RNeXML: a package for reading and writing richly annotated phylogenetic, character, and trait data in R

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9 Abstract

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- 1. NeXML is a powerful and forward-compatible exchange standard recently proposed to better meet the expanding needs for machine-proof phylogenetic data and metadata sharing. Here we present the RNeXML package, which aims to provide to users of the popular R programming language easy programmatic access to reading and writing NeXML documents, including rich metadata, in a way that interfaces seamlessly with the extensive library of phylogenetic tools already available in the R ecosystem.
 - 2. Wherever possible, we designed RNeXML to map NeXML document contents, whose arrangement is influenced by the format's normative XML Schema definition, to their most intuitive or useful representation in R. To make NeXML's powerful facility for recording semantically rich machine-readable metadata accessible to R users, we designed a functional programming interface to it that hides the semantic web standards leveraged by NeXML from R users who are unfamiliar with them.
 - 3. RNeXML can read any NeXML document that validates, and it generates valid NeXML documents from phylogeny and character data in various R representations in use. The metadata programming interface at a basic level aids fulfilling data documentation best practices, and at an advanced level preserves NeXML's nearly limitless extensibility, for which we provide a fully working demonstration. Furthermore, to lower the barriers to sharing well-documented phylogenetic data, RNeXML has started to integrate with taxonomic metadata augmentation services on the web, and with online repositories for data archiving.
 - 4. RNeXML allows R's rich ecosystem to read and write data in the NeXML format through an interface that is no more involved than reading or writing data from other, less powerful data

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formats. It also provides an interface designed to feel familiar to R programmers and to be consistent with recommended practices for R package development, yet that retains the full power for users to add their own custom data and metadata to the phylogenies they work with, without introducing potentially incompatible changes to the exchange standard.

5 Introduction

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Users of the popular statistical and mathematical computing platform R (R Core Team 2014) enjoy a wealth of readily installable comparative phylogenetic methods and tools (O'Meara 2014). Exploiting the opportunities arising from this wealth for complex and integrative comparative research questions relies on the ability to reuse and integrate previously generated or published data and metadata. The expanding data exchange needs of the evolutionary research community are rapidly outpacing the capabilities of most current and widely used data exchange standards (Vos et al. 2012), which were all developed a decade or more ago. This has resulted in a radiation of different data representations and exchange standard "flavors" that are no longer interoperable at the very time when the growth of available data and methods has made that interoperability most valuable. In response to the unmet needs for standardized data exchange in phylogenetics, a modern XML-based exchange standard, called NeXML, has recently been developed (Vos et al. 2012). NeXML comprehensively supports current data exchange needs, is predictably machine-readable, and is forward compatible.

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The exchange problem for phylogenetic data is particularly acute in light of the challenges in finding and sharing phylogenetic data without the otherwise common loss of most data and metadata semantics (Stoltzfus et al. 2012; Drew et al. 2013; Cranston et al. 2014). For example, the still popular NEXUS file format (Maddison et al. 1997) cannot consistently represent horizontal gene transfer or ambiguity in reading a character (such as a DNA sequence base pair). This and other limitations have led to modifications of NEXUS in different ways for different needs, with the unfortunate result that NEXUS files generated by one program can be incompatible with another (Vos et al. 2012). Without a formal grammar, software based on NEXUS files may also make inconsistent assumptions about tokens, quoting, or element lengths. Vos et al. (2012) estimates that as many as 15% of the NEXUS files in the CIPRES portal contain unrecoverable but hard to diagnose errors.

A detailed account of how the NeXML standard addresses these and other relevant challenges can be found in Vos et al. (2012). In brief, NeXML was designed with the following important properties. First,

NeXML is defined by a precise grammar that can be programmatically **validated**. This ensures that any
NeXML file can be expected to be parsed and read without errors by any NeXML-compliant software,
provided the file passes validation. Second, NeXML is **extensible**: a user can define representations of
new, previously unanticipated information (as we will illustrate) without violating its defining grammar.
Third and most importantly, NeXML is **rich in computable semantics**. The formal semantics used
in NeXML is machine-readable in a predictable manner, and also enables machine reasoning about
the data and metadata (see Parr et al. (2011)), especially when metadata attributes and values are
anchored in appropriately axiomatized ontologies.

