

Analysing the extent of cell type information present in Wikidata: A case study on PanglaoDB

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Abstract

[Wikidata](#), a freely editable knowledge graph database, presents a great opportunity for the integration of biomedical knowledge, its well thought linked data model can improve significantly the handling and distribution of scientific information. On the other hand, Wikidata is still lacking in various aspects, in particular to what pertains to cell type information. This study aims to analyse how cell type knowledge is currently modelled in Wikidata and how it differs from other types of biological information, using, as a reference point, metadata from the well known single cell RNA sequencing database, [PanglaoDB](#).

Keywords: wikidata, knowledge graph, cell type, ontology.

Introduction

Wikidata

Wikidata [1] is an open, freely editable, knowledge graph database within the semantic web [2] that stores knowledge across a multitude of domains, such as arts, history, chemistry and biology, using an item-property-value linked data model (Figure 1). It is easy to use and edit, by both humans and machines, with a rich web user interface and wrapper packages available in common programming languages such as R and Python. All the data within Wikidata is linked and inherently public domain, thus, it presents a great opportunity to make scientific data more FAIR (Findable, accessible, interoperable and reusable), as well as provides the necessary tools to curate and develop ontologies.

hepatocyte

(Q827450)

Item Identifier

Item Label

liver cell type

Hepatocyte | hepatocytes

edit

In more languages

Configure

Language	Label	Description	Alternative labels
English	hepatocyte	liver cell type	Hepatocyte hepatocytes
Portuguese	Hepatócito	tipo celular	
German	Hepatozyt	Hauptzelltyp der Leber	Hepatocyt Leberepithelzelle Hepatozyten Leberzelle
Italian	Epatocito	No description defined	Epatociti Epatocita

All entered languages

Statements

Statement

instance of

Property

cell type

Value

edit

0 references

+ add reference

+ add value

Figure 1: Wikidata item example, showing item hepatocyte (Q827450), the labels change according to the user's language, but each item has a universal identifier, called QID.

Several advances towards biological data integration and biological data analysis in Wikidata have been made before, yielding positive results [3] [4] and showcasing it's potential for bioinformatics-related analyses, such as drug repurposing and ID conversion [4]. Wikidata has been proposed as a unified base to gather and distribute biomedical knowledge, with more than 50 000 human gene items indexed and hundreds of biomedical-related properties [5]. However, as of August 2020, cell type information is still very scarce, with only 264 items being categorized as "instances of cell types (Q189118)" (<https://w.wiki/b2w>), of those, only nine have a "Cell Ontology ID"[6] (P7963) associated, and most have a varying amount of statements (Table 1). As an additional problem, there are also 23 items being categorized as "instances of cell (Q7868)" (<https://w.wiki/b2x>), illustrating the absence of any formal data model.

Table 1: As of August 2020, Wikidata items regarding cell types have a varying amount of information, with most having very few statements.

Cell type Item	Number of statements
red blood cell (Q37187)	48
myocyte (Q428914)	18
mesenchymal cell (Q66568500)	2

PanglaoDB

PanglaoDB [7] [8] is a public database that contains data and metadata on hundreds of single-cell RNA sequencing experiments, providing extensive information on cell types, genes and tissues, as well as manually and community curated cell type markers (Tables 2 and 3). It also provides a rich web user interface for easy data acquisition, including database dumps for bulk downloads.

Table 2: Database statistics for each species in PanglaoDB, as of 31st of August, 2020.

	Mus musculus	Homo sapiens
Samples	1063	305
Tissues	184	74
Cells	4,459,768	1,126,580
Cell Clusters	8,651	1,748

Table 3: Metadata statistics for PanglaoDB, gathered from their [last update on August, 2019](#).

	Number
Cell types	215 (uniquely named)
Tissues	240 (+6 germ layers)
Organs	29
Species	2 (Homo sapiens and Mus musculus)
Genes	110292

Methodology

Data acquisition

Gene data from Wikidata was acquired using the Wikidata Query Service [9] - <https://w.wiki/bWc> for *Homo sapiens* genes and <https://w.wiki/bWe> for *Mus musculus* genes.

