

# Intégration de données hétérogènes Bases de données non structurées

Master 2

Bioinformatique et Biologie des Systèmes

# Programme de la journée

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- Matin
  - Apperçu sur
    - Big data, NoSQL, base de données de graphes, neo4j
  - Prise en main de neo4j
    - Construction d'une base de connaissances (graphe) sur les gènes/protéines d'un organisme, *Escherichia coli*
    - données d'expression
    - données d'interaction protéine-protéine
    - données phylogénomique
    - annotations Gene Ontology
- Après midi
  - Concepts
    - caractérisation d'un ensemble
  - Application
    - Annotations GO sur-représentées parmi les ensembles de gènes/protéines présentant à la fois
      - coexpression,
      - interaction protéine-protéine,
      - conservation du voisinage génomique

# Tendances et observations

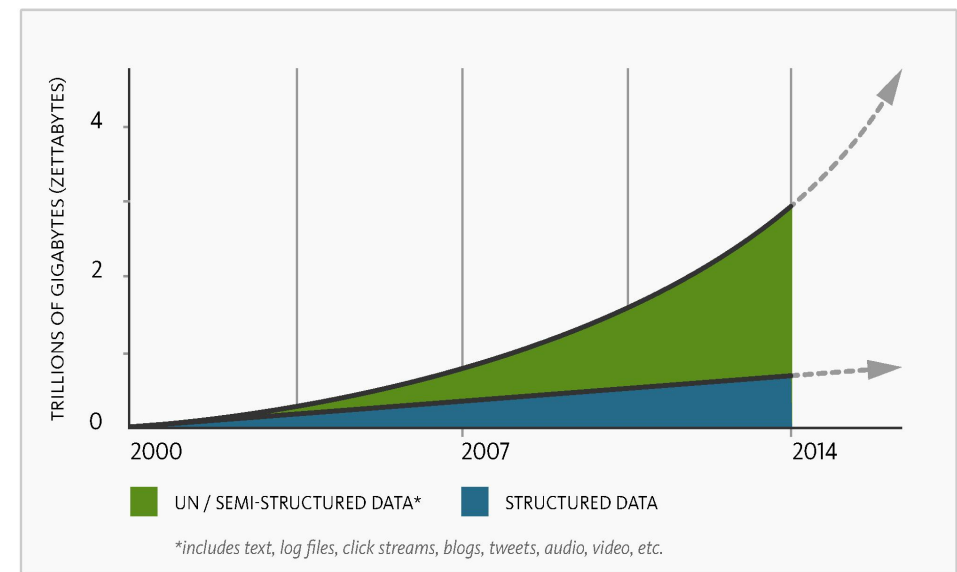
## Big data

- “Big data is high **volume**, high **velocity**, and/or high **variety** information assets that require new forms of processing to enable enhanced decision making, insight discovery and process optimization.” (Gartner, 2012)

## Données non structurées

- Pareja-Tobes et al. 2015 : "Sometimes analysis can be scheduled (annotating a genome for a research project), while in other cases it is inherently impossible to even predict when you will need to run them (think of analyzing data coming from an outbreak). The resources needed or their characteristics vary wildly depending on the specific analysis. We are thus in a context where in general **we do not know in advance what we need to access**, how fast, when, and how."

## NoSQL Databases



# NoSQL databases

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## What is “NoSQL”?

- term used in late 90s for a different type of technology:  
Carlo Strozzi: [http://www.strozzi.it/cgi-bin/CSA/tw7/l/en\\_US/NoSQL/](http://www.strozzi.it/cgi-bin/CSA/tw7/l/en_US/NoSQL/)
- “Not Only SQL”? (but many RDBMS are also “not just SQL”)
- “NoSQL is an accidental term with no precise definition”. first used at an informal meetup in 2009 in San Francisco (presentations from Voldemort, Cassandra, Dynomite, HBase, Hypertable, CouchDB, and MongoDB)

## NoSQL: Database technologies that are (mostly)

- Not using the relational model (nor the SQL language)
- Designed to run on **large clusters** (horizontally scalable)
- **No schema** - fields can be freely added to any record
- Open source
  - Based on the needs of 21st century web estates

## Other characteristics (often true):

- easy replication support (fault-tolerance, query efficiency)
- simple API
- eventually consistent (not ACID)

