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# Combining systems and synthetic biology for in vivo enzymology

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# Abstract

Enzymatic parameters are classically determined in vitro, under reaction conditions that are far from those encountered in cells, casting doubt on their physiological relevance. In this study, we developed a generic approach combining tools from synthetic and systems biology to measure enzymatic parameters in vivo. In the context of a synthetic pathway aiming to produce phytoene and lycopene in Saccharomyces cerevisiae, we focused on four membrane-associated enzymes, a phytoene synthase and three phytoene desaturases, which are difficult to study in vitro. We designed, built, and analyzed a collection of yeast strains mimicking substantial variations in substrate concentration (> 200-fold change) by strategically manipulating the copy number and expression strength of the gene encoding geranyl-geranyl pyrophosphate (GGPP) synthase. We successfully determined in vivo-equivalent Michaelis-Menten parameters ( $K_M$ ,  $V_{max}$  and  $k_{cat}$ ) for GGPP-converting phytoene synthase from quantitative metabolomics, fluxomics and proteomics data, highlighting differences between in vivo and in vitro parameters. Then, leveraging the versatility of the same set of strains to cover a wide range of phytoene concentrations, we successfully extracted enzymatic parameters for two of the three phytoene desaturases. Our approach demonstrates the feasibility of assessing enzymatic parameters along a pathway of interest directly in vivo, providing a novel perspective on the kinetic characteristics of enzymes in real cellular conditions.

# Introduction

Enzymes catalyze most of the chemical reactions in living systems. A comprehensive interpretation of enzymatic parameters is therefore of paramount importance to grasp the complexity of cellular metabolism. The vast amount of data collected for thousands of enzymes has contributed to significant progress in our understanding of the remarkable chemical capabilities of biocatalysts and of their roles in cellular reactions.

Enzymatic reactions are traditionally analyzed *in vitro*, under dilute conditions, using pure or semi-pure protein samples in buffer solution. In contrast, the cellular medium is most accurately viewed as a heterogeneous, dense, crowded gel, containing various types of macromolecules and cellular lipidic organelles, with potential partitioning effects and variations in substrate and/or product diffusion coefficients (1). The parameters determined in classical enzymology experiments may therefore not be representative of *in vivo* reaction rates and equilibrium constants (2, 3). While some progress has been made in implementing and understanding viscosity and crowding effects in *in vitro* enzymatic assays, these conditions do not mimic the intrinsic complexity of the cellular environment (2).

In vivo enzymology seems to be the obvious approach to measure enzymatic parameters inside cells. Early attempts were made using the enzyme photolyase, for which both in vitro and in vivo parameters were determined (4, 5). Meanwhile, the *in vivo* V<sub>max</sub> values of ten central carbon metabolism enzymes determined in the early 90s revealed significant differences between in vivo and in vitro assays for heteromeric protein complexes (6). The past ten years have seen growing interest from systems biology in determining the *in vivo*  $k_{cat}$  of native enzymes in model organisms. The *in vivo* apparent  $k_{cat}$ of E. coli enzymes have been determined independently by two research groups, leveraging advances in absolute protein quantification and high throughput metabolomics (7, 8). These values were obtained by dividing the flux of enzymatic reactions by the absolute abundance of the corresponding enzymes. Both studies found correlations between in vivo and in vitro data (with correlation coefficients around 0.6), suggesting that this method could serve as an alternative to *in vitro* assays. While this systems biology approach has provided valuable information, it cannot be employed to determine apparent  $K_{MS}$  in vivo. Zotter et al. measured the activity and affinity of TEM1- $\beta$  lactamase in mammalian cells in vivo with confocal microscopy, using a fluorescently tagged enzyme and a fluorescent substrate and product (9). They observed that the catalytic efficiency  $(k_{cat}/K_{M})$  of this enzyme differs in vitro and in vivo due to substrate attenuation, indicating that in vitro data are not always indicative of in vivo function. Zotter et al. is probably the most comprehensive in vivo enzymology study to date, but this approach cannot be generalized because of a lack of universally appropriate fluorescent substrates/products for all enzymes. In a recent investigation of the *in vivo* kinetic parameters of thymidylate kinase (TmK), an interesting finding was the difference in TmK's activity pattern when the substrate (thymidine monophosphate, TMP) was supplied in the media versus provided by internal metabolism, with Michaelis-Menten kinetics in the former and Hill-like kinetics in the latter. The authors hypothesized that the limited diffusion of TMP might be due to its confinement in a putative metabolon in E. coli (10).

*In vivo* enzymology is a particularly attractive prospect for membrane and multimeric proteins (6), which are tedious to purify and for which activity assays are difficult to optimize, mostly because artificial membranes are required (11–14). In the industrially important carotenoid pathway for example, despite the expression of numerous carotene biosynthesis enzymes, our understanding of their enzymatic behavior remains limited because many are membrane-associated proteins. While

kinetic parameters for some phytoene synthases, sourced from plants or bacteria, have been established *in vitro*, many enzymatic assays require the co-expression of geranylgeranyl pyrophosphate (GGPP) synthase to attain activity, possibly because of the amphiphilic nature of the substrate or the requirement of membranes (12, 14). Another example of the difficulty of *in vitro* assays is phytoene desaturase, for which discernible enzyme activity has only been achieved in engineered environments (e.g. liposomes) with cell-purified substrate (13, 15–17).