Data from PanglaoDB was acquired through their metadata database dump repository[10].

All data used was handled using the Pandas[11] library, with the Seaborn[12] and Matplotlib[13] libraries being used for plotting.

Reconciliation and matching

The metadata from PanglaoDB on cell types, tissues (including germ layers) and organs was matched to Wikidata items using the reconciler[14] library, further matching was done using a custom stemming function on the item labels, via PorterStemmer from the NLTK library [15]. Matches were considered perfect if the reconciliation service or the stemming function returned a value of “match” equals to “True”. Matches were manually analysed for false matches, such as items with same labels but used for different concepts.

Gene data was matched manually using a Pandas [11] inner merge, since both data sources contained identifiers, which should be the same.

Item quality assessment

Wikidata items were assessed for their quality by their number of statements, which were acquired using a custom wrapper on the MediaWiki API [16] and, in the case of gene data, via Wikidata’s own query service, as stated in the Data acquisition section.

Furthermore, items were also assessed by the presence of external identifiers - all of which are Wikidata properties: Ensembl Gene[17] (P594) and Entrez Gene[18] (P351) IDs for genes, Cell Ontology[6] (P7963) IDs for cell types and Uberon[19] (P1554) IDs for organs and tissues.

Results

Entities from PanglaoDB, that is, cell types, genes, tissue types and organs, were matched with Wikidata items, matching summary can be seen on Table 1. Only *Homo sapiens* genes and Organs reconciled more than 50%. In the case of genes, this is probably due to the Gene Wiki initiative [20], a long-running project to improve biological information in Wikipedia and its sister-projects, including Wikidata. This is further illustrated by Figure 2, in which we can see that all *Mus musculus* gene items - and nearly all *Homo sapiens* items - analysed had the Entrez ID alternative identifier present - Most of the data from the Gene Wiki project came from NCBI, creator and maintainer of Entrez.

Nevertheless, there are still many gene items without an “Ensembl Gene ID” property, showcasing the need for further work in migrating this important source of information.

In the case of Organ data, there was a high number of matches both due to the fact that there were only a few number of items, but also since most Organ entities have Wikipedia pages, that are, therefore, cross-linked using Wikidata, requiring the creation of these items.

Regarding alternative identifiers, what was observed for genes cannot be said for histological entities, while there is significant progress in integrating UBERON IDs, there is near to no items with a Cell Ontology ID property (Figure 1).

Table 1: Summary of the matched entities from PanglaoDB.

	# of total items	# of unique matches	# of matched items	% of total items that were matched	% of matches that were perfect	% of matches that don't have P31
Cells	215	81	85	37.67%	38.82%	55.29%
Tissues	246	79	87	32.11%	62.06%	37.93%
Organs	29	22	30	75.86%	53.33%	46.66%
Human Genes	58216	35423	35427	60.84%	NA	NA
Mouse Genes	53793	25124	25127	46.70%	NA	NA