### Flexible scalability

- **horizontal scalability** instead of vertical

### Dynamic schema of data

- different levels of flexibility for different types of DB

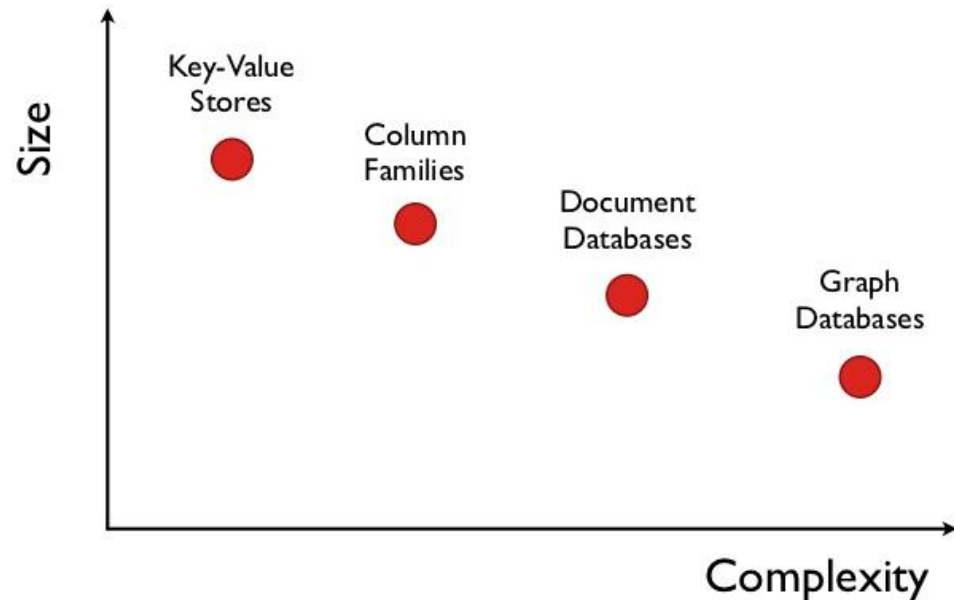
### Efficient reading

- spend more time storing the data, but read fast
- keep relevant information together

### Cost saving

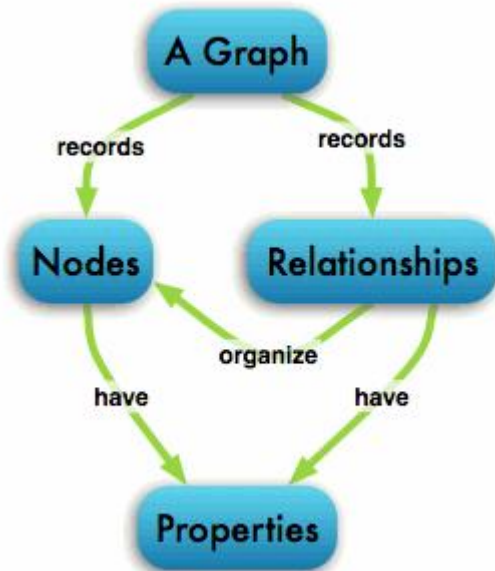
- designed to run on commodity hardware
- typically open-source (with a support from a company)

- MapReduce programming model
  - running over a distributed file system
- **Key-value stores**
- **Document databases**
- **Column-family stores**
- **Graph databases**



source: <http://www.slideshare.net/emileifrem/nosql-east-a-nosql-overview-and-the-benefits-of-graph-databases>

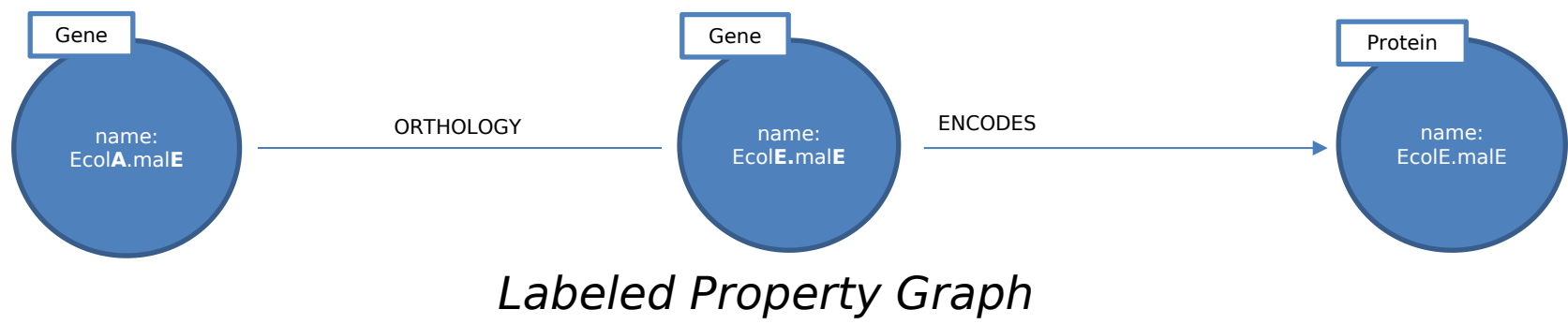
- Open source graph database
  - The most popular
- Initial release: 2007
- Written in: Java
- OS: cross-platform
- Stores data as nodes connected by directed, typed relationships
  - With properties on both
  - Called the “property graph”



- reliable – with full ACID transactions
- durable and fast – disk-based, native storage engine
- scalable – up to several billion nodes/relationships/properties
- highly-available – when distributed (replicated)
- **expressive** – powerful, human readable graph query language
- fast – powerful traversal framework
- **embeddable** - in Java program
- **simple** – accessible by REST interface & Java API

