Breakout Session 1: Track B

Improving FAIRness and AI/ML Readiness of Bioconductor Data Resources

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Improving FAIRness and AI/ML readiness of Bioconductor data resources

Cancer Genomics: Integrative and Scalable Solutions in R/Bioconductor

2024.03.27 Sehyun Oh, PhD (Speaker) Levi Waldron, PhD (PI)



GRADUATE SCHOOL OF PUBLIC HEALTH & HEALTH POLICY

Challenge



Data class from *Bioconductor*

SummarizedExperiment . Samples Samples (Columns) colData(se) se[, se\$dex == "trt"] Features (Rows) metadata(se) rowRanges(se) metadata(se)\$modelFormula rowData(se) assays(se) subsetByOverlaps(se, roi) assay(se, n = 2) assay(subsetByOverlaps(se, roi)) assay(se[, se\$dex == "trt"])





Our project aims to build the first large-scale, platform-independent, curated, ML-ready data repository for diverse omics and associated non-omics data.

Two target Bioconductor data packages:

1) *curatedMetagenomicData* package (human microbiome)

 \rightarrow 142 attributes collected on 22,599 samples from 93 studies

2) *cBioPortalData* packages (cancer genomics):

 \rightarrow 3,733 attributes collected on 189,439 samples from 375 studies

Methods

- 1. [Data export] datasets in R objects into a language-agnostic format
 - hdf5 for assays, csv for metadata, json for manifest
 - New Aim for the U24 renewal
- 1. [Metadata harmonization] sample-level metadata curation/harmonization
 - Major variables: disease/condition, treatment, age, race/ethnicity, sex
 - Compress, consolidate, complete, and correct
- 1. [Resource distribution] data/harmonized metadata in language-agnostic format
 - Public (e.g., Zenodo) and commercial (e.g., GCP, Azure) for storage
 - User-facing website and API
- 1. [Use case] use cases of AI/ML-application
 - Cross-studies analysis
 - Analyses In different languages

Metadata Harmonization



Improvement in data harmonization

- 1. **Compression**: the number of original attributes merged/combined into a new curated attribute
- 2. Consolidation: the number of unique values for a given attribute
- 3. Completeness: the proportion of non-missing values
- *4.* **Correction rate**: the proportion of the values updated during the curation



curatedMetagenomicData





• original o curated

cMD - Correction



- 12 curated attributes contain completely new values
- control;disease;target_condition



cBioPortalData





Curation Tool

- Current features: automatically fetch info from SRA, validation to the schema, use ontologies
- Ongoing features: version control, automated updates, etc.

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2			Υ							
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4		v.	v			v	v			



OmicsMLRepoR

- An R software package for users to access and manipulate metadata
- Inputs are all in non-R-specific formats, such as csv and tsv
- Major functionalities:
 - 1. Manipulate metadata tables: get long/short/wide/narrow forms of metadata table
 - 2. Metadata search through ontology tree: *searchMetadata* and *ontoTraverse*





- Low quality information
 - not covered by the existing ontologies
 - arbitrary abbreviations
 - not self-sufficient information
- Low level or metadata harmonization even some large, widely-used, curated public databases
- Maintain the quality of incoming datasets



Future Plans

- 1. Release the harmonized version of metadata for *curatedMetagenomicData cBioPortalData*
- 2. Release OmicsMLRepoR package
- 3. Automated metadata harmonization tool
- 4. Implement *DataConductor*, a unified resource for enhanced data accessibility and integration, which will be featured by:
 - Data in language-agnostic formats
 - Metadata JSON schema
 - User-friendly API and faceted data portal





Summary and Conclusions

- 1. We identified very heterogeneous, sparse, and inaccurate metadata from large, widely used public omics data resources and even from 'curated' data resource.
- 2. We did successful metadata harmonization and curation of two major Bioconductor resources on metagenomics and cancer genomics data.
- 3. We developed the user-friendly metadata schema based on the already collected information.
- 4. We developed the manual metadata curation framework and supporting tools.
- 5. Preliminary work under this Supplementary contributes to develop a new Aim for our renewal application of the parent U24.
- 6. It is critical to continuously improve and maintain the metadata quality to support diverse use cases of omics data, including AI/ML application.



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<u>Team</u>

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