” is appended to the default size, e.g. “`float64!`”, then the default size is also the required minimum size.

**Attention:**

- **TODO** Check that the list of allowable dtypes is complete
- **TODO** Check that the behavior described for type bit lengths is current

### `dims`

Optional key describing the names of the dimensions of the array stored as value of the attribute. If the attribute stores an array, `dims` specifies the list of dimensions. If no `dims` is given, then attribute stores a scalar value.

In case there is only one option for naming the dimensions, the key defines a single list of strings:

```
...
dims:
- dim1
- dim2
```

In case that the attribute may have different forms, this will be a list of lists:

```
...
dims:
- - num_times
- - num_times
- num_channels
```

Each entry in the list defines an identifier/name of the corresponding dimension of the array data.

### `shape`

Optional key describing the shape of the array stored as the value of the attribute. The description of `shape` must match the description of dimensions in so far as if we name two dimensions in `dims` than we must also specify the `shape` for two dimensions. We may specify `null` in case that the length of a dimension is not restricted. E.g.:

```
...
shape:
- null
- 3
```

Similar to `dims` `shape` may also be a list of lists in case that the attribute may have multiple valid shape options, e.g.:

```
...
shape:
- - 5
- - null
- 5
```

### **required**

Optional boolean key describing whether the attribute is required. Default value is True.

### **value**

Optional key specifying a fixed, constant value for the attribute. Default value is None, i.e., the attribute has a variable value to be determined by the user (or API) in accordance with the current data.

### **default\_value**

Optional key specifying a default value for attributes that allow user-defined values. The default value is used in case that the user does not specify a specific value for the attribute.

---

**Note:** Only one of either `value` or `default_value` should be specified (or neither) but never both at the same time, as `value` would always overwrite the `default_value`.

---

## Links

The link specification is used to specify links to other groups or datasets. In HDF5 it is recommended that links be stored as soft links. The link specification is a dictionary with the following form:

```
links:
- doc: Link to target type
  name: link name
  target_type: type of target
```

### Link specification keys

#### **target\_type**

`target_type` specifies the key for a group in the top level structure of a namespace. It is used to indicate that the link must be to an instance of that structure.

#### **doc**

`doc` specifies the documentation string for the link and should describe the purpose and use of the linked data. The `doc` key is required.

#### **name**

Optional key specifying the `name` of the link.

## Datasets

Datasets are specified as part of lists stored in the key `datasets` as part of group specifications. The specification of a datasets is described in YAML as follows:

```
- datasets:
  - name: fixed name of the dataset
    default_name: default name of the dataset
    doc: Required description of the dataset
    neurodata_type_def: Optional new neurodata_type for the group
    neurodata_type_inc: Optional neurodata_type the group should inherit from
    quantity: Optional quantity identifier for the group (default=1).
    linkable: Boolean indicating whether the group is linkable (default=True)
    dtype: Required string describing the data type of the dataset
    dims: Optional list describing the names of the dimensions of the dataset
    shape: Optional list describing the shape (or possibel shapes) of the dataset
    attributes: Optional list of attribute specifications describing the attributes_
    ↪of the group
```

The specification of datasets looks quite similar to attributes and groups. Similar to attributes, datasets describe the storage of arbitrary n-dimensional array data. However, in contrast to attributes, datasets are not associated with a specific parent group or dataset object but are (similar to groups) primary data objects (and as such typically manage larger data than attributes). The key/value pairs that make up a dataset specification are described in more detail next in Section [Section 1.8.1](#).

### Dataset specification keys

#### `name`

String with the optional fixed name for the dataset

---

**Note:** Every dataset must have either a unique fixed name or a unique `neurodata_type` to enable the unique identification of datasets when stored on disk.

---

#### `default_name`

Default name of the group.

---

**Note:** Only one of either `name` or `default_name` (or neither) should be specified as the fixed name given by `name` would always overwrite the behavior of `default_name`.

---

#### `doc`

The value of the dataset specification `doc` key is a string describing the dataset. The `doc` key is required.

---

**Note:** In earlier versions (before version 1.2a) this key was called `description`

---

### `neurodata_type_inc` and `neurodata_type_def`

Same as for groups. See [Section 1.5.1.4](#) for details.

### `quantity`

Same as for groups. See [Section 1.5.1.5](#) for details.

### `linkable`

Boolean describing whether the this group can be linked.

### `dtype`

String describing the data type of the dataset. Same as for attributes. See [Section 1.6.1.3](#) for details.

### `shape`

List describing the shape of the dataset. Same as for attributes. See [Section 1.6.1.5](#) for details.

### `dims`

List describing the names of the dimensions of the dataset. Same as for attributes. See [Section 1.6.1.4](#) for details.

### `attributes`

List of attribute specifications describing the attributes of the group. See [Section \\\_required](#) for details.

```
attributes:  
- ...
```

## Relationships

---

**Note:** Future versions will add explicit concepts for modeling of relationships, to replace the implicit relationships encoded via shared dimension descriptions and implicit references in datasets in previous versions of the specification language.

