

A dataset of small molecules triggering transcriptional and translational cellular responses

Mathilde Koch, Amir Pandi, Baudoin Delepine, Jean-Loup Faulon

▶ To cite this version:

Mathilde Koch, Amir Pandi, Baudoin Delepine, Jean-Loup Faulon. A dataset of small molecules triggering transcriptional and translational cellular responses. Data in Brief, 2018, 17, pp.1374-1378. 10.1016/j.dib.2018.02.061. hal-02622874

HAL Id: hal-02622874 https://hal.inrae.fr/hal-02622874v1

Submitted on 26 May 2020

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.





Contents lists available at ScienceDirect

Data in Brief





Data Article

A dataset of small molecules triggering transcriptional and translational cellular responses



Mathilde Koch ^a, Amir Pandi ^a, Baudoin Delépine ^{a,b,c}, Jean-Loup Faulon ^{a,b,c,d,*}

- ^a Micalis Institute, INRA, AgroParisTech, Université Paris-Saclay, 78350 Jouy-en-Josas, France
- ^b UMR 8030 Genomics Metabolics, Systems and Synthetic Biology Lab, CEA, CNRS, University of Evry-val-d'Essonne, University Paris-Saclay, Évry, France
- ^c CEA, DRF, IG, Genoscope, Évry 91000, France
- ^d SYNBIOCHEM Centre, Manchester Institute of Biotechnology, University of Manchester, 131 Princess Street, Manchester M1 7DN, UK

ARTICLE INFO

Article history: Received 30 January 2018 Received in revised form 13 February 2018 Accepted 21 February 2018 Available online 27 February 2018

ABSTRACT

The aim of this dataset is to identify and collect compounds that are known for being detectable by a living cell, through the action of a genetically encoded biosensor and is centred on bacterial transcription factors. Such a dataset should open the possibility to consider a wide range of applications in synthetic biology. The reader will find in this dataset the name of the compounds, their InChI (molecular structure), the publication where the detection was reported, the organism in which this was detected or engineered, the type of detection and experiment that was performed as well as the name of the biosensor. A comment field is also provided that explains why the compound was included in the dataset, based on quotes from the reference publication or the database it was extracted from. Manual curation of ACS Synthetic Biology abstracts (Volumes 1 to 6 and Volume 7 issue 1) was performed as well as extraction from the following databases: Bionemo v6.0 (Carbajosa et al., 2009) [1], RegTransbase r20120406 (Cipriano et al., 2013) [2], RegulonDB v9.0 (Gama-Castro et al.,

^{*} Corresponding author at: Micalis Institute, INRA, AgroParisTech, Université Paris-Saclay, 78350 Jouy-en-Josas, France. E-mail address: jean-loup.faulon@inra.fr (J.-L. Faulon).

2016) [3], RegPrecise v4.0 (Novichkov et al., 2013) [4] and Sigmol v20180122 (Rajput et al., 2016) [5].

© 2018 The Authors. Published by Elsevier Inc. This is an open access article under the CC BY license

(http://creativecommons.org/licenses/by/4.0/).

Specifications Table

Subject area Biology

Synthetic biology More specific sub-

iect area

Type of data Table

How data was

Database extraction from Bionemo v6.0, RegTransbase r20120406, RegulonDB acquired v9.0, RegPrecise v4.0 and Sigmol v20180122 as well as manual curation ACS

Synthetic Biology abstracts (Volumes 1 to 6 and Volume 7 issue 1)

Data format Experimental Analysed Not applicable

factors

Experimental Not applicable

features Data source

location

https://github.com/brsynth/detectable_metabolites

Data accessibility Data is with this article and on GitHub at https://github.com/brsynth/detect

able_metabolites

Value of the data

- This dataset provides a basis for the development of new biosensing circuits for synthetic biology and metabolic engineering applications, e.g. the design of whole-cell biosensor, high-throughput screening experiments, dynamic regulation of metabolic pathways, transcription factor engineering or creation of sensing-enabling pathways.
- This dataset provides a unique source of a broad number of compounds that can be detected and acted upon by a cell, increasing the possibility of orthogonal circuit design from the few usual compounds used in those applications.
- The manually curated section provides information on where the biosensor has been first reported and successfully used, enabling the reader to select trustworthy information for his application of choice.
- Detectable compounds can be searched by both by name and chemical similarity.
- This dataset is an update of [10.6084/m9.figshare.3144715.v1].