# Base de données orientée graphe (*Graph databases*)

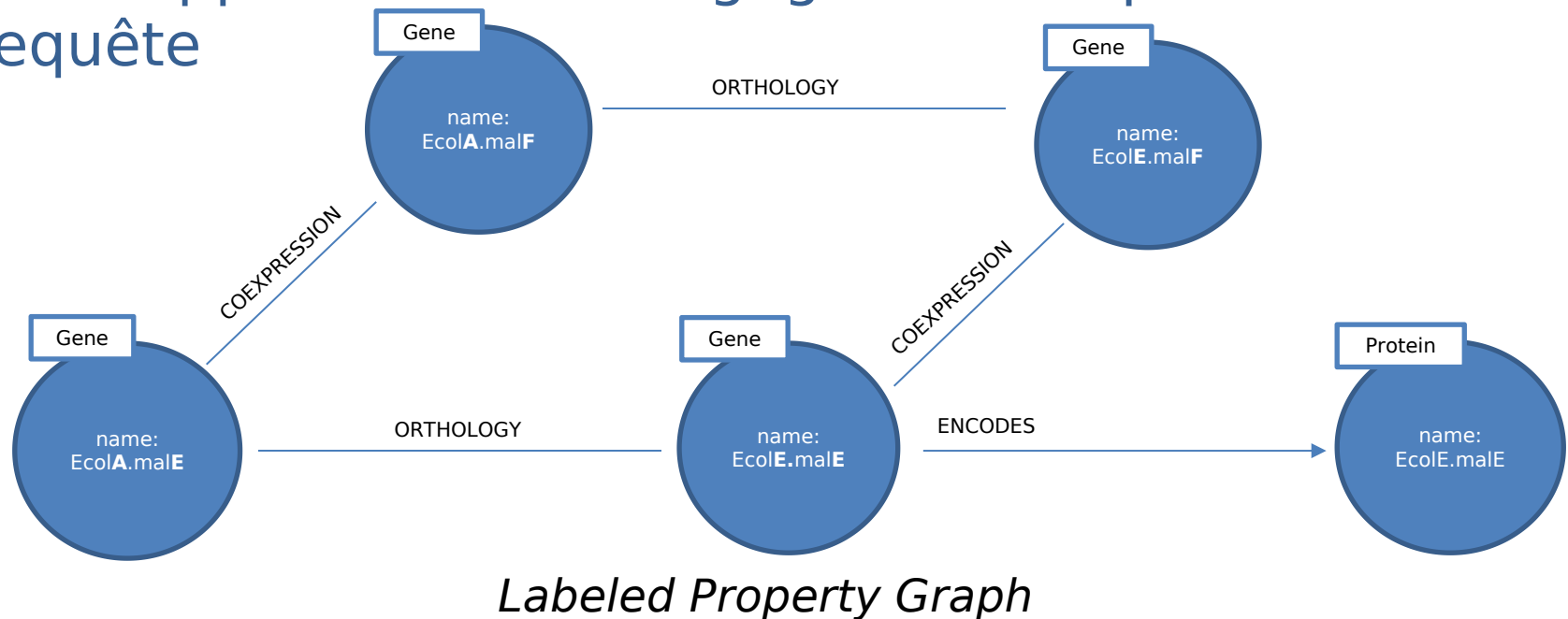
- Principes
  - Représenter les données en tant qu'objets reliés par des relations
  - Chaque objet ou relation peut avoir des attributs qui lui sont propres
  - Développement d'un langage de manipulation et de requête
- Neo4j : Labeled property graph



(EcolA.malE:Gene) - [ :ORTHOLOGY ] - (Ecole.malE:Gene) - [ :ENCODES ] ->(Ecole.malE:Protein)

# Base de données orientée graphe (*Graph databases*)

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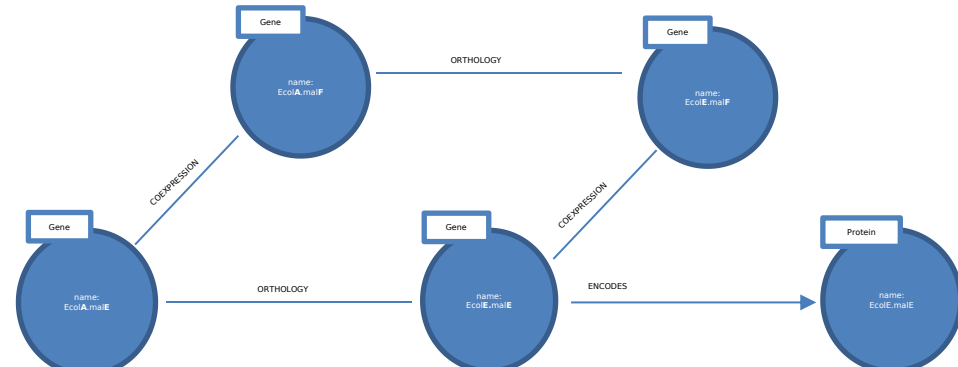
```
(Ecole.malE:Gene) - [:COEXPRESSION] - (EcolA.malF) - [:ORTHOLOGY] -  

(EcolA.malF:Gene) - [:COEXPRESSION] - (EcolA.malE:Gene) - [:ORTHOLOGY] - (Ecole.malE:Gene)  

- [:ENCODES] -> (EcolE.malE:Protein)
```

## Labeled property graph

Un graphe avec propriétés étiquetées est constitué de sommets, relations, propriétés et étiquettes :