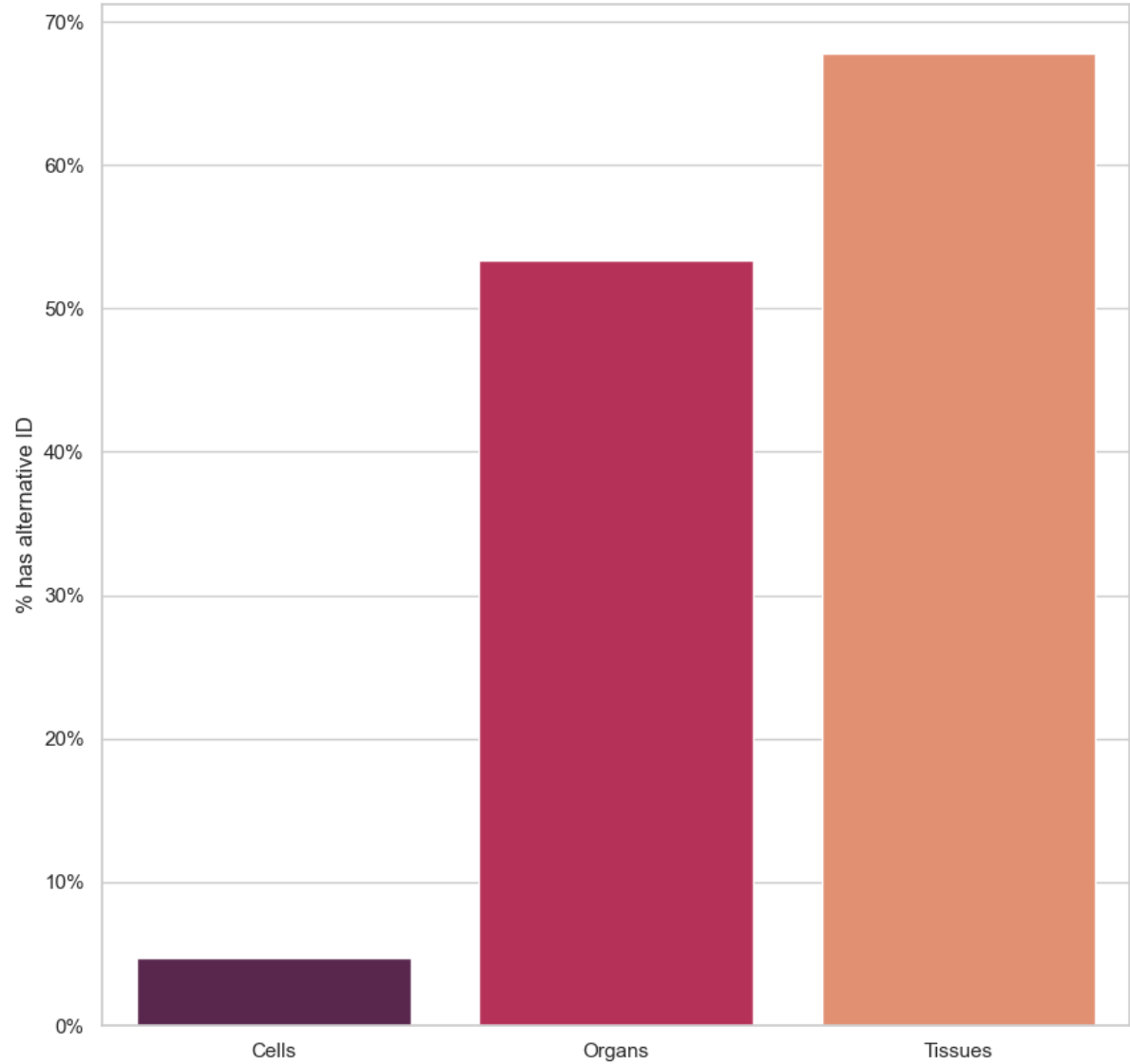


Figure 1: Percentage of matched histological items that had alternative identifiers, UBERON IDs for Tissues and Organs, Cell Ontology IDs for Cell types.