---



### Version 1.2a (April, 2017)

#### Summary

- **Simplify reuse of neurodata\_types:**

- Added new key: ``neurodata_type_def`` and `````neurodata_type_inc`` (which in combination replace the keys ``neurodata_type``, ``include`` and ``merge``). See below for details.
- Removed key: ``include``
- Removed key: ``merge``
- Removed key: ``merge+``
- Removed key: ``neurodata_type`` (replaced by `neurodata_type_inc`` and `neurodata_type_def``)
- Removed ``_properties`` key. The primary use of the key is to define abstract specifications. However, as format specifications don't implement functions but define a layout of objects, any spec (even if marked abstract) could still be instantiated and used in practice without limitations. Also, in the current instantiation of NWB-N this concept is only used for the ``Interface`` type and it is unclear why a user should not be able to use it. As such this concept was removed.
- To improve compliance of NWB-N inheritance mechanism with established object-oriented design concepts, the option of restricting the use of subclasses in place of parent classes was removed. A subclass is always also a valid instance of a parent class. This also improves consistency with the NWB-N principle of a minimal specification that allows users to add custom data. This change affects the ``allow_subclasses`` key of links and the `subclasses`` option of the removed ``include`` key.

- **Improve readability and avoid collision of keys by replacing values encoded in keys with dedicated key/value pairs:**

- **Explicit encoding of names and types:**

- \* Added ``name`` key
- \* Removed `<...>` name identifier (replaced by empty ``name`` key)
- \* Added ``groups`` key (previously groups were indicated by “/” as part of object’s key)
- \* Added ``datasets`` key (previously datasets were indicated by missing “/” as part of the object’s key)
- \* Added ``links`` key (previously this was a key on the group and dataset specification). The concept of links is with this now a first-class type (rather than being part of the group and dataset specs).
- \* Removed `link` key on datasets as this functionality is now fully implemented by the `links` key on groups.
- \* Removed / flag in keys to identify groups (replaced by ``groups`` and ``datasets`` keys)

– **Explicit encoding of quantities:**

- \* Added new key ``quantity`` (which replaces the ``quantity_flag``). See below for details.
- \* Removed ``quantity_flag`` as part of keys
- \* Removed `Exclude_in`` key. The key is currently not used in the NWB core spec. This feature is superseded by the ability to overwrite the ``quantity`` key as part of the reuse of ``neurodata_types``
- Removed ``\_description`` key. The key is no longer need because name conflicts with datasets and groups are no longer possible since the name is now explicitly encoded in a dedicated key/value pair.

• **Improve human readability:**

- Added support for YAML in addition to JSON
- Values, such as, names, types, quantities etc. are now explicitly encoded in dedicated key/value pairs rather than being encoded as regular expressions in keys.

• **Improve direct interpretation of data:**

- Remove ``references`` key. This key was used in previous versions of NWB to generate implicit data structures where datasets store references to part of other metadata structures. These implicit data structures violate core NWB principles as they hinder the direct interpretation of data and cannot be interpreted (neither by human nor program) based on NWB files alone without having additional informaton about the specification as well. Through simple reorganization of metadata in the file, all instances of these implicit data structures were replaced by simple links that can be interpreted directly.

• **Simplified specification of dimensions for datasets:**

- Renamed ``dimensions`` key to ``dims``
- Added key ``shape`` to allow the specification of the shape of datasets
- **Removed custom keys for defining structures as types for dimensions:**
  - \* ``unit`` keys from previous structured dimensions are now ``unit`` attributes on the datasets (i.e., all values in a dataset have the same units)
  - \* The length of the structs are used to define the length of the corresponding dimension as part of the ``shape`` key
  - \* ``alias`` for components of dimensions are currently encoded in the dimensions name.

- **Added support for default vs. fixed name for groups and datasets:**
  - Added `default_name` key for groups and dataset to allow the specification of default names for objects that can have user-defined names (in addition to fixed names via `name`). Attributes can only have a fixed name since attributes can not have a `neurodata_type` and can, hence, only be identified via their fixed name.
- **Updated specification of fixed and default values for attributes to make the behavior of keys explicit:**
  - **Specifying attribute values:**
    - \* Added `default_value` key for attributes to specify a default value for attributes
    - \* Removed `const` key for attributes which was used to control the behavior of the `value` key, i.e., depending on the value of `const` the `value` key would either act as a fixed or default value. By adding the `default_value` key this behavior now becomes explicit and the behavior of the `value` key no longer depends on the value of another key (i.e., the `const` key)
- **Improved governance and reuse of specifications:**
  - The core specification documents are no longer stored as `.py` files as part of the original Python API but are released as separate YAML (or optionally JSON) documents in a separate repository
  - All documentation has been ported to use reStructuredText (RST) markup that can be easily translated to PDF, HTML, text, and many other forms.
  - Documentation for source codes and the specification are auto-generated from source to ensure consistency between sources and the documentation
- **Avoid mixing of format specification and computations:**
  - Removed key ``autogen`` (without replacement). The `autogen` key was used to describe how to compute certain derived datasets from the file. This feature was problematic with respect to the guiding principles of NWB for a couple of reasons. E.g., the resulting datasets were often not interpretable without the provenance of the autogeneration procedure and autogeneration itself and often described the generation of derived data structures to ease follow-on computations. Describing computations as part of a format specification is problematic as it creates strong dependencies and often unnecessary restrictions for use and analysis of data stored in the format. Also, the reorganization of metadata has eliminated the need for `autogen` in many cases. A `autogen` features is arguably the role of a data API or intermediary derived-quantity API (or specification), rather than a format specification.
- **Others:**
  - Removed key ``\_custom`` (without replacement). This feature was used only in one location to provide user hints where custom data could be placed, however, since the NWB specification approach explicitly allows users to add custom data in any location, this information was not binding.

