See discussions, stats, and author profiles for this publication at: https://www.researchgate.net/publication/339313738

#### Synthetic control devices for gene regulation in filamentous fungus Penicillium chrysogenum (STF/CRISPRa)

Poster · February 2020

ATIONS		reads 16	
*	and the second sec		
2	Laszlo Mozsik		Roel A L Bovenberg
	University of Groningen	22	Royal DSM
	3 PUBLICATIONS 9 CITATIONS		104 PUBLICATIONS 2,834 CITATIONS
	SEE PROFILE		SEE PROFILE
	Arnold J.M. Driessen		
2	University of Groningen		
	546 PUBLICATIONS 24,582 CITATIONS		
	SEE PROFILE		

Some of the authors of this publication are also working on these related projects:

 Project
 (Multi)drug resistance View project

 Project
 OMNIYEAST View project



university of groningen

faculty of mathematics and natural sciences molecular microbiology and synthetic biology



# Synthetic control devices for gene regulation



## in Penicillium chrysogenum

### László Mózsik<sup>1\*</sup>, Zsofia Buttel<sup>1</sup>, Roel A.L. Bovenberg<sup>2,3</sup>, Arnold J. M. Driessen<sup>1</sup>, Yvonne Nygård<sup>4</sup>

<sup>1</sup> Department of Molecular Microbiology, GBB, University of Groningen, Groningen, NL; <sup>2</sup> DSM Biotechnology Center, DSM, Delft, NL; <sup>3</sup> Synthetic Biology and Cell Engineering, GBB, University of Groningen, Groningen, NL; <sup>4</sup> Department of Biology and Biological Engineering, Division of Industrial Biotechnology, Chalmers University of Technology, Göteborg, SE **\*E-mail: L.Mozsik@rug.nl** 

#### **Introduction**

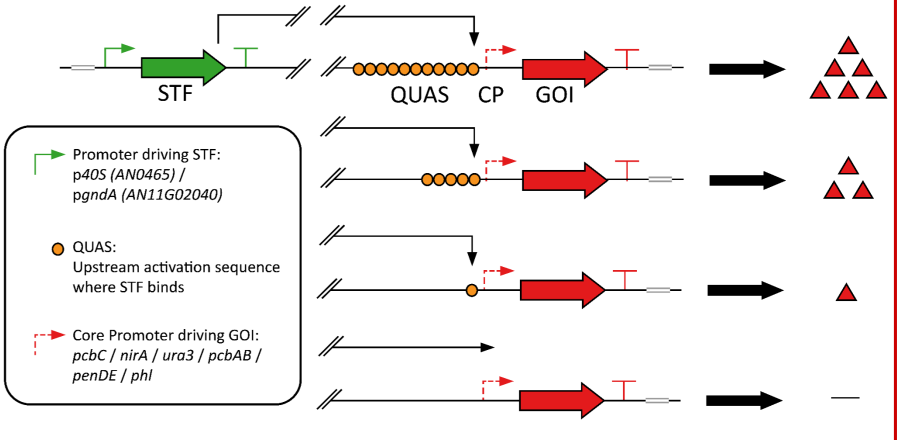
Synthetic biology aims at controlled gene regulation that can lead to increased production of chemicals and pharmaceuticals. In this work synthetic control devices were developed for *Penicillium chrysogenum*, a model filamentous fungus and industrially relevant cell factory.

In the synthetic transcription factor (STF) the QF DNA-binding domain of the transcription factor of the quinic acid gene cluster of *Neurospora crassa*<sup>2</sup> is fused to the VP16 activation domain. This synthetic transcription factor controls the expression of genes under a synthetic promoter containing quinic acid upstream activating sequence (QUAS) elements, where it binds prior to a core promoter (CP) (**Fig.1.**).

