



a python utility to study and analyse cancer-associated metabolites using knowledge graphs

- Difficult to navigate, abundant metabolomics data
- Need to automate analysis
- A search-engine-like interface



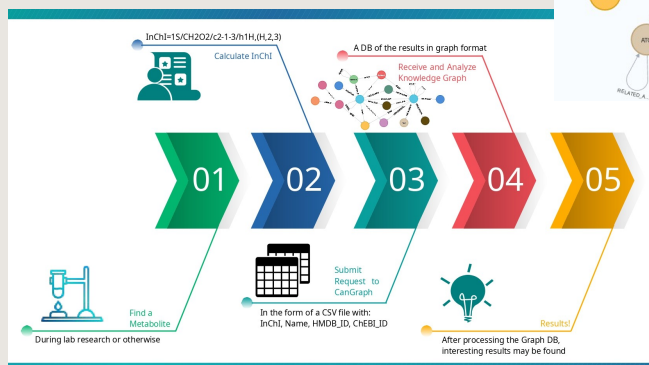
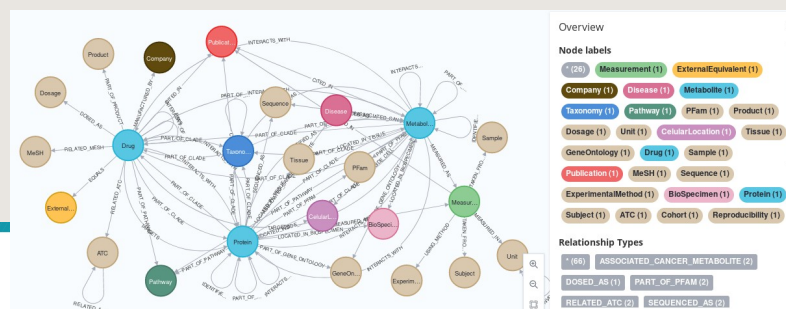
Seven top-notch databases



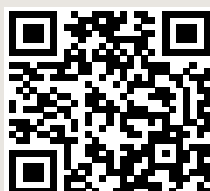
Pulled together using



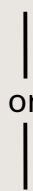
And unified using a graph schema



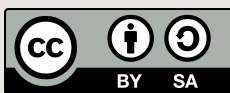
So that newly discovered metabolites can be more easily identified!



See the official website



Read the full publication on my blog



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