

Semantic-based methods for morphological descriptions: An applied example for Neotropical species of genus *Lepidocyrtus* Bourlet, 1839 (Collembola: Entomobryidae)

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Abstract

The production of semantic annotations has gained renewed attention due to the development of anatomical ontologies and the documentation of morphological data. Two methods are proposed in this production, differing in their methodological and philosophical approaches: class-based method and instance-based method. The first, the semantic annotations are established as class expressions, while in the second, the annotations incorporate individuals. An empirical evaluation of the above methods was applied in the morphological description of Neotropical species of the genus *Lepidocyrtus* (Collembola: Entomobryidae: Lepidocyrtinae). The semantic annotations are expressed as RDF triple, which is a language most flexible than the Entity-Quality syntax used commonly in the description of phenotypes. The morphological descriptions were built in Protégé 5.4.0 and stored in an RDF store created with Fuseki Jena. The semantic annotations based on RDF triple increase the interoperability and integration of data from diverse sources, e.g., museum data. However, computational challenges are present, which are related with the development of semi-automatic methods for the generation of RDF triple, interchanging between texts and RDF triple, and the access by non-expert users.

Keywords

class-based methods, instance-based methods, resource description framework (RDF), Neotropical region, Hexapoda

Introduction

In recent years, anatomical ontologies have gained attention in the formalization of semantic-based morphological descriptions, which encompass the standardization of anatomical terminology and interoperability between morphological data coming from diverse sources (Dahdul et al. 2015). This formalization implies new computational tools and common language that allows building of semantic annotations parsable by a non-human reader. A review of formalized language and software developed for semantic annotations was realized by Yoder et al. (2018). Three standardized languages are employed by the World Wide Web Consortium (W3C): 1) XML together with the introduction of CharaParser (Cui 2012, Cui et al. 2016), 2) NeXML linked to phenotypic descriptions in OWL (Balhoff et al. 2013), and 3) Resource Description Framework (RDF) (Vogt 2017). Software most used for semantic annotations is Phenex, a platform that employs the Entity-Quality syntax (EQ model), which could be translated to character statement (Balhoff et al. 2010):

chaeta (Entity): ciliated (Quality)

chaeta (character): ciliated (character state)

This software was based initially in NeXML language, incorporating after mx, a web-based application to gather information about specimens, and building descriptive matrices (Balhoff et al. 2013). Nevertheless, NeXML is challenged for non-expert users and mx is disabled currently. On the other hand, the class-based method presupposes the identification *a priori* of comparative homologs (Vogt 2017). The EQ model is a class-based method, where the generalizations and phenotypes are described within the definition of ontology class (Vogt 2019), containing TBox assertions, axioms that defining classes, and the relations between them (De Giacomo and Lenzerini 1996). This model is employed in morphological descriptions and building of character statements (Balhoff et al. 2012, Burks et al. 2016, Trietsch et al. 2018, Yoder et al. 2018, Tarasov 2019)

A second approach called “semantic instance anatomy” is developed by Vogt (2017) and Vogt (2019). In this method, each anatomical entity and its qualities are represented by individual resources, being themselves instances (Vogt 2019), containing ABox assertions, axioms on individual objects or instances (De Giacomo and Lenzerini 1996). While the class-based method contains metadata from different organisms, “semantic instance anatomy” contains metadata for an organism documented in a separate semantic graph (Vogt 2019). The documentation is made through ontology classes from Ontology for Biomedical Investigations (OBI) to specify how the specimens are preserved, and from Biological Collection Ontology to related collection information as catalog number or

collection number (Vogt 2019). Both methods employ predicates that describe topological relations between anatomical entities, and relations between anatomical entities and their qualities.