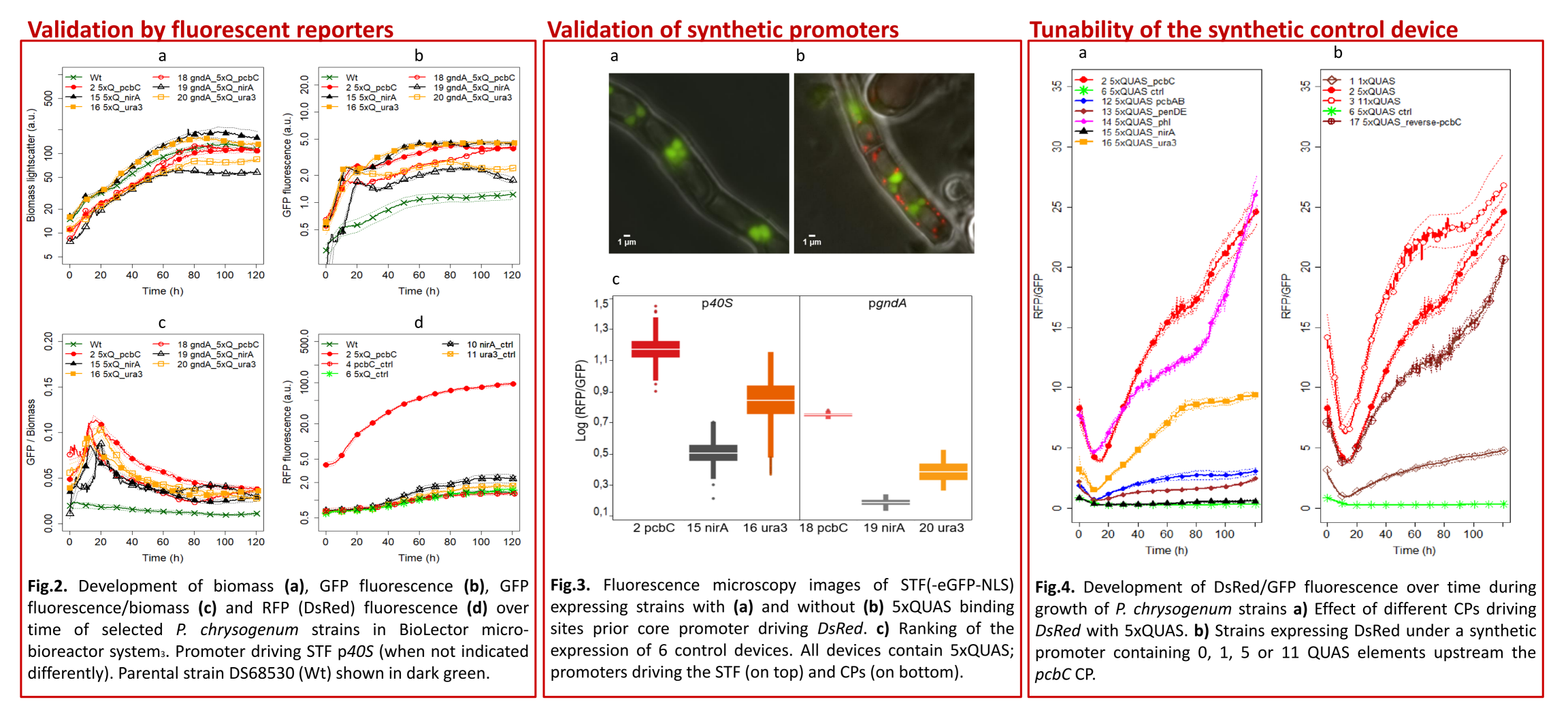
#### Results

The strength of the control device can be altered by altering the expression of the transcription factor, the core promoter upstream the QUAS or the number of QUAS elements (**Fig.2.-4.**).

The versatility of the control device was demonstrated by fluorescent reporters (eGFP-NLS, DsRed-SKL) and its application was confirmed by synthetically controlling the production of Penicillin V (Fig.5.).



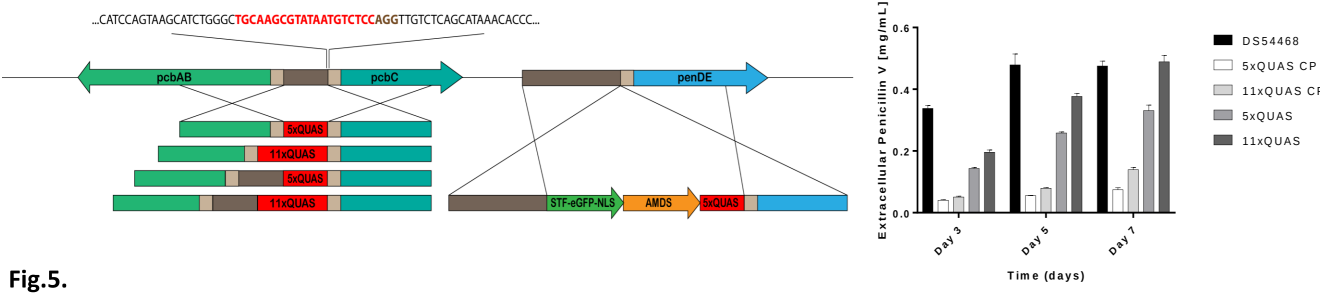
#### **Fig.1.** Schematic representation of the control devices.