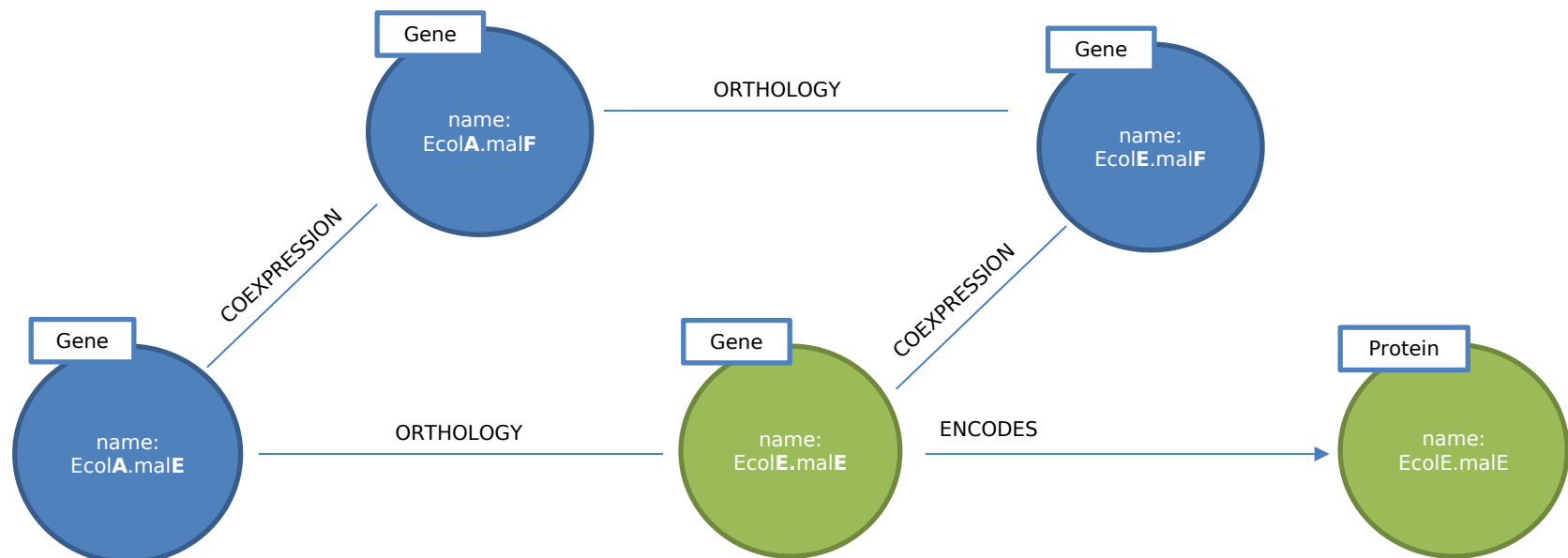
- Propriétés des sommets : de type clé/valeur
- Étiquettes des sommets : une ou plusieurs afin de les regrouper (Gene, Protein)
- Relations : orientées, peuvent avoir des propriétés comme les sommets.

## Langage de requête, exemple : Cypher

```
MATCH (g:Gene) - [:ENCODES] -> (p:Protein)  
WHERE g.name='Ecole.E.malE'  
RETURN g, p
```

ou

```
MATCH (g:Gene {name: 'Ecole.E.malE'}) - [:ENCODES] -> (p:Protein)  
RETURN g, p
```



## Cypher Language

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- Neo4j graph query language
  - For querying and updating
- Declarative – we say **what** we want
  - Not how to get it
  - Not necessary to express traversals
- Human-readable
- Inspired by SQL and SPARQL
- *Still growing* = syntax changes are often

<http://neo4j.com/docs/stable/cypher-query-lang.html>

- **MATCH**: The graph **pattern** to match
- **WHERE**: **Filtering** criteria
- **RETURN**: What to return
- **CREATE/MERGE**: Create/Update nodes and relationships.
- **DELETE**: Remove nodes, relationships, properties
- **SET**: Set values to properties
- **WITH**: Divides a query into multiple parts