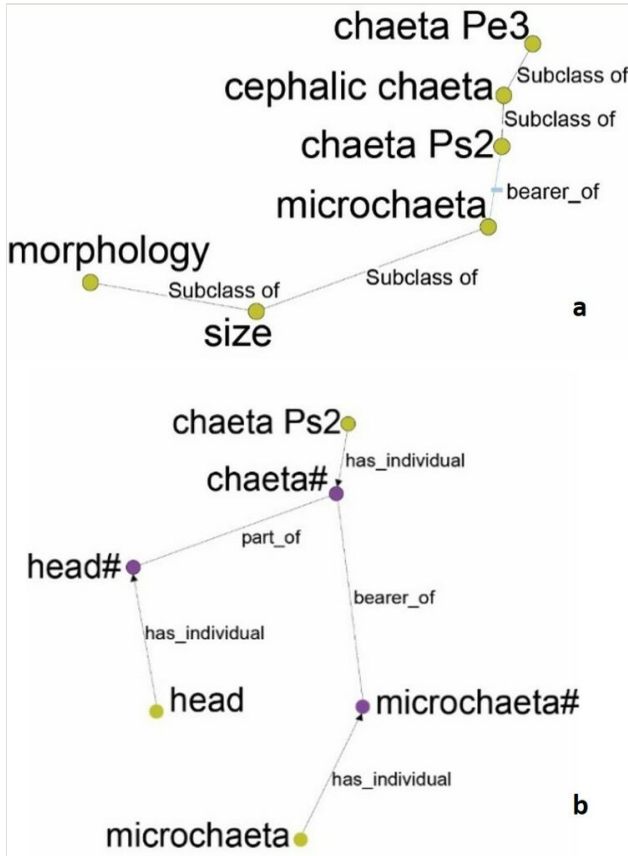


Figure 1. [doi](#)

Graphs of internal structure of **a)** class-based and **b)** instance-based methods. The orange points represent classes within the descriptive template that in the example refers to the chaeta Ps2 part of cephalic chaetae. The purple points represent instances or individuals for each class and are named with provisional labels. The SubClassOf relation links classes, and the has_individual relation links class and individual instanced.

Philosophical differences rely on objects that are described in each method, classes, and instances (Fig. 1), and how these are approached epistemologically. While the classes are generalizations, the instances are the result of direct observation of objects (parts of organisms) that exist in the reality and independent of human cognition (Mahner and Bunge 1997). The development of semantic-based morphological descriptions has been extensive within Hymenoptera (Mikó et al. 2015, Mikó et al. 2016, Silva and Feitosa 2019). Of course, the application of these semantic methods is strongly related to the building of anatomical ontologies, but these are scarce and only available for few taxa, Hymenoptera

Ontology Anatomy (HAO) (Yoder et al. 2010), Mosquito Gross Anatomy Ontology (TGMA) (Grumbling and Strelets 2006), Drosophila Gross Anatomy (FBBT) (Grumbling and Strelets 2006), Spider Ontology (SPD) (Ramírez and Michalik 2019), Tick Anatomy Ontology (TADS) (Topalis et al. 2008), and Ontology for the Anatomy of the Insect SkeletoMuscular system (AISM) (Girón et al. 2021). Recently, Collembola Anatomy Ontology (CLAO) (Gonzalez-Montaña 2021) has been developed (available at www.ontobee.org), offering an opportunity in the application of semantic methods for morphological descriptions in Collembola. The goal of this paper is a demonstration of the applicability of semantic-based methods in morphological descriptions for the Neotropical species of genus *Lepidocyrtus* Bourlet, 1839, the second larger genus within Entomobryidae, with almost 512 species worldwide (Bellinger et al. 2021) and 40 species described for the Neotropical Region (Mari Mutt and Bellinger 1990).

Material and methods

Descriptive templates

Unfortunately, approaches to the production of semantic annotations in a standardized language are scarce or not available for non-expert users. A method is the building of semantic spreadsheet templates where classes, individuals, and properties are declared, with which an ontology is built. A good example is developed by Girón et al. 2021 found at https://github.com/insect-morphology/aism/blob/master/AISM_template_examples.tsv, where the description of cercus through axioms is made for various insects orders. Another application is proto.morphdbase.de, which allows the declaration of individuals, but currently not available. In this paper, the morphological descriptions are based on the building of a “descriptive template”, which is a .owl file to hold the anatomical terms (e. g. chaetotaxy) manually imported from CLAO. The import involves the search and the declaration within the template of terms related with anatomical entities described for *Lepidocyrtus*. Optional tools to import terms could be useful as ODK (<https://github.com/INCATools/ontology-development-kit/>) or Ontofox (<http://ontofox.hegroup.org>), increasing the performance in this step, but the terms curation must be done by an expert in the taxon. The “descriptive template” has classes and subclasses organized in a hierarchical structure, where class expressions or direct instances are the semantic annotations for each species. For instance, class expression for the absence of scales on antennae is: “**antenna** *SubClassOf* not (*bearer_of* some **scaled**)”, while the presence of scales is expressed as “**antenna** *SubClassOf* (*bearer_of* some **scaled**)”.