To make the capabilities of NeXML available to R users in an easy-to-use form, and to lower the hurdles to adoption of the standard, we present RNeXML, an R package that aims to provide easy programmatic access to reading and writing NeXML documents, tailored for the kinds of use-cases that will be common for users and developers of the wealth of evolutionary analysis methods within the R ecosystem.

73 The RNeXML package

The RNeXML package is written entirely in R and available under a Creative Commons Zero public domain waiver. The current development version can be found on Github at https://github.com/ropensci/RNeXML, and the stable version can be installed from the CRAN repository. RNeXML is part of the rOpenSci project. Users of RNeXML are encouraged to submit bug reports or feature requests in the issues log on Github, or the phylogenetics R users group list at r-sig-phylo@r-project.org for help. Vignettes with more detailed examples of specific features of RNeXML are distributed with the R package and serve as a supplement to this manuscript. Each of the vignettes can be found at http://ropensci.github.io/RNeXML/.

Representation of NeXML documents in R

Conceptually, a NeXML document has the following components: (1) phylogeny topology data, (2)
character or trait data in matrix form, (3) operational taxonomic units (OTUs), and (4) metadata.
To represent the contents of a NeXML document (currently in memory), RNeXML defines the nexml
object type. This type therefore holds phylogenetic trees as well as character or trait matrices, and
all metadata, which is similar to the phylogenetic data object types defined in the phylobase package

- (NESCENT R Hackathon Team 2014), but contrasts with the more widely used ones defined in the ape package (Paradis *et al.* 2004), which represents trees alone.
- When reading and writing NeXML documents, RNeXML aims to map their components to and from,
- 91 respectively, their most widely used representations in R. As a result, the types of objects accepted
- 92 or returned by the package's methods are the phylo and multiPhylo objects from the ape package
- 93 (Paradis et al. 2004) for phylogenies, and R's native data.frame list structure for data matrices.
- 94 Reading phylogenies and character data
- The method nexml read() reads NeXML files, either from a local file, or from a remote location via
- of its URL, and returns an object of type nexml:

```
nex <- nexml_read("components/trees.xml")</pre>
```

- The method get trees list() can be used to extract the phylogenies as an ape::multiPhylo object,
- which can be treated as a list of ape::phylo objects:

```
phy <- get_trees_list(nex)</pre>
```

The get_trees_list() method is designed for use in scripts, providing a consistent and predictable return type regardless of the number of phylogenies a NeXML document contains. For greater convenience in interactive use, the method get_trees() returns the R object most intuitive given the arrangement of phylogeny data in the source NeXML document. For example, the method returns an ape::phylo object if the NeXML document contains a single phylogeny, an ape::multiPhylo object if it contains multiple phylogenies arranged in a single trees block, and a list of ape::multiPhylo objects if it contains multiple trees blocks (the capability for which NeXML inherits from NEXUS).

If the location parameter with which the nexml_read() method is invoked is recognized as a URL, the method will automatically download the document to the local working directory and read it from there. This gives convenient and rapid access to phylogenetic data published in NeXML format on the web, such as the content of the phylogenetic data repository TreeBASE (Piel et al. 2002). For example, the following plots a tree in TreeBASE (using ape's plot function):

```
tb_nex <- nexml_read(
"https://raw.github.com/TreeBASE/supertreebase/master/data/treebase/S100.xml")
tb_phy <- get_trees_list(tb_nex)
plot(tb_phy[[1]])</pre>
```