1. Data

The aim of this dataset is to identify and collect compounds that are known for being detectable by a living cell, through the action of a genetically encoded biosensor and is centred on bacterial transcription factors. The dataset should allow the synthetic biology community to consider a wide range of applications. The reader will find in this dataset the name of the compounds, their InChI (molecular structure), the publication where the detection was reported, the organism in which this was detected or engineered, the type of detection and experiment that was performed as well as the name of the biosensor. A comment field is also provided that explains why the compound was included in the dataset, based on quotes from the reference publication or the database it was extracted from. Manual curation of ACS Synthetic Biology abstracts (Volumes 1 to 6 and Volume 7 issue 1) was

performed as well as extraction from the following databases: Bionemo v6.0 [1], RegTransbase r20120406 [2], RegulonDB v9.0 [3], RegPrecise v4.0 [4] and Sigmol v20180122 [5].

This dataset is available online on GitHub to allow for further updates as well as community contributions

2. Experimental design, materials and methods

- Manual curation of ACS Synthetic Biology (Volume 1–6 and Volume 7 issue 1):
 All abstracts of ACS Synthetic Biology (Volume 1–6 and Volume 7 issue 1) were read and information relevant to this dataset was extracted from those abstracts. The aim of this manual curation was to establish a list of detectable compounds whose detection method was already successfully implemented in a synthetic circuit, providing a good basis for further implementation for synthetic biologists.
- Bionemo v6.0 [1]:

The SQL request used to create this dataset is:

SELECT DISTINCT substrate.id_substrate, minesota_code, name FROM substrate

INNER JOIN complex_substrate ON complex_substrate.id_substrate = substrate.id_substrate

INNER JOIN complex ON complex.id_complex=complex_substrate.id_complex WHERE activity='REG';

• RegTransbase r20120406 [2]:

The SQL request used to create this dataset is:

SELECT DISTINCT a.pmid, e.name, r.name

FROM regulator2effectors AS re

INNER JOIN exp2effectors AS ee ON ee.effector_guid=re.effector_guid

INNER JOIN dict_effectors AS e ON e.effector_guid=ee.effector_guid

INNER JOIN regulators AS r ON r.regulator_guid = re.regulator_guid

INNER JOIN articles AS a ON a.art_guid = ee.art_guid

ORDER BY e.name;

RegTransbase was not maintained anymore at the time of writing of this manuscript.

• *RegulonDB v9.0* [3]:

The SQL request used to create this dataset is:

SELECT c.conformation_id, c.final_state, e.effector_id, e.effector_name, tf.transcription_factor_id, tf. transcription_factor_name, p.reference_id, xdb.external_db_name

FROM effector AS e

INNER JOIN conformation_effector_link AS mm_ce ON mm_ce.effector_id = e.effector_id

LEFT JOIN conformation AS c ON c.conformation_id = mm_ce.conformation_id

LEFT [OIN transcription_factor AS tf ON tf.transcription_factor_id = c.transcription_factor_id

Table 1 Contribution of each data source.

Source	Compounds without structure	Compounds with structure	Unique compounds with structure
RegPrecise	136	418	73
BioNemo	5	499	8
RegTransBase	683	2057	63
RegulonDB	12	245	23
Sigmol	2	175	135
ACS Synthetic Biology	44	287	73
All sources	882	3681	729

The first column contains the data source, the second column the number of compounds found without a structure in that source, the third column the number of compounds with a structure (InChI) and the last column the number of compounds with a structure found only in that source.

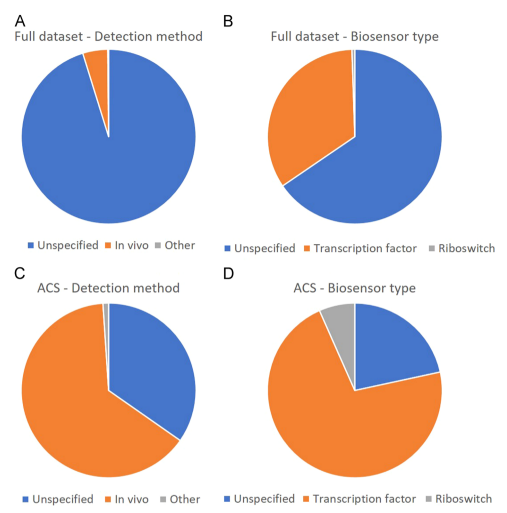


Fig. 1. Type of experiment and biosensor type in the full dataset and the manually curated dataset. A: Full dataset – detection method. B: Full dataset – biosensor type. C: ACS dataset – detection method. D: ACS dataset – biosensor type. A and C: other in detection method corresponds to *in silico*, *in vivo* and *cell-free* detections. C and D: ACS dataset is the dataset obtained from manual curation of ACS Synthetic Biology with compounds that have available structures.