Continuing with the example above, the class **antenna** represents an individuals set that shares some property, e. g., part of the head capsule. The individuals are described object property or data property assertion could be made as:

“**antenna** *has_individual* some antenna123”, where the individual labeled as “antenna123” is the instance of the class **antenna**.

and

“antenna123 *part_of* headcapsule123”, where “*part_of* abdominalsegment1” is the object property assertion and “headcapsule123” is a provisional label for an individual that is an instance of the class **head capsule**.

Under the class-based method, the template has a “layer”, the class expressions, and under the instance-based method, the template has two “layers”, the class expressions, and individuals by class.

The morphological descriptions follow the HYA model proposed by Wirkner et al. 2017, including the phenotypic categories: 1. qualitative phenotypes, 2. presence/absence phenotypes, 3. count phenotypes, and 4. relative measurement phenotypes. Terminology related to the description of phenotypes were taken from Phenotype And Trait Ontology (PATO) (Table 1): anatomical side, color pattern, length, morphology (shape, size, and texture), and number; the spatial object properties are enlisted in the Table 2. Semantic annotations are declared through OWL class expressions and Manchester Syntax (<http://www.w3.org/TR/owl2-manchester-syntax/>) and built in Protégé 5.4.0 Musen 2015.

Table 1.

Predicates or relations employed during the expression of RDF triple statement under class-based and instance-based methods. Relation Ontology (RO); Phenotype And Trait Ontology (PATO).

Term	definition	Ontology identifier
globular	A spheroid quality inhering in a bearer by virtue of the bearer's resembling a ball	PATO_0001499
cylindrical	A convex 3-D shape quality inhering in a bearer by virtue of the bearer's exhibiting a consistently sized round cross section	PATO_0001873
absent	A quality denoting the lack of an entity.	PATO_0000462
rounded	A shape quality inhering in a bearer by virtue of the bearer's being such that every part of the surface or the circumference is equidistant from the center	PATO_0000411
Protruding	A quality inhering in a bearer by virtue of the bearer's extending out above or beyond a surface or boundary.	PATO_0001598
truncated	A shape quality inhering in a bearer by virtue of the bearer's terminating abruptly by having or as if having an end or point cut off	PATO_0000936
lanceolate	A shape quality inhering in a bearer by virtue of the bearer's being shaped like a lance-head, considerably longer than wide, tapering towards the tip from below the middle; attached at the broad end	PATO_0001877
serrated	A shape quality inhering in a bearer by virtue of having sharp straight-edged teeth pointing to the apex.	PATO_0001206
asymmetrically curved	A curvature quality inhering in a bearer by virtue of the bearer's being curved asymmetrically.	PATO_0001848

Term	definition	Ontology identifier
domed	A curvature quality inhering in a bearer by virtue of the bearer's having a shape resembling a dome.	PATO_0001789
increased curvature	A curvature which is relatively high.	PATO_0001592
distributed	A spatial pattern inhering in a bearer by virtue of the bearer's being spread out or scattered about or divided up.	PATO_0001566
subulate	A shape quality inhering in a bearer by virtue of the bearer's being linear, very narrow, tapering to a very fine point from a narrow base.	PATO_0001954
filamentous	A shape quality inhering in a bearer by virtue of the bearer's having thin filamentous extensions at its edge.	PATO_0001360
semicircular	A 2-D shape quality inhering in a bearer by virtue of the bearer's having shape or form of half a circle.	PATO_0002232
branched	A branchiness quality inhering in a bearer by virtue of the bearer's having branches.	PATO_0000402
increased amount	An amount which is relatively high.	PATO_0000470
decreased amount	An amount which is relatively low.	PATO_0001997
right side of	A spatial quality inhering in a bearer by virtue of the bearer's being located on right side of a another entity.	PATO_0001793
left side of	A spatial quality inhering in a bearer by virtue of the bearer's being located on left side of from the a another entity.	PATO_0001792
aligned with	An alignment quality inhering in a bearer by virtue of the bearer's being in a proper spatial positioning with respect to an additional entity.	PATO_0001653
normal	A quality inhering in a bearer by virtue of the bearer's exhibiting no deviation from normal or average.	PATO_0000461