The method get_characters() obtains character data matrices from a nexml object, and returns them as a standard data.frame R object with columns as characters and rows as taxa:

```
nex <- nexml_read("components/comp_analysis.xml")
get_characters(nex)</pre>
```

113		log snout-vent length	reef-dwelling
114	taxon_8	-3.2777799	0
115	taxon_9	2.0959433	1
116	taxon_10	3.1373971	0
117	taxon_1	4.7532824	1
118	taxon_2	-2.7624146	0
119	taxon_3	2.1049413	0
120	taxon_4	-4.9504770	0
121	taxon_5	1.2714718	1
122	taxon_6	6.2593966	1
123	taxon_7	0.9099634	1

A NeXML data matrix can be of molecular (for molecular sequence alignments), discrete (for most morphological character data), or continuous type (for many trait data). To enable strict validation 125 of data types NeXML allows, and if their data types differ requires multiple data matrices to be 126 separated into different "blocks". Since the data.frame data structure in R has no such constraints, the 127 get_characters() method combines such blocks as separate columns into a single data.frame object, 128 provided they correspond to the same taxa. Otherwise, a list of data.frames is returned, with list 129 elements corresponding to characters blocks. Similar to the methods for obtaining trees, there is also a 130 method get_characters_list(), which always returns a list of data.frames, one for each character 131 block. 132

133 Writing phylogenies and character data

The method nexml_write() generates a NeXML file from its input parameters. In its simplest invocation, the method writes a tree to a file:

```
data(bird.orders)
nexml_write(bird.orders, file = "birds.xml")
```

The first argument to nexml_write() is either an object of type nexml, or any object that can be coerced to it, such as in the above example an ape::phylo phylogeny. Alternatively, passing a multiPhylo object would write a list of phylogenies to the file.

In addition to trees, the nexml_write() method also allows to specify character data as another parameter. The following example uses data from the comparative phylogenetics R package geiger (Harmon et al. 2008).

142 Validating NeXML

File validation is a central feature of the NeXML format which ensures that any properly implemented
NeXML parser will always be able to read the NeXML file. The function takes the path to any NeXML
file and returns TRUE to indicate a valid file, or FALSE otherwise, along with a display of any error
messages generated by the validator.

```
nexml_validate("geospiza.xml")
```

147 [1] TRUE

The nexml_validate() function performs this validation using the online NeXML validator (when a network connection is available), which performs additional checks not expressed in the NeXML schema itself (Vos et al. 2012). If a network connection is not available, the function falls back on the schema validation method from the XML package (Lang 2013).

152 Creating and populating nexml objects

Instead of packaging the various components for a NeXML file at the time of writing the file, RNeXML also allows users to create and iteratively populate in-memory nexml objects. The methods to do this are add_characters(), add_trees(), and add_meta(), for adding characters, trees, and metadata, respectively. Each of these functions will automatically create a new nexml object if not supplied with an existing one as the last (optional) argument.

For example, here we use add_trees() to first create a nexml object with the phylogeny data, and then add the character data to it:

```
nexObj <- add_trees(geospiza$phy)
nexObj <- add_characters(geospiza$dat, nexObj)</pre>
```

The data with which a nexml object is populated need not share the same OTUs. RNeXML automatically adds new, separate OTU blocks into the NeXML file for each data matrix and tree that uses a different set of OTUs.

Other than storage size, there is no limit to the number of phylogenies and character matrices that can be included in a single NeXML document. This allows, for example, to capture samples from a posterior probability distribution of inferred or simulated phylogenies and character states in a single NeXML file.

Data documentation and annotation with built-in metadata

NeXML allows attaching ("annotating") metadata to any data element, and even to metadata themselves.

Whether at the level of the document as a whole or an individual data matrix or phylogeny, metadata

can provide bibliographic and provenance information, for example about the study as part of which

the phylogeny was generated or applied, which data matrix and which methods were used to generate it.