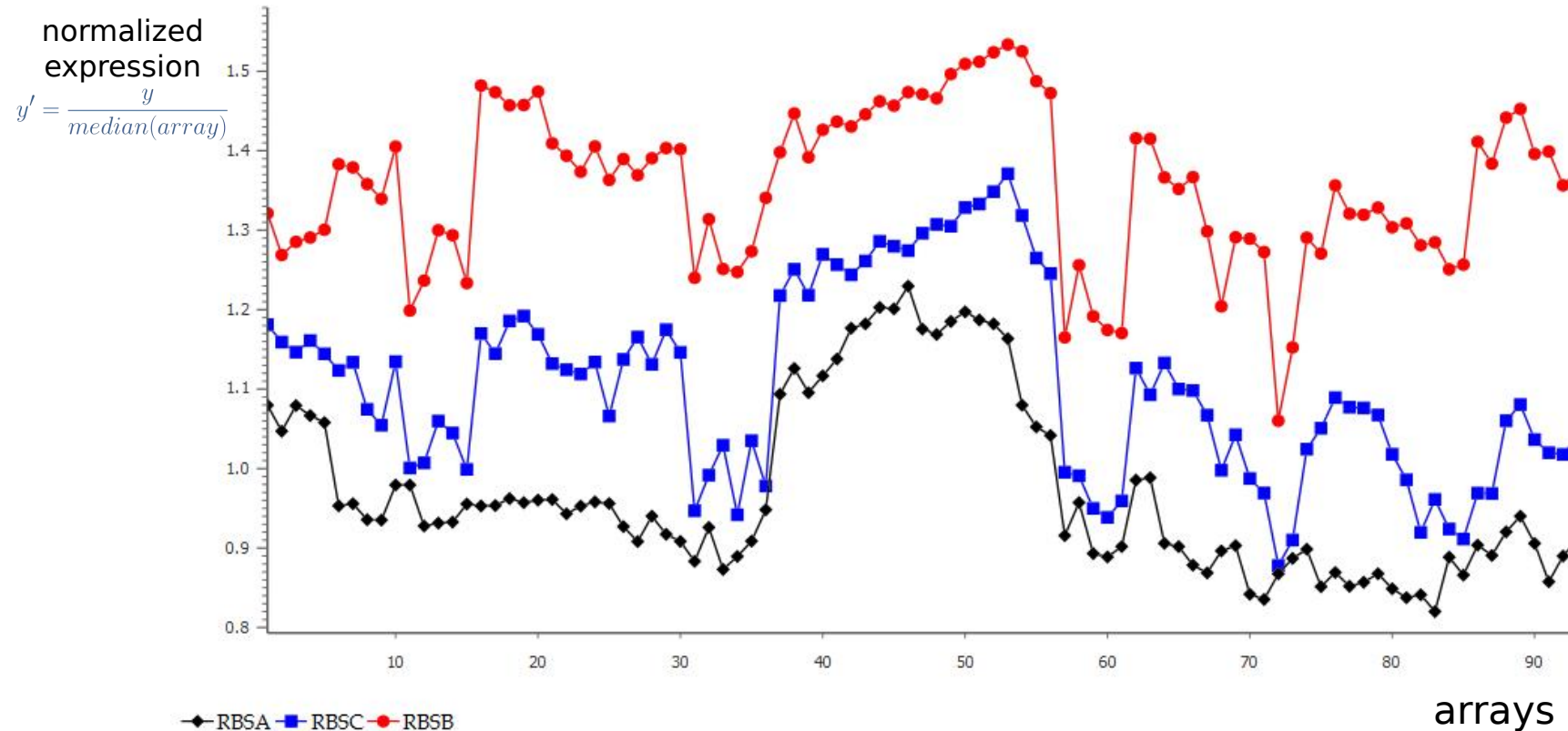
## Access to Neo4j

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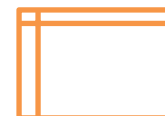
- Embedded database in Java system
- Language-specific connectors
  - Libraries to connect to a running Neo4j server (python, R, ...)
- Cypher query language
  - Standard language to query graph data
- HTTP REST API
- etc.

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# Gene expression

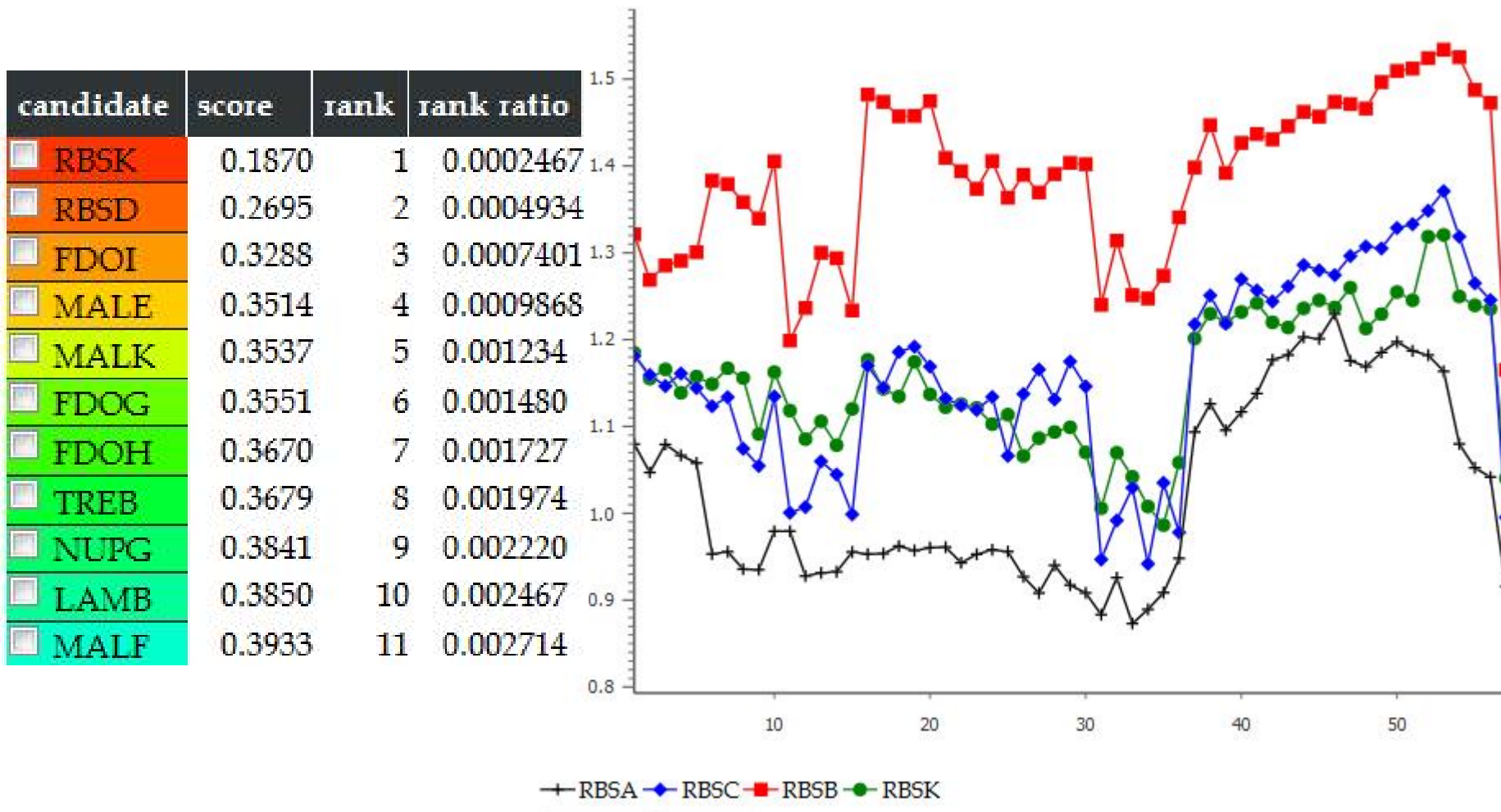


- a gene: set of expression values in various experimental conditions
- a pair of genes: dissimilarity index based on Pearson's correlation coefficient
- score : average dissimilarity