RDF repository

An RDF store was built with Apache Jena Fuseki, an HTTP interface for querying RDF graphs, which can be explored in a browser as <http://localhost:3030/query.html>, employing SPARQL according to W3C recommendations (The World Wide Web Consortium). Apache Jena Fuseki was chosen by the simplicity in the installation, in contrast with another web API with SPARQL endpoint, for instance, Openlink Virtuoso, Ontotext, or Neo4j. This created RDF store only holds class-based morphological descriptions for each described species.

Table 2.

Predicates or relations employed during the expression of RDF triple statement under class-based and instance-based methods. Relation Ontology (RO); Phenotype And Trait Ontology (PATO). The relation *bearer_of* is a alternative term of *has_characteristic* (RO:0000053), which could be available.

Imported relation property	Ontology	PURL
has_part	RO	http://purl.obolibrary.org/obo/BFO_0000051
adjacent_to	RO	http://purl.obolibrary.org/obo/RO_0002220
aligned_with	RO	http://purl.obolibrary.org/obo/RO_0002001
anterior_to	PATO	http://purl.obolibrary.org/obo/PATO_0001632
bearer_of	RO	http://purl.obolibrary.org/obo/RO_0000053
decreased_in_magnitude_relative_to	PATO	http://purl.obolibrary.org/obo/pato#decreased_in_magnitude_relative_to
external_to	RO	http://purl.obolibrary.org/obo/PATO_0002483
increased_in_magnitude_relative_to	PATO	http://purl.obolibrary.org/obo/pato#increased_in_magnitude_relative_to
internal_to	Not available	
is_approximately_equivalent_to	RO	http://purl.obolibrary.org/obo/RO_0002603
lateral_to	PATO	http://purl.obolibrary.org/obo/PATO_0001193
located_in	PATO	http://purl.obolibrary.org/obo/PATO_0002261
part_of	RO	http://purl.obolibrary.org/obo/BFO_0000050
posterior_to	PATO	http://purl.obolibrary.org/obo/PATO_0001633

Results and discussion

Descriptive templates

Semantic-based morphological descriptions were made for 22 species, whose files in RDF/XML format are available at <https://github.com/luis-gonzalez-m/Lepidocyrtus-RDF-Store>. These descriptions have an average of 592 anatomical terms, of which 260 are referred for the chaetotaxy. The RDF triple (see below) expresses (a) part-whole relation between anatomical entities (Fig. 2) or (b) between anatomical entities and their qualities. The “absences” are represented as (c) negations, following the principle of Open Word Assumption (Antoniou and van Harmelen 2008). The importance of this assumption relies on the fact that it is not possible to make a full morphological description of an organism (Wirkner et al. 2017). The term “absence” usually contains evolutive assumptions in a phylogenetic context to explain changes of homologs or appearance of evolutive novelties (Serenó 2007, Willmann 2016), but at descriptive level it is considered as a quality. The above expressions are equivalent to the EQ format and have the logical structure for the character statements (Serenó 2007) and discussed by Göpel and Wirkner 2018.