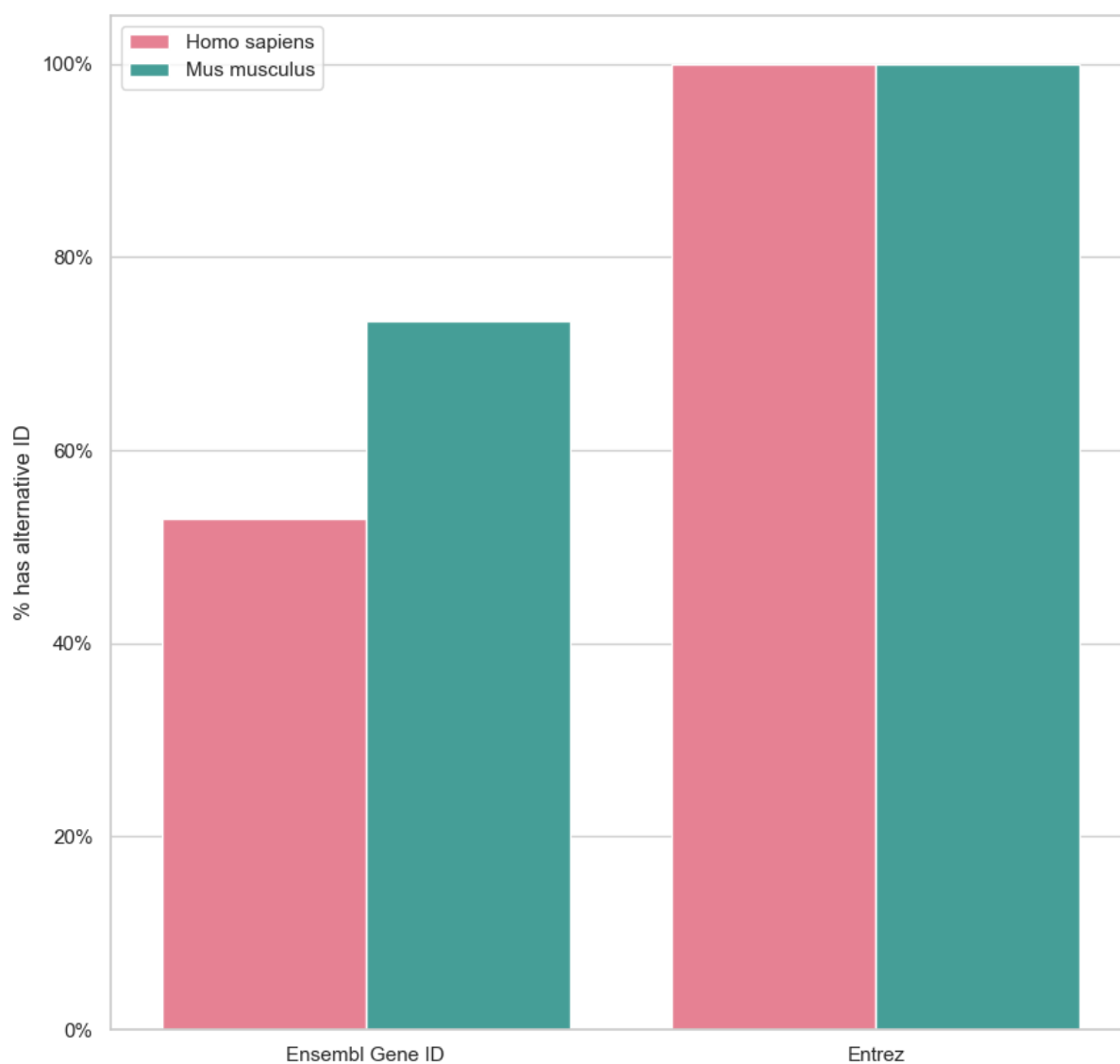


Figure 2: Percentage of matched gene items that had alternative identifiers, Entrez ID and Ensembl Gene ID, divided by species.