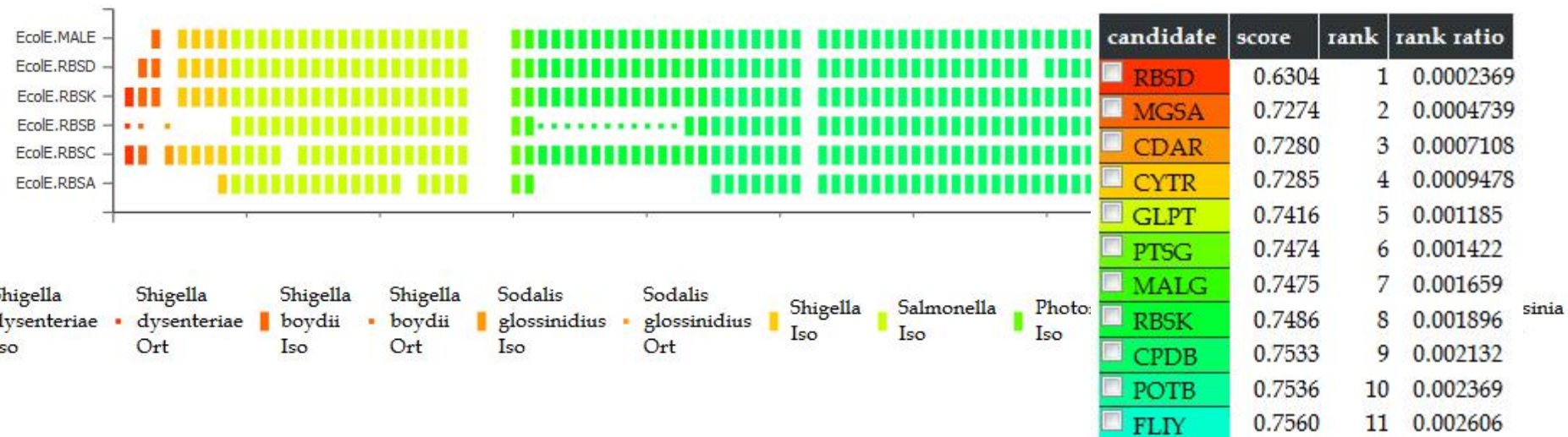
gene pairwise dissimilarity matrix

- training: rbsA, rbsB, rbsC in *E. coli* K-12

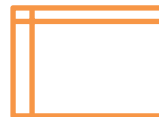


# Phylogenetic profiles

training: rbsA, rbsB, rbsC



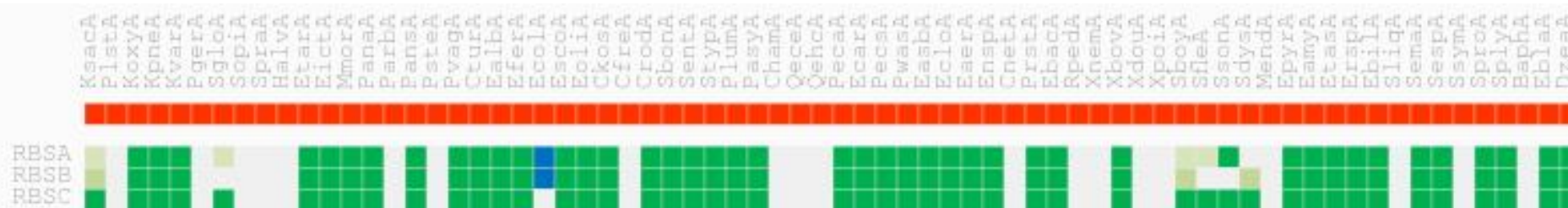
- a gene: presence/absence of orthologs 1:1 in other genomes
- pair of genes: dissimilarity index based on the Jaccard index
- score: average dissimilarity



gene pairwise dissimilarity matrix

## Phylogenetic data

- Phylogenetic profiles



- gene pairwise distance matrix computation
    - Hypothesis: genes located near each other in a set of genomes are likely to be functionaly related
    - $g_1'$  and  $g_2'$  orthologs 1:1 of gene<sub>1</sub> and gene<sub>2</sub> in another genome  $i$
    - Probability that the distance  $D_i$  is smaller than the observed distance  $d_i$

$$p_i = \Pr(D_i \leq d_i) = \frac{2d_i}{N_i - 1}$$

- For a set of  $M$  other genomes

$$d = \Pr(D_1 \leq d_1, \dots, D_M \leq d_M) = \prod_{i=1}^M p_i$$

- $M$  depends on the pair of genes considered
  - $d$  not comparable between genes (e.g.  $0.1^6 = 10^{-6}$  vs.  $0.5^{20} = 9.5 \cdot 10^{-7}$ )
  - normalization: log transformation, z-score, average of distance matrix and its transpose