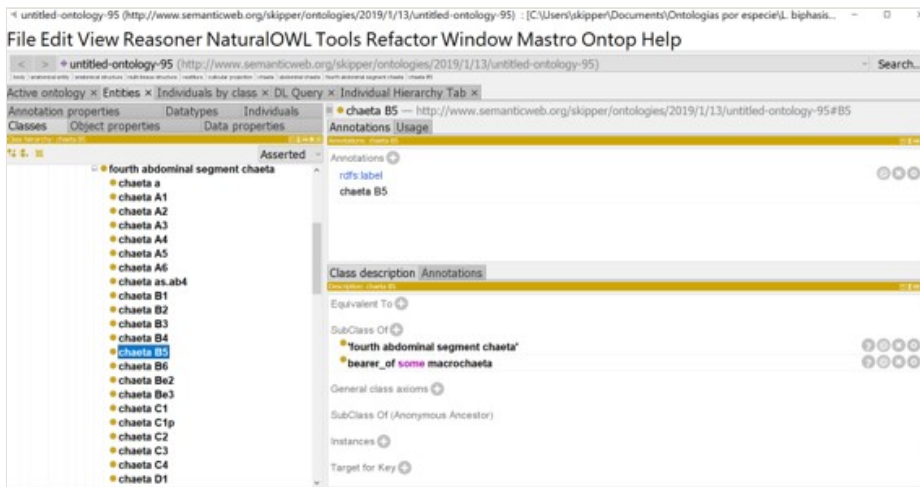


Figure 2. doi

Screenshot of Protégé showing panels used in the class-based method. Left-hand side, some anatomical terms that composed a descriptive template for the species *Lepidocyrtus biphasis* Mari Mutt, 1986. Right-hand side, description for the class “chaeta B5”.

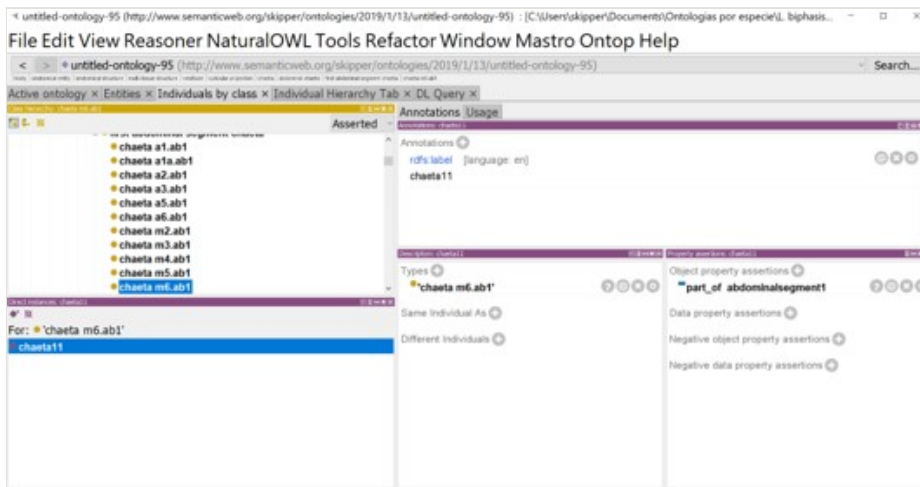


Figure 3. doi

Screenshot of Protégé showing panels used in the class-based method. Left-hand side, the class “chaeta m6.ab1”, the chaeta “m6” located on the abdominal segment 1, and right-hand side, the individual identified by the label “chaeta11” and object property assertion expressed for the species *Lepidocyrtus biphasis* Mari Mutt, 1986.

chaeta am6.ab3 part_of some abdominal segment 3.....(a)

chaeta am6.ab3 bearer_of some macrochaeta.....(b)

chaeta am6.ab3 not (part_of some abdominal segment 3).....(c)

Under the instance-based method, the above is most complex because a second “layer” must be added (Fig. 3), where each class instantiates an individual. For instance, the statement (d) is expressed as:

chaeta145 part_of abdominalsegment3.....(d)

An object property assertion with the provisional labels “chaeta145” and “abdominalsegment3” to name the individuals or parts of organisms perceived in the reality. Some examples of RDF triples and descriptive statements are present in the Table 3.