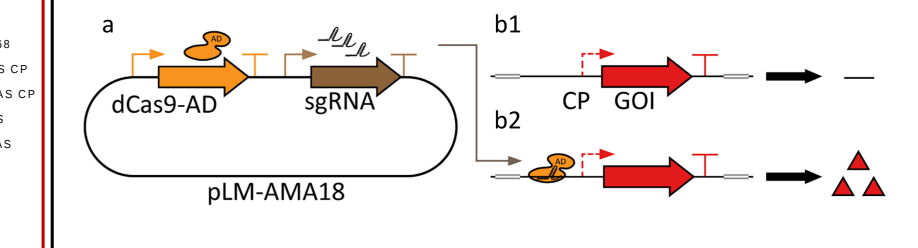


#### Validation by Penicillin V production

#### **Envisioned silent promoter activation with CRISPRa**

а





#### Fig.6.

a) Schematic representation of CRISPR/Cas9 based integration₄ and co-transformation of the synthetic control device and QUAS elements into the penicillin cluster of DS54468. Designed sgRNA targets boundary between the promoter of *pcbC*.
 b) Extracellular Penicillin V production of strains where the penicillin biosynthesis cluster is under control of the control device and parental strain DS54468 in shake flask cultivation in penicillin producing medium. Data shows 3 independent cultures, measured in replicates.
 a) Schematic representation of the synthetic control device and parental strain DS54468 in shake flask cultivation in penicillin biosynthesis of interest biosynthesis of interest biosynthesis of interest biosynthesis of strains where the penicillin biosynthesis of interest biosynthesis biosynthesis of interest biosynthesis of interest biosynthesis of interest biosynthesis biosynthesis of interest biosynthesis of interest biosynthesis biosynthesynthesis biosynthesis biosy

a) Schematic representation of plasmid delivery of the CRISPRa components (dCas9m4-AD and sgRNA) where AD stands for transcriptional activator domain.
b1) Representation of synthetic silent promoter: no transcription from the gene of interest (GOI) without the integrated QUAS sequences (Figure 1).
b2) sgRNA guided dCas9-AD binding upstream the synthetic silent promoter, promoting transcriptional activation.

#### **Conclusions/Outlook**

- Modular, synthetic control devices were developed for P. chrysogenum and their function was demonstrated with fluorescent reporters and Penicillin V production.
- The strength of the control devices can be altered by altering the expression of the STF, the core promoter upstream the QUAS or the amount of QUAS elements, leading
  to expression ranging from barely detectable to similar the highest expressed native genes.
- We anticipate that these well-characterized and robustly performing control devices are highly useful tools in the development of filamentous fungi as production hosts.
- Silent promoter activation was envisioned with the CRISPRa system. This genome editing free transcriptional regulatory tool could be further expand the fungal toolbox of synthetic regulatory systems and could be used to interrogate transcriptionally silent fungal gene clusters.

#### References:

1) Mózsik, L., Büttel, Z., Bovenberg, R.A.L., Driessen, A. J. M., & Nygård, Y. I.; Synthetic control devices for gene regulation in Penicillium chrysogenum. Microb Cell Fact 18, 203. (2019)

2) Potter, C. J., Tasic, B., Russler, E. V., Liang, L., and Luo, L. ; The Q System: A Repressible Binary System for Transgene Expression, Lineage Tracing and Mosaic Analysis, Cell 141, 536-548. (2010)

3) Polli, F., Meijrink, B., Bovenberg, R. A. L., and Driessen, A. J. M.; New promoters for strain engineering of Penicillium chrysogenum, Fungal Genet Biol 89. (2016)

4) Pohl, C., Mózsik, L., Driessen, A. J. M., Bovenberg, R. A. L., & Nygård, Y. I.; Genome Editing in Penicillium chrysogenum Using Cas9 Ribonucleoprotein Particles. Synthetic biology: Methods and Protocols (pp. 213-232). (2018)

#### This research was supported by the QuantFung Project (Seventh Framework No. 607332) and ALERT Project (Marie Skłodowska-Curie No. 713482).