# Template to prepare preprints and manuscripts using markdown and github actions

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**Purpose:** This template provides a series of scripts to render a markdown document into an interactive website and a series of PDFs.

**Motivation:** It makes collaborating on text with GitHub easier, and means that we never need to think about the output.

**Internals:** GitHub actions and a series of python scritpts. The markdown is handled with pandoc.

- NCBITaxonomy. jl is a package designed to facilitate the reconciliation and cleaning of taxonomic names,
- using a local copy of the NCBI taxonomic backbone (Federhen 2012; Schoch et al. 2020); The basic search
- <sup>3</sup> functions are coupled with quality-of-life functions including case-insensitive search and custom fuzzy
- 4 string matching to facilitate the amount of information that can be extracted automatically while allowing
- 5 efficient manual curation and inspection of results. NCBITaxonomy. jl works with version 1.6 of the Julia
- 6 programming language (Bezanson et al. 2017), and relies on the Apache Arrow format to store a local copy
- of the NCBI raw taxonomy files. The design of NCBITaxonomy. jl has been inspired by similar efforts, like
- 8 the R package taxadb (Norman et al. 2020), which provides an offline alternative to packages like taxize
- 9 (Chamberlain & Szöcs 2013).

## 5 Statement of need

- <sup>11</sup> Unambiguously identifying species is a far more challenging task than it may appear. There are a vast
- number of reasons for this. Different databases keep different taxonomic "backbones," i.e. different data
- structures in which names are mapped to species, and organised in a hierarchy. Not all names are unique
- identifiers to groups. For example, Io can either refer to a genus of plants from the aster family, or to a
- genus of molluscs; the genus Mus (of which the house mouse Mus musculus is a species), contains a
- sub-genus also named Mus. Conversely, the same species can have several names, which are valid
- synonyms: for example, the domestic cow *Bos taurus* admits *Bos primigenius taurus* as a valid synonym.
- 18 Taxonomic nomenclature also changes regularly, with groups being split, merged, or moved to a new
- position in the tree of life; this is, notably, a common occurrence with viral taxonomy, each subsequent
- version of which can differ markedly from the last; compare, e.g Lefkowitz et al. (2018) to Walker et al.
- 21 (2020).
- To add to the complexity, one must also consider that most taxa names are at some point manually typed,
- which has the potential to introduce additional mistakes in raw data; it is likely to expect that such
- 24 mistakes may arise when attempting to write down the (perfectly valid) names of the bacterial isolate
- 25 known as Myxococcus llanfairpwllgwyngyllgogerychwyrndrobwllllantysiliogogogochensis, or of the crowned
- 26 slaty flycatcher Griseotyrannus aurantioatrocristatus. These mistakes are more likely when dealing with
- 27 hyper-diverse samples, like plant census (Dauncey et al. 2016; Wagner 2016; Conti et al. 2021). In addition
- 28 to binomial names, the same species can be known by many vernacular (common) names, which are

- language or even region-specific: Ovis aries, for example, has valid English vernaculars including lamb,
- 30 sheep, wild sheep, and domestic sheep.
- 31 All these considerations are actually important when matching species names both within and across
- datasets. Let us consider the following species survey of individual fishes, European chub, Cyprinus
- cephalus, Leuciscus cephalus, Squalius cephalus: all are the same species (S. cephalus), referred to as one of
- the vernacular (European chub) and two formerly accepted names now classified as synonyms. A
- cautious estimate of diversity based on the user-supplied names would give n = 4 species, when there is in
- 36 fact only one.
- A package with the ability to handle the sources of errors outlined above, and especially while provide an
- <sup>38</sup> authoritative classification, can accelerate the work of consuming large volumes of biodiversity data. For
- example, this package was used in the process of developing the CLOVER database (Gibb et al. 2021) of
- 40 host-virus associations, by reconciling the names of viruses and mammals from four different sources,
- where all of the issues described above were present.

### 42 Overview of functionalities

- 43 An up-to-date version of the documentation for NCBITaxonomy, jl can be found online at
- 44 https://docs.ecojulia.org/NCBITaxonomy.jl/stable/, including examples and a documentation of
- every method. The package is released under the MIT license. Contributions can be made in the form of
- 46 issues (bug reports, questions, features suggestions) and pull requests. The package can be checked out
- and installed anonymously from the central Julia repository:

```
# This line should go in the Julia configuration file - note that the path
# will be created if it doesn't exist, and will be used to store the
# raw taxonomic table
ENV["NCBITAXONOMY_PATH"] = joinpath(homedir(), "data", "NCBITaxonomy.jl")
Pkg.add("NCBITaxonomy") # Dowloading the files may take a long time
```

- The package will download the most recent version of the NCBI taxonomy database, and transform in into
- a set of Apache Arrow files ready for use. Note that the NCBITAXONOMY\_PATH can specified on a per-project
- basis, and as long as the package is not re-built, the local set of tables downloaded from NCBI will not
- 51 change; this way, users can re-run an analysis with a guarantee that the underlying taxonomic backbone
- 52 has not changed.