Metadata can also be attached to very specific elements of the data, such as specific traits, individual

OTUs, nodes, or even edges of the phylogeny.

As described in Vos et al. (2012), to encode metadata annotations NeXML uses the "Resource Description Framework in Annotations" (RDFa) (Prud'hommeaux 2014). This standard provides for a strict machine-readable format yet enables future backwards compatibility with compliant NeXML

- parsers (and thus RNeXML), because the capacity of a tool to *parse* annotations is not predicated on understanding the meaning of annotations it has not seen before.
- To lower the barriers to sharing well-documented phylogenetic data, RNeXML aims to make recording useful and machine-readable metadata easier at several levels.
- First, when writing a NeXML file the package adds certain basic metadata automatically if they are absent, using default values consistent with recommended best practices (Cranston *et al.* 2014).
- Currently, this includes naming the software generating the NeXML, a time-stamp of when a tree was produced, and an open data license.
- Second, RNeXML provides a simple method, called add_basic_metadata(), to set metadata attributes commonly recommended for inclusion with data to be publicly archived or shared (Cranston *et al.* 2014). The currently accepted parameters include title, description, creator, pubdate, rights, publisher, and citation. Behind the scenes the method automatically anchors these attributes in common vocabularies (such as Dublin Core).
- Third, RNeXML integrates with the R package taxize (Chamberlain & Szöcs 2013) to mitigate one of
 the most common obstacles to reuse of phylogenetic data, namely the misspellings and inconsistent
 taxonomic naming with which OTU labels are often fraught. The taxize_nexml() method in RNeXML
 uses taxize to match OTU labels against the NCBI database, and, where a unique match is found, it
 annotates the respective OTU with the matching NCBI identifier.
- Data annotation with custom metadata
- The RDFa standard adopted by NeXML (see Vos et al. (2012)) allows for arbitrary custom extensions to meet ongoing scientific needs without requiring modifications (or "extensions") of the syntax. Specifically, every metadata annotation consists of a metadata property and a value of the property; the bearer of the property is determined by where the annotation appears in the NeXML document (technically, the containing element in the XML document model). This structure corresponds directly to Resource Description Framework (RDF) subject-predicate-object triples, and therefore the only requirement for custom metadata is that they be cast as RDF triples.
- The RNeXML interface described above for built-in metadata annotations perform this cast automatically behind the scenes, including mapping metadata attributes to terms in requisite common vocabularies (such as Dublin Core for "title", "creator", etc.) or ontologies. Giving metadata properties (and

where applicable, values) as vocabulary or ontology terms rather than simple text strings is crucial for allowing machines to not only parse but also interpret and potentially reason over their semantics. 207 To achieve this benefit for custom metadata extensions, the user necessarily needs to handle certain 208 technical details from which the RNeXML API shields her otherwise, in particular the globally unique 209 identifiers (normally HTTP URIs) of metadata terms and vocabularies. To be consistent with XML 210 terminology, RNeXML calls vocabulary URIs namespaces, and their abbreviations prefixes. For example, 211 the namespace for the Dublin Core Metadata Terms vocabulary is "http://purl.org/dc/elements/1.1/". 212 Using its common abbreviation "dc", a metadata property "dc:title" expands to the identifier "http: 213 //purl.org/dc/elements/1.1/title". This URI resolves to a human and machine-readable (depending on 214 access) definition of precisely what the term title in Dublin Core means. In contrast, just using the 215 text string "title" could also mean the title of a person, a legal title, the verb title, etc. URI identifiers of metadata vocabularies and terms are not mandated to resolve, but if machines are to derive the 217 maximum benefit from them, they should resolve to a definition of their semantics in RDF. 218

RNeXML includes methods to obtain and manipulate metadata properties, values, identifiers, and namespaces. The get_namespaces() method accepts a nexml object and returns a named list of namespace prefixes and their corresponding identifiers known to the object:

```
birds <- nexml_read("birds.xml")
prefixes <- get_namespaces(birds)
prefixes["dc"]</pre>
```

```
dc dc "http://purl.org/dc/elements/1.1/"
```