## Currently unsupported features:

- ``_required`` : The current API does not yet support specification and verification of constraints previously expressed via `_required`.
- Relationships are currently available only through implicit concepts, i.e., by sharing dimension names and through implicit references as part of datasets. The goal is to provide explicit mechanisms for describing these as well as more advanced relationships.
- ``dimensions_specification`` : This will be implemented in later version likely through the use of relationships.

## YAML support

To improve human readability of the specification language, Version 1.2a now allows specifications to be defined in YAML as well as JSON (Version 1.1c allowed only JSON).

### ``quantity``

Version 1.1c of the specification language used a ``quantity_flag`` as part of the name key of groups and datasets to the quantity

- `!` - Required (this is the default)
- `?` - Optional
- `^` - Recommended
- `+` - One or more instances of variable-named identifier required
- `*` - Zero or more instances of variable-named identifier allowed

Version 1.2a replaces the ``quantity_flag`` with a new key ``quantity`` with the following values:

value	required	number of instances
<code>`zero_or_more` or `*`</code>	optional	unlimited
<code>`one_or_more` or `+`</code>	required	unlimited but at least 1
<code>`zero_or_one` or `?`</code>	optional	0 or 1
<code>`1`, `2`, `3`, ...</code>	required	Fixed number of instances as indicated by the value

### ``merge` and `include``

To simplify the concept ``include`` and ``merge``, version 1.2a introduced a new key ``neurodata_type_def`` which describes the creation of a new `neurodata_type`. The combination ``neurodata_type_def`` and ``neurodata_type_inc`` simplifies the concepts of merge (i.e., inheritance/extension) and inclusion and allows us to express the same concepts in an easier-to-use fashion. Accordingly, the keys ``include``, ``merge`` and ``merge+`` have been removed in version 1.2a. Here a summary of the basic cases:

neuro- data_type_inc	neuro- data_type_def	Description
not set	not set	define standard dataset or group without a type
not set	set	create a new <code>neurodata_type</code> from scratch
set	not set	include (reuse) <code>neurodata_type</code> without creating a new one (include)
set	set	merge/extend <code>neurodata_type</code> and create a new type (merge)

### ``structured_dimensions``

The definition of structured dimensions has been removed in version 1.2a. The concept of structs as dimensions is problematic for several reasons: 1) it implies support for defining general tables with mixed units and data types which are currently not supported, 2) they easily allow for colliding specification where mixed units are assigned to the same value, 3) they are hard to use and unsupported by HDF5. Currently structured dimensions, however, have been used only to encode information about “columns” of a dataset (e.g., to indicate that a dimension stores x,y,z values). This information was translated to the `dims`` and `shape`` keys and `unit`` attributes. The more general concept of structured dimensions will be implemented in future versions of the specification language and format likely via support for modeling of relationships or support for table data structures (stay tuned)

``autogen``

The ``autogen`` key has been removed in 1.2a without replacement.

## Version 1.1c (Oct. 7, 2016)

- Original version of the specification language generated as part of the NWB pilot project



### Contents

- *Credits*
  - *Authors*
    - \* *NWB-N: Version 1.1.x and later*
    - \* *NWB-N: Version 1.0.x and earlier*
  - *Acknowledgments*
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  - *Copyright*
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## Authors

### NWB-N: Version 1.1.x and later

Version 1.1.x of the NWB-N format, Version 1.2.x and later of the specification language, and Version 1.x and later of the NWB storage documents have been created by Oliver Ruebel and Andrew Tritt et al. in collaboration with original members of the NWB pilot project at the University of California in Berkeley and San Francisco and at the Allen Institute for Brain Science.

## NWB-N: Version 1.0.x and earlier

Version 1.0.5g (and earlier) of the NWB file format and Version 1.1c (and earlier) of the specification language were created by Jeff Teeters et al. as part of the first NWB pilot project. The documents governing the specification language and format have been adopted from the final versions released by the original NWB pilot project.

## Acknowledgments

### 2016 - 2017

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**Note: #TODO Add Acknowledgement for LBL Kavli project and acknowledge other projects as well**

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### 2015 - 2016

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## CHAPTER 4

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### Indices and tables

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- `modindex`
- `search`