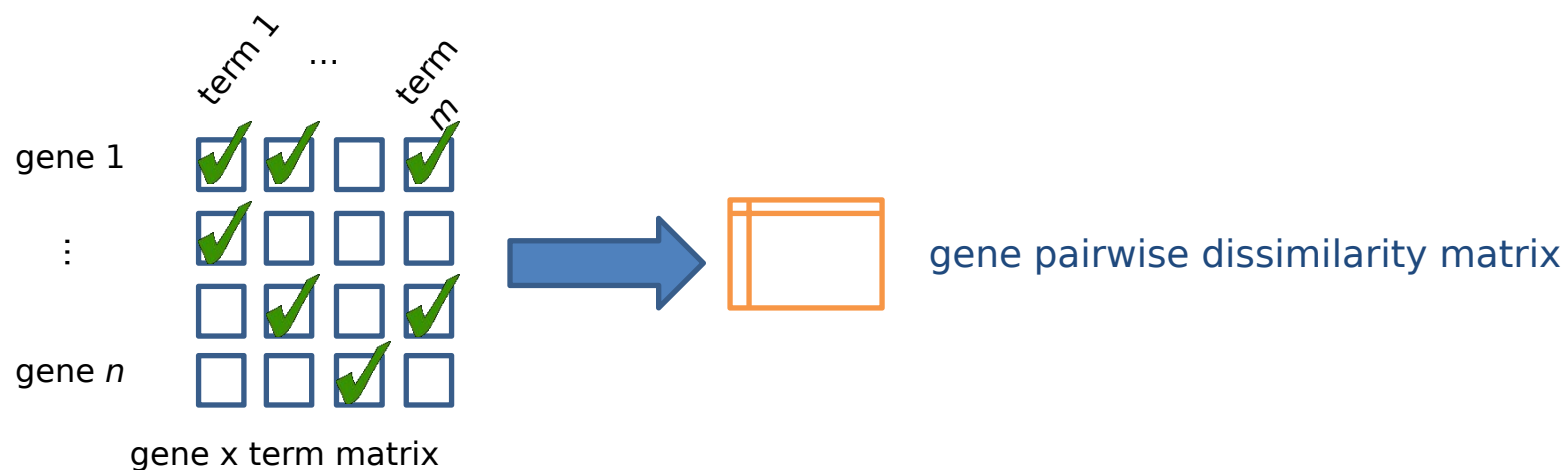
## Phylogenetic data: genome selection

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- Reference genomes should not be too evolutionary close to the genome of interest
- Reference genomes should not be redundant in order not to introduce biases
- Need to estimate the relevance of a genome with respect to
  - A genome of interest: is it not too closely related? is it informative?
  - A set of already selected reference genomes: redundancy *vs.* additional signal
- Parameters
  - Rearrangements
    - Significance of genes proximity on the chromosome
  - Core genome size
    - Maximize the coverage of the genome of interest

## Approaches:

- gene-term matrix: distance between rows
  - manhattan/euclidean, Jaccard, ...
- **but:** same weight for each GO-term
- based on GO-term similarity
  - adapt weight to information content



## Similarité entre Termes GO

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- Node/term information content

$$IC(term) = -\log p(term) \quad \text{with } p(\text{term}) = \text{freq}(\text{term})$$

- $MICA(t_1, t_2)$ : Maximum Information Common Ancestor

$$MICA(t_1, t_2) = \arg \max IC(t_i), t_i \in ancestors(t_1, t_2)$$

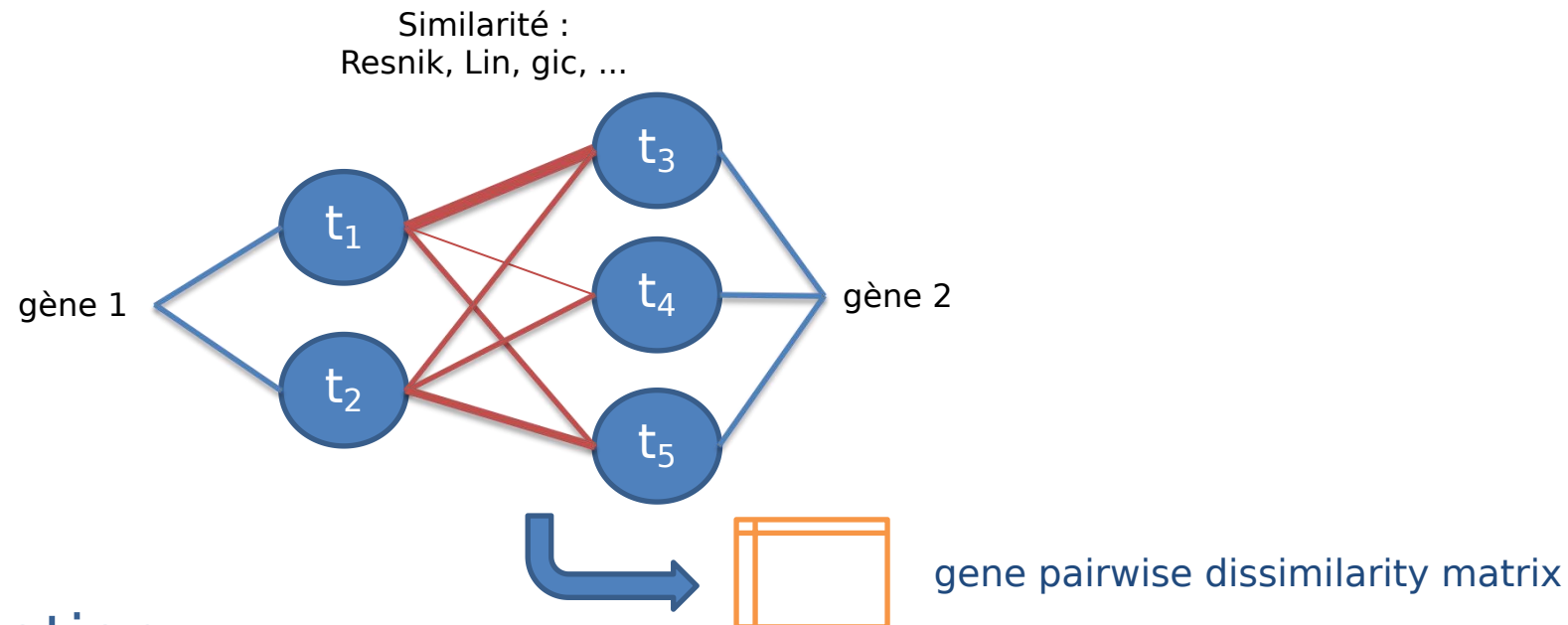
- $sim_{res}(t_1, t_2) = IC(MICA(t_1, t_2))$  [Resnik, 1995]