Table 3. Examples of RDF triple and descriptive statement.	
RDF triple statement	Descriptive statement (natural language)
chaeta A0.h bearer_of some microchaeta	chaeta A0, size: microchaeta
chaeta A0.h bearer_of some macrochaeta	chaeta A0, size: macrochaeta
chaeta A0.h has_part exactly 1 microchaeta	chaeta A0, number: 1
chaeta Ps4 part_of some cephalic chaeta	chaeta Ps4: present
chaeta a3.ab1 part_of some abdominal segment 1 and (anterior_to some chaeta m3.ab1)	chaeta a3, position: anterior to chaeta m3
chaeta a2.ab2 bearer_of some triangular	chaeta a2, shape: triangular
chaeta a2.ab3 aligned_with some chaeta m2.ab3	chaeta a3, alignment: aligned with chaeta m2
chaeta as.ab2 bearer_of some length) and (inheres_in some chaeta m3e.ab2) and (is_approximately_equivalent_to some length)	chaeta as, length: equal to chaeta m3e
chaeta D1p bearer_of some smooth	chaeta D1p, texture: smooth
dental tubercle bearer_of some domed	dental tubercle, curvature: domed

RDF repository

RDF store for biological data is oriented mainly to molecular data with Uniprot (<https://uniprot.org>) and Bio2RDF (<https://bio2rdf.org>), while for morphological data, RDF stores have not been developed. When the RDF is available, the next step is the creation of a semantic web service to put the semantic data on the web (Wollbrett et al. 2013). In SPARQL query the clauses WHERE and SELECT permit to extract information stored in RDF stores. The WHERE clause obtains data out of the dataset and SELECT names what parts of the dataset are pulled (DuCharme 2013). These clauses return RDF triples as the answer for the query.

A list of RDF triple, where the subclass relation is retrieved for the chaetae that composed the cephalic chaeta (Table 4) is obtained through the code in the example 1. Likewise, a list of RDF triple with the displayed relations *bearer_of*, and *posterior_to*, is showed in the

Table 5 through the code in the example 2, being a descriptive block for the species *L. americanus* Cipola & Bellini, 2019:

Table 4. Example 1 (Output)			
ID	SUBJECT	PREDICATE	OBJECT
1	chaeta A1.h	subClassOf	cephalic chaeta
2	chaeta A2.h	subClassOf	cephalic chaeta
3	chaeta A3.h	subClassOf	cephalic chaeta
4	chaeta A4.h	subClassOf	cephalic chaeta
5	chaeta A5.h	subClassOf	cephalic chaeta

Table 5. Example 2 (Output)			
ID	SUBJECT	PREDICATE	OBJECT
1	chaeta A1	bearer_of	microchaeta
2	chaeta A1.h	bearer_of	microchaeta
3	chaeta A3.h	posterior_to	chaeta A5.h
4	chaeta a4.lb	bearer_of	smooth
5	chaeta A5.lb	bearer_of	serrated

Example 1 (Table 4)

```
PREFIX rdfs: <http://www.w3.org/2000/01/rdf-schema#>
PREFIX rdf: <http://www.w3.org/1999/02/22-rdf-syntax-ns#>
PREFIX owl: <http://www.w3.org/2002/07/owl#>
SELECT DISTINCT ?s ?p ?o
WHERE { ?s rdfs:subClassOf ?o;
}
```

Example 2 (Table 5)

```
PREFIX rdfs: <http://www.w3.org/2000/01/rdf-schema#>
PREFIX rdf: <http://www.w3.org/1999/02/22-rdf-syntax-ns#>
PREFIX owl: <http://www.w3.org/2002/07/owl#>
SELECT DISTINCT ?s ?p ?o
WHERE { ?s rdf:type owl:Class ;
  rdfs:subClassOf [ rdf:type owl:Restriction ;
    owl:onProperty ?p ;
    owl:someValuesFrom ?o; ] ;
}
```

```
<!-- http://purl.obolibrary.org/obo/L.americanus.owl#e.lb -->
<owl:Class rdf:about="http://purl.obolibrary.org/obo/L.americanus.owl#e.lb">
```

```

<rdfs:subClassOf rdf:resource="http://purl.obolibrary.org/obo/L.americanus.owl#labial_chaeta"/>
<rdfs:subClassOf>
  <owl:Restriction>
    <owl:onProperty rdf:resource="http://purl.obolibrary.org/obo/L.americanus.owl#bearer_of"/>
    <owl:someValuesFrom rdf:resource="http://purl.obolibrary.org/obo/L.americanus.owl#ciliated"/>
  </owl:Restriction>
</rdfs:subClassOf>
<rdfs:label>chaeta e.lb</rdfs:label>
</owl:Class>