LEFT JOIN object_ev_method_pub_link AS x ON x.object_id=c.conformation_id OR x.object_id=tf. transcription_factor_id OR x.object_id=e.effector_id

LEFT JOIN publication AS p ON p.publication_id = x.publication_id

LEFT JOIN external_db AS xdb ON xdb.external_db_id = p.external_db_id

WHERE c.interaction_type IS Null OR c.interaction_type! = 'Covalent';

• *RegPrecise v4.0* [4]:

The RegPrecise website was accessed (version v4.0) and all relevant data was extracted from the effector pages of the website.

• Sigmol v20170216 [5]:

Sigmol was accessed on 16/02/2017 and all effector data was retrieved from the unique *Quorum Sensing Signaling Molecule* page. In the "detected by" column, we provide the class of signaling compounds the compound belongs to. The comment field reads 'Extracted from Sigmol v20170216 – Uniq_QSSM_"number".

2.1. Data overview

In Table 1 are presented some characteristics of each data source: number of compounds without a structure from this source, total number of compounds with a structure from this source and number of compounds with a structure found only in this source. The last column in particular shows that around half the compounds are found in more than one data source.

Fig. 1 shows the repartition of the type of experiment (*in vivo*, unspecified or other), as well as the repartition of Biosensor type (Transcription factor, riboswitch or unspecified) in the full dataset and the manually curated dataset from ACS Synthetic Biology.

Acknowledgements

This work was supported by the French National Research Agency (ANR-15-CE21-0008), the Biotechnology and Biological Sciences Research Council, Centre for Synthetic Biology of Fine and Speciality Chemicals (BB/M017702/1); Synthetic Biology Applications for Protective Materials (EP/N025504/1. M.K is supported by DGA (French Ministry of Defense) and Ecole Polytechnique. BD was supported by Structure et Dynamique des Systemes Vivants Doctoral School, Universite Paris Saclay.

Competing financial interests

None declared.

Appendix A. Supplementary material

Supplementary data associated with this article can be found in the online version at doi:10.1016/j. dib.2018.02.061.

Transparency document. Supporting information

Transparency data associated with this article can be found in the online version at https://doi.org/10.1016/j.dib.2018.02.061.

References

- [1] G. Carbajosa, A. Trigo, A. Valencia, I. Cases, Bionemo: molecular information on biodegradation metabolism, Nucleic Acids Res. 37 (2009) D598–D602. http://dx.doi.org/10.1093/nar/gkn864.
- [2] M.J. Cipriano, P.N. Novichkov, A.E. Kazakov, D.A. Rodionov, A.P. Arkin, M.S. Gelfand, I. Dubchak, RegTransBase a database of regulatory sequences and interactions based on literature: a resource for investigating transcriptional regulation in prokaryotes, BMC Genom. 14 (2013) 213. http://dx.doi.org/10.1186/1471-2164-14-213.
- [3] S. Gama-Castro, H. Salgado, A. Santos-Zavaleta, D. Ledezma-Tejeida, L. Muñiz-Rascado, J.S. García-Sotelo, K. Alquicira-Hernández, I. Martínez-Flores, L. Pannier, J.A. Castro-Mondragón, A. Medina-Rivera, H. Solano-Lira, C. Bonavides-Martínez, E. Pérez-Rueda, S. Alquicira-Hernández, L. Porrón-Sotelo, A. López-Fuentes, A. Hernández-Koutoucheva, V. Del Moral-Chávez, F. Rinaldi, J. Collado-Vides, RegulonDB version 9.0: high-level integration of gene regulation, coexpression, motif clustering and beyond, Nucleic Acids Res. 44 (2016) D133–D143. http://dx.doi.org/10.1093/nar/gkv1156.
- [4] P.S. Novichkov, A.E. Kazakov, D.A. Ravcheev, S.A. Leyn, G.Y. Kovaleva, R.A. Sutormin, M.D. Kazanov, W. Riehl, A.P. Arkin, I. Dubchak, D.A. Rodionov, RegPrecise 3.0 a resource for genome-scale exploration of transcriptional regulation in bacteria, BMC Genom. 14 (2013) 745. http://dx.doi.org/10.1186/1471-2164-14-745.
- [5] A. Rajput, K. Kaur, M. Kumar, SigMol: repertoire of quorum sensing signaling molecules in prokaryotes, Nucleic Acids Res. 44 (2016) D634–D639. http://dx.doi.org/10.1093/nar/gkv1076.