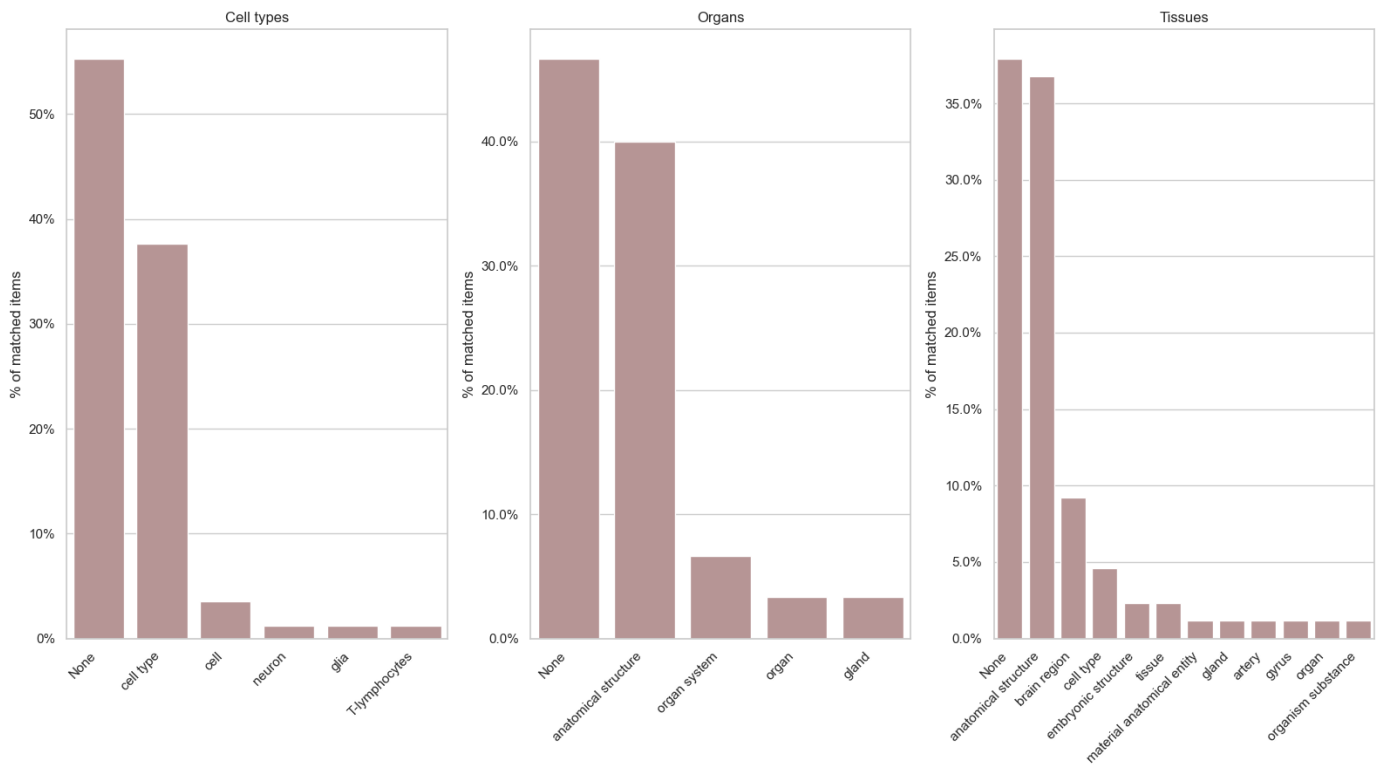


Figure 3: Percentage of reconciled entities, divided by which item type they belong to. Most reconciled items don't count with the P17 property.

A significant proportion of the matches we could acquire for histological data didn't contain in their data model an "instance of" (P17) property, this illustrates an extremely concerning fact: Although we could still match around 30 percent of the data - in the case of Cell types and Tissues - this data was probably "low-quality", that is, hard to find and even harder to obtain insights from, we can affirm this since the P17 property is the basis for most items in Wikidata, it's the most intuitive way to perform queries against their database and to annotate their items.

Furthermore, there is a significant disparity between histological data and gene data: while we could only match around 37% of Cell types from PanglaoDB, and of those 55% didn't have P17, we matched 60% of *Homo sapiens* genes, and all of them had P17. This disparity is not clearly shown when looking exclusively at the number of statements for these items (Figures 4 and 5), but it shows there is still a great amount of missing information for biological data, in particular in regards to cell types.