- $sim_{lin}(t_1, t_2) = IC(MICA(t_1, t_2))/(IC(t_1) + IC(t_2))$  [Lin, 1998]

- $sim_{gic}(t_1, t_2) = \frac{\sum_{t \in \{GO(t_1) \cap GO(t_2)\}} IC(t)}{\sum_{t \in \{GO(t_1) \cup GO(t_2)\}} IC(t)}$  [Pesquita et al., 2008]

## Possibilités :

- Similarité moyenne des termes communs au 2 gènes
  - Similarité maximale, ex :  $t_1-t_3$
  - Best Match Average (bma), ex :  $\text{ave}(t_1-t_3, t_2-t_5)$



## Application :

- Performances légèrement meilleures obtenues que les autres avec la combinaison Resnik + similarité maximale
  - à confirmer sur d'autres jeux de données ou d'autres contextes

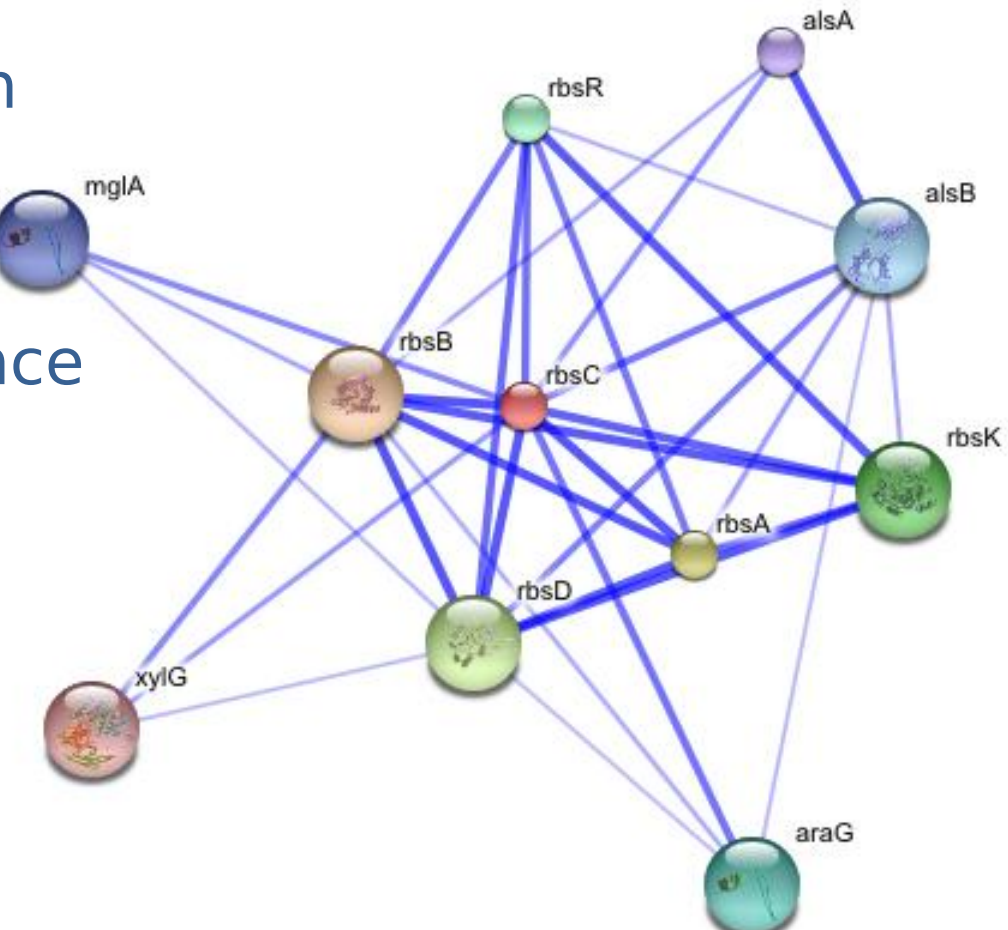
# Interactions

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- all pairs shortest path
- a pair of gene:  
shortest path length
- score: average distance

training: rbsA, rbsB, rbsC

candidate	score	rank	rank ratio
RBSK	1.000	2	0.0005136
RBSD	1.000	2	0.0005136
RBSR	1.000	2	0.0005136
ALSB	1.333	5	0.001284
ALSC	1.333	5	0.001284
YPHD	1.333	5	0.001284
MGLC	1.667	10.5	0.002696
XYLG	1.667	10.5	0.002696
ALSA	1.667	10.5	0.002696
YTFT	1.667	10.5	0.002696



from STRING  
<http://string-db.org>