#### 53 Improved name matching

- Name finding is primarily done through the taxon function, which admits either a unique NCBI identifier
- 55 (e.g. taxon(36219) for the bogue Boops boops), a string (taxon("Boops boops")), or a data frame with a
- restricted list of names (see the next section). The taxon method has additional arguments to perform
- fuzzy matching in order to catch possible typos (taxon("Boops bops"; strict=false)), to perform a
- lowercase search (useful when alphanumeric codes are part of the taxon name, like for some viruses), and
- to restrict the the search to a specific taxonomic rank.
- The lowercase search can be a preferable alternative to fuzzy string matching. Consider the string
- 61 Adeno-associated virus 3b it has three names with equal distance (under the Levensthein string
- 62 distance function):

```
julia> similarnames("Adeno-associated virus 3b"; threshold=0.95)
3-element Vector{Pair{NCBITaxon, Float64}}:
   Adeno-associated virus - 3 (ncbi:46350) => 0.96
   Adeno-associated virus 3B (ncbi:68742) => 0.96
Adeno-associated virus 3A (ncbi:1406223) => 0.96
```

- 63 Depending on the operating system, either of these three names can be returned; compare to the output of
- a case insensitive name search:

```
julia> taxon("Adeno-associated virus 3b"; casesensitive=false)
Adeno-associated virus 3B (ncbi:68742)
```

This returns the correct name.

#### 66 Name matching output and error handling

- 67 The taxon function will either return a NCBITaxon object (made of a name and id), or throw either a
- NameHasNoDirectMatch (with instructions about how to possible solve it, using the similarnames
- function), or a NameHasMultipleMatches (listing the possible valid matches, and suggesting to use
- alternativetaxa to find the correct one). Therefore, the common way to work with the taxon function
- would be to wrap it in a try/catch statement:

```
try
  taxon(name)
  # Additional operations with the matched name

catch err
  if isa(err, NameHasNoDirectMatch)
    # What to do if no match is found
  elseif isa(err, NameHasMultipleMatches)
    # What to do if there are multiple matches
  else
    # What to do in case of another error that is not NCBITaxonomy specific
  end
end
```

- These functions will not demand any user input in the form of key presses (though they can be wrapped in
- additional code to allow it), as they are intended to run on clusters without supervision. The taxon
- function has good scaling using muliple threads. For convenience in rapidly getting a taxon for
- 75 demonstration purposes, we also provide a string macro, whereby e.g. ncbi "Procyon lotor" will return
- the taxon object for the raccoon.

#### 77 Name filtering functions

- As the full NCBI names table has over 3 million entries at the time of writing, we have provided a number
- <sub>79</sub> of functions to restrict the scope of names that are searched. These are driven by the NCBI divisions. For
- example of = mammalfilter(true) will return a data frame containing the names of mammals, inclusive

- of rodents and primates, and can be used with e.g. taxon(nf, "Pan"). This has the dual advantage of
- making search faster, but also of avoiding matching on names that are shared by another taxonomic group
- 83 (which is not an issue with *Pan*, but is an issue with *e.g. Io* as mentioned in the introduction).
- Note that the use of a restricted list of names can have significant performance consequences: compare,
- 85 for example, the time taken to return the taxon Pan (ID 9596) in the entire database, in all mammals, and
- 86 in all primates:

Names list	Fuzzy matching	Time (ms)	Allocations	Memory allocated
all	no	23	34	2 KiB
	yes	105	2580	25 MiB
mammalfilter(true)	no	0.55	32	2 KiB
	yes	1.9	551	286 KiB
<pre>primatefilter()</pre>	no	0.15	33	2 KiB
	yes	0.3	92	27 KiB

- 87 Clearly, the optimal search strategy is to (i) rely on name filters to ensure that search are conducted within
- 88 the appropriate NCBI division, and (ii) only rely on fuzzy matching when the strict or lowercase match
- sales fails to return a name, as fuzzy matching can result in order of magnitude more run time and memory
- 50 footprint. These numbers were obtained on a single Intel i7-8665U CPU (@ (1.90GHz). Using
- "chimpanzees" as the search string (the NCBI recognized vernacular for *Pan*) gave qualitatively similar
- 92 results, suggesting that there is no performance cost associated with working with synonyms or verncular
- 93 input data.

#### 94 Quality of life functions

- 95 In order to facilitate working with names, we provide the authority function (gives the full taxonomic
- <sup>96</sup> authority for a name), synonyms (to get alternative valid names), vernacular (for English common
- names), and rank (for the taxonomic rank).

#### 98 Taxonomic lineages navigation

- 99 The children function will return all nodes that are directly descended from a taxon; the descendants
- function will recursively apply this function to all descendants of these nodes, until only terminal leaves
- are reached. The parent function is an "upwards" equivalent, giving that taxon from which a taxon
- descents; the lineage function chains calls to parent until either taxon(1) (the taxonomy root) or an
- arbitrary ancestor is reached.
- The taxonomicdistance function (and its in-place equivalent, taxonomicdistance!, which uses
- memory-efficient re-allocation if the user needs to change the distance between taxonomic ranks) uses the
- Shimatani (2001) approach to reconstruct a matrix of distances based on taxonomy, which can serve as a
- 107 rough proxy when no phylogenies are available.
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