The get_metadata() method returns, as a named list, the metadata annotations for a given nexml object at a given level, with the whole NeXML document being the default level ("all" extracts all metadata objects):

```
meta <- get_metadata(birds)
otu_meta <- get_metadata(birds, level="otu")</pre>
```

The returned list does not include the data elements to which the metadata are attached. Therefore, a different approach, documented in the metadata vignette, is recommended for accessing the metadata attached to data elements.

The meta() method creates a new metadata object from a property name and content (value). For example, the following creates a modification date metadata object, using a property in the PRISM vocabulary:

```
modified <- meta(property = "prism:modificationDate", content = "2013-10-04")</pre>
```

- Metadata annotations in NeXML can be nested within another annotation, which the meta() method accommodates by accepting a parameter children, with the list of nested metadata objects (which can themselves be nested) as value.
- The add_meta() function adds metadata objects as annotations to a nexml object at a specified level,
 with the default level being the NeXML document as a whole:

```
birds <- add_meta(modified, birds)</pre>
```

If the prefix used by the metadata property is not among the built-in ones (which can be obtained using get_namespaces()), it has to be provided along with its URI as the namespaces parameter. For example, the following uses the SKOS vocabulary (which currently is not built-in) to add a note to the trees in the nexml object:

- Alternatively, additional namespaces can also be added in batch using the add_namespaces() method.
- By virtue of subsetting the S4 nexml object, RNeXML also offers fine control of where a meta element is added, for which the package vignette on S4 subsetting of nexml contains examples.
- Because NeXML expresses all metadata using the RDF standard, and stores them compliant with RDFa, they can be extracted as an RDF graph, queried, analyzed, and mashed up with other RDF data,
- local or on the web, using a wealth of off-the-shelf tools for working with RDF (see Prud'hommeaux
- 248 (2014) or Hartig (2012)). Examples for these possibilities are included in the RNeXML SPARQL vignette,

and the package also comes with a demonstration that can be run from R using the following command:

demo("sparql", "RNeXML")).

251 Using metadata to extend the NeXML standard

NeXML was designed to prevent the need for future non-interoperable "flavors" of the standard in response to new research directions. Its solution to this inevitable problem is a highly flexible metadata system without sacrificing strict validation of syntax and structure.

Here we illustrate how RNeXML's interface to NeXML's metadata system can be used to record and share a type of phylogenetic data not taken into account when NeXML was designed, in this case stochastic character maps (Huelsenbeck et al. 2003). Such data assign certain parts (corresponding to time) of each branch in a time-calibrated phylogeny to a particular "state" (typically of a morphological characteristic). The current de-facto format for sharing stochastic character maps, created by simmap (Bollback 2006), a widely used tool for creating such maps, is a non-interoperable modification of the standard Newick tree format.

To express these data as a custom metadata extension, we create metadata annotations for the edge elements in the phylogeny topology. The metadata need to describe the character state being assigned, and the duration (in terms of branch length) that the edge spends in that state. The NeXML standard already defines state elements for discrete character traits, which, as most elements in NeXML, also have unique identifiers by which they can be referenced.

In the following proposal, we term each series of character state assignments for an edge a reconstruction, and for each state assignment comprising a reconstruction, we provide the length, the identifier of the character state, and an ordinal property to record the chronological order of the assignment. In this manner, a single edge containing one state change could be annotated as follows. (the identifier values in the proposal, "cr1", "s1", and "s2", are identifiers local to a – fictitious – NeXML document, based on the example simmap tree illustrated in Revell (2012)):

```
m <- meta("simmap:reconstructions", children = c(
    meta("simmap:reconstruction", children = c(
    meta("simmap:char", "cr1"),
    meta("simmap:stateChange", children = c(</pre>
```

```
meta("simmap:order", 1),
    meta("simmap:length", "0.2030"),
    meta("simmap:state", "s2"))),

meta("simmap:char", "cr1"),
    meta("simmap:stateChange", children = c(
        meta("simmap:order", 2),
        meta("simmap:length", "0.0022"),
        meta("simmap:state", "s1")))
))))
```