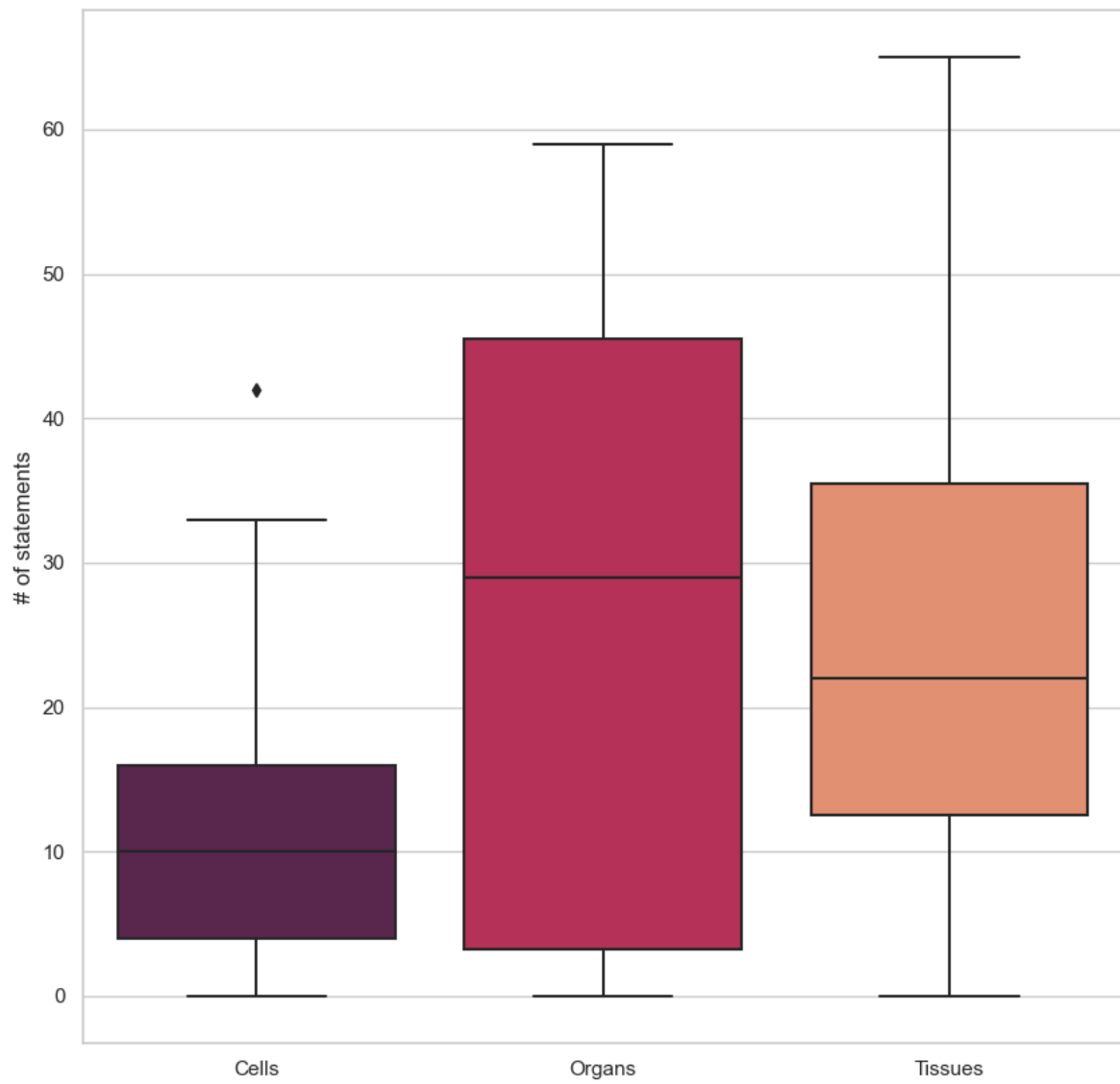


Figure 4: The distribution of the number of statements of the matched histological entities. Cell types performed the lowest.