```

The querying of individuals is similar to querying ontology classes in Fuseki, but the main difference is the querying of individuals or objects in named graphs. For example, the SPARQL query to access named graph follows the general syntax (Vogt 2019):

```

SELECT*
WHERE {
GRAPH <123> # 123 is the named graph
}

```

Semantic-based methods for the documenting of morphological data have gained interest in recent years, motivated by the potential application in phylogenetics (Vogt et al. 2009, Balhoff et al. 2013, Tarasov 2019). This application involves the coding and extraction of the character and character states, and recognition of historical and serial homologies across anatomical entities employing computational reasoning (Cui 2012, Mabee et al. 2019). However, the development of computational tools for the generation of phenotypic annotations from RDF triple has received little attention. There are several limitations to its implementation: 1. these methods require datasets such as anatomic ontology, which are not available for most taxonomic groups, 2. these methods are unknown or less preferable than morphological descriptions in natural language, 3. the attention of character statement coding is relevant in comparison with the expression of descriptive statements, which are framed within taxonomic tradition or publication requirements, and 4. technical limitations. In this study, for example, the descriptive statements were generated manually, but other options such as the declaration of RDF triples in spreadsheets and importing to ontology editor may reduce the time-consuming demand. Also, the extraction of phenotypic annotations from texts has important advances (Wood et al. 2003, Thessen et al. 2012, Dahdul et al. 2015, Cui 2012). For instance, the incorporation of Natural Language Processing (NLP) during the extraction of phenotype data sets could increase the efficiency of this task (Dahdul et al. 2015). However, the generation of phenotypic annotations from RDF triple, employing techniques of Natural Language Generation (NLG) has not been explored or applied in morphological descriptions.

The instance-based method, “semantic anatomy instance”, is most complex when the Abox assertions are built with Protégé because individuals by class need to be specified, resulting in a template composed of two layers: ontology class and instanced objects. Recently proto.morphdbase.de incorporates instance-based methods, promising to increase the use of semantic-based tools during morphological descriptions. This application is complementary to other tools that integrate morphological data with

multimedia representations, for instance, MorphoNet (Leggio et al. 2019), MorphoBank (O’Leary and Kaufman 2011), and Phenotools (Eliason et al. 2019). Proto.morphdbase.de has two principal components: semantic graph generated for each organism and accessibility to metadata supplied from scientific publication and regulated by FAIR Guiding Principles to manage stored data (Wilkinson et al. 2016, Vogt 2019).

Currently, there is an imperative need to document biological diversity, which implies the use of computational tools for the processing of different data generated in the biology domain. Unfortunately, this urgency is directed mostly to the storing and processing of molecular data, while the morphology is continually displaced. Morphological descriptions are a useful source of data but due to their nature and complexity requires “creative” solutions, new automatic or semi-automatic methods that permit the interchange between natural language employed commonly in published morphological descriptions and RDF triple syntax. The use of ontologies uncovers the subtle process between morphological data (expressed by RDF triple statement) to character statement, where the character state (properties) arises from the comparison between species and before the building of character matrix.

Initiatives about morphological descriptions that employ standardized languages are not new (Dallwitz 1984, Paterson et al. 2004, Cui 2008). Recently, Cui et al. 2020 developed an author-driven method, where the author or taxonomist defines what ontological classes are necessary for their descriptions. These classes are included by the software engineer in an ontology, this means, the author is immersed in the building of the ontology, expressing semantic relations and conflicts in the use of terminology. Although the ability on the management of RDF syntax during phenotypic annotation could be cumbersome, the use of structured syntax in morphological descriptions contributes to making them available to computational analysis.

However, the taxonomic tradition has an important weight in the language employed during morphological descriptions and is taxon-dependent. It is necessary to reconcile the needs of taxonomists and friendly tools to incorporate these methods. It is not the goal to evaluate the multiple RDF store available, which differs in properties as storage size, querying time, and applicability (Frey et al. 2019). In this study, the building of descriptive templates is suggested, being an intuitive approach to manage RDF triplet and its storage. The semantic-based methods are a tool that increases the management, processing, and comparison of morphological data that could contribute to its integration in the taxonomic work with a friendly computational approach.

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Conflicts of interest

None

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