The example uses a prefix, simmap, to group the newly introduced metadata properties in a vocabulary,

for which the add_namespace() method can be used to give a URI as an identifier:

```
nex <- add_namespaces(c(simmap = "https://github.com/ropensci/RNeXML/tree/master/inst/simmap.md"</pre>
```

Here the URI does not resolve to a fully machine-readable definition of the terms and their semantics,

but it can nonetheless be used to provide at least a human-readable informal definition of the terms.

277 Because the above representation does not modify the NeXML format, it should be readable – even if

278 not necessarily interpretable as a stochastic character mapping – by any compliant NeXML parser. In

²⁷⁹ addition, it includes a direct and machine-readable link to metadata documentation.

²⁸⁰ Writing out metadata annotations as in the above proposal may seem tedious. However, by virtue of

281 RNeXML's API it is straightforward to define functions that encapsulate these details. As a demonstration,

282 RNeXML comes with functions simmap_to_nexml() and nexml_to_simmap() that map bidirectionally

between NeXML files with simmap metadata according to the above proposal, and the extension of the

ape::phylo class devised by Revell (2012) in the R package phytools. The RNeXML simmap vignette

285 illustrates how this allows seamlessly generating and reading NeXML files containing simmap data, and

using the various functions from phytools designed for simmap objects (Revell 2012).

Publishing NeXML files from R

Data archiving is increasingly required by scientific journals, including in evolutionary biology, ecology,

289 and biodiversity (e.g. Rausher et al. (2010)). The effort involved with preparing and submitting

properly annotated data to archives remains a notable barrier to the broad adoption of data archiving
and sharing as a normal part of the scholarly publication workflow (Tenopir et al. 2011; Stodden 2014).

In particular, the majority of phylogenetic trees published in the scholarly record are inaccessible or lost
to the research community (Drew et al. 2013).

One of RNeXML's aims is to promote the archival of well-documented phylogenetic data in scientific data
repositories, in the form of NeXML files. To this end, the method nexml_publish() provides an API
directly from within R that allows data archival to become a step programmed into data management
scripts. Initially, the method supports the data repository Figshare (http://figshare.com):

```
doi <- nexml_publish(birds, visibility = "public", repository="figshare")</pre>
```

Figshare also supports "private" visibility, allowing to securely backup data to a repository and to share
them with collaborators prior to public release.

RNeXML allows R's ecosystem to read and write data in the NeXML format through an interface that is

300 Conclusions and future directions

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no more involved than reading or writing data from other phylogenetic data formats. It also carries 302 immediate benefits for its users compared to other formats. For example, comparative analysis R 303 packages and users frequently add their own metadata annotations to the phylogenies they work with, 304 such as annotations of species, stochastic character maps, trait values, model estimates and parameter values. RNeXML affords R the capability to harness machine-readable semantics and an extensible 306 metadata schema to capture, preserve, and share these and other kinds of information, all through an 307 API instead of having to understand in detail the schema underlying the NeXML standard. To assist users in meeting the rising bar for best practices in data sharing in phylogenetic research (Cranston et 309 al. 2014), RNeXML captures metadata information from the R environment to the extent possible, and 310 applies reasonable defaults. 311 The goals for continued development of RNeXML revolve primarily around better interoperability with other existing phylogenetic data representations in R, such those found in the phylobase package 313 (NESCENT R Hackathon Team 2014); and better integration of the rich metadata semantics found in 314 ontologies defined in the Web Ontology Language (OWL), including programmatic access to machine 315 reasoning with such metadata.

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