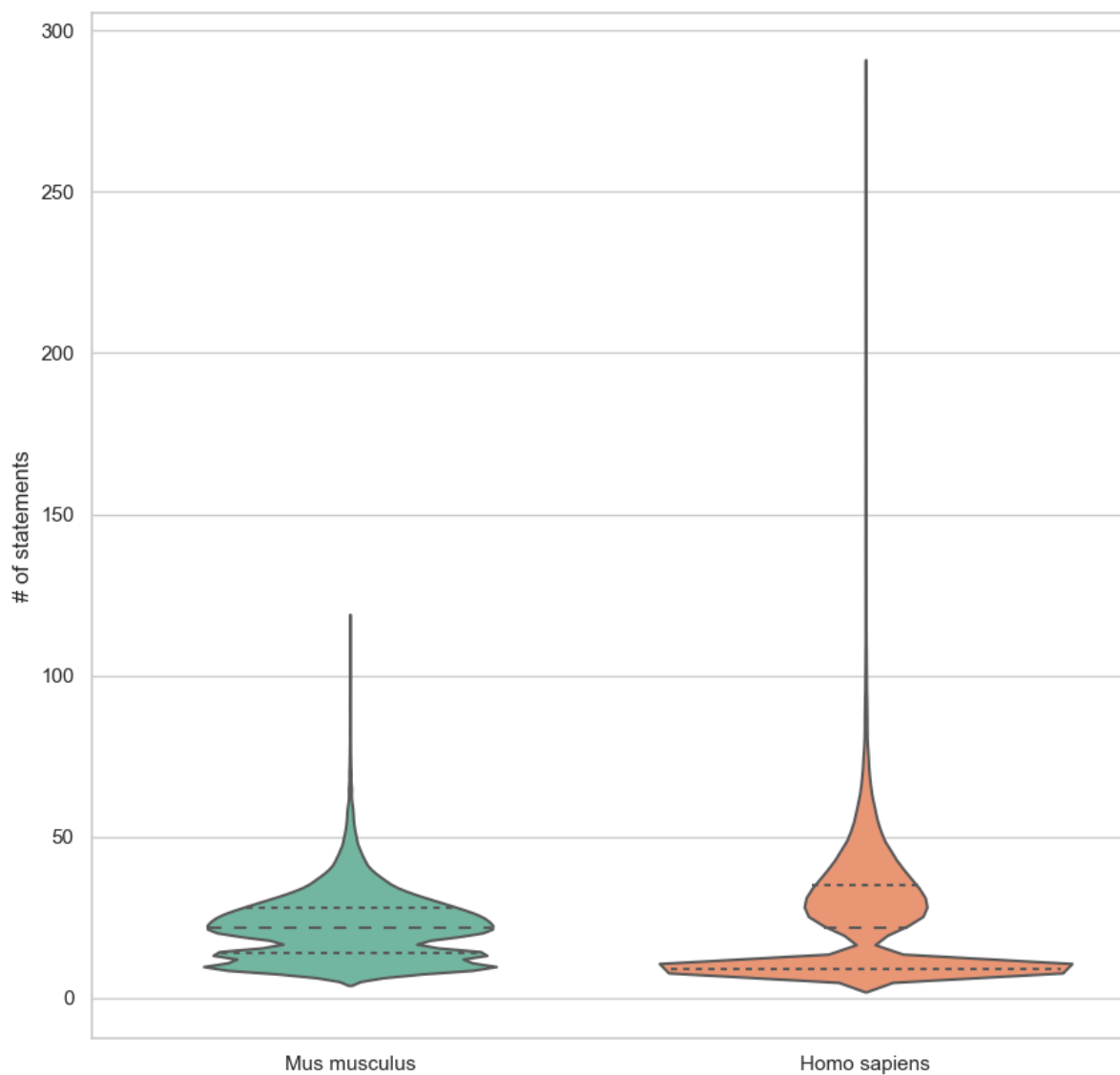


Figure 5: The distribution of the number of statements for matched gene items, divided by species.

General Ideas

Temporary file containing ideas for the project. Interesting references and concepts.

med2rdf[[21](#)] is a project to migrate biomedical knowledge bases to RDF format, facilitating integration with the semantic web.

15 years ago, in the original Cell Ontology paper, they mention the idea to integrate their knowledge with gene expression databases, something not done as far as we know [[22](#)]

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