This study combines synthetic and systems biology tools to develop an original *in vivo* enzymology approach. Synthetic biology offers a remarkable set of genetic engineering tools to precisely modulate the activity of enzymes in synthetic pathways, enabling *in vivo* control of substrate concentrations for the studied enzymatic reactions. In turn, systems biology can provide quantitative data on these reaction (fluxes and enzyme, substrate and product concentrations) and computational tools to model their behavior. We applied this approach to investigate a synthetic carotenoid production pathway in *Saccharomyces cerevisiae*, with industrial applications ranging from foods to pharmaceuticals.

# Results

#### General principles of the proposed in vivo enzymology method

Enzymes are usually characterized in terms of their affinity ( $K_{\rm M}$ ) and activity (maximal reaction rate  $V_{max}$  and turnover number  $k_{cat}$ ) (Figure 1). These parameters are typically determined *in vitro* by varying the substrate concentration across a relatively broad range (about two orders of magnitude) and measuring the reaction rate for each substrate concentration (18). Different mathematical formulas, such as the Michaelis-Menten equation, have been derived to estimate enzymatic parameters from these data. We propose a novel approach wherein the substrate concentration is varied directly within cells (Figure 1), by modulating the concentration of the enzyme producing the substrate of the studied reaction. Levels of the substrate-producing enzyme are varied using different combinations of promoter strengths and gene copy numbers, and translate into a wide range of substrate concentrations (2-3 orders of magnitude, as detailed in the following sections). In our experimental setup, each substrate concentration is achieved using a specifically engineered yeast strain. In each of these strains, the gene encoding the enzyme of interest is expressed at a given level, mirroring the conditions of in vitro experiments. Engineered strains are then grown under steady-state conditions (i.e., exponential growth), where reaction fluxes and metabolite concentrations remain constant (19). This stability allows for the accurate measurement of product formation fluxes (equivalent to reaction rates), intracellular substrate concentrations, and enzyme concentrations. The data obtained can then be combined to calculate apparent in vivo kinetics parameters, denoted  $K_{1/2}^{cell}$ ,  $V_{max}^{cell}$  and  $k_{cat}^{cell}$  in analogy with the classical  $K_{M}$ ,  $V_{max}$  and  $k_{cat}$  parameters (Figure 1).



**Figure 1.** General principle of the proposed strategy for *in vivo* enzymology and comparison between *in vitro* and *in vivo* enzymology approaches.

#### Construction of yeast strains to investigate a synthetic carotenoid production pathway

We evaluated our strategy by investigating a synthetic carotenoid pathway in yeast (S. cerevisiae), an industrially important chassis in biotechnology, which includes two membrane-interacting enzymes that are challenging to investigate in vitro (12, 15, 16, 20): phytoene synthase (CrtB) and phytoene desaturase (Crtl) (Figure 2A). Phytoene synthase, the first enzyme in the carotenoid pathway and considered the bottleneck of carotenoid biosynthesis in plants (20), condenses two molecules of geranylgeranyl pyrophosphate (GGPP) head-to-head to form phytoene. Phytoene synthases have proved difficult to express in soluble and active forms in microorganisms (12, 14, 20). In vitro rates and affinity constants have been determined for both plant and bacterial phytoene synthases but these parameters may have been altered by the conditions of the *in vitro* assays—adding detergents for the bacterial enzyme and co-expressing a GGPP synthase or using semi-crude extracts for the plant enzyme (12, 14, 20–23). Following this enzymatic step, three insaturations are introduced in phytoene to produce lycopene. In non-photosynthetic bacteria and fungi, three reaction steps are catalyzed by a single enzyme (Crtl in bacteria and CarB in fungi) (24), whereas in plants and cyanobacteria, four different enzymes are involved in the conversion of phytoene to lycopene (25). Given that these desaturation steps are a major bottleneck in microbial carotenoid biosynthesis, determining in vivo enzymatic constants is an industrially relevant challenge (10, 21, 22).

To analyze these enzymes, we assembled a set of strains covering a broad range of substrate concentrations (**Supplementary Table S1**). To create the collection of strains with different intracellular levels of GGPP, the substrate of phytoene synthase (**Figure 2A**), we first increased GGPP production by modifying the native terpene pathway: strong constitutional expression of *ERG20* and of the truncated form of *HMG1* and deletion of the main phosphatase gene involved in GGPP and FPP pyrophosphate group removal (*DPP1*), yielding the strain yENZ015 (**Figure 2A** and **Supplementary Figure S1**) (26, 27). Three different integration cassettes containing different microbial GGPP synthases (CrtE) expressed under different constitutive promoters were integrated into the yENZ015 genome, yielding a series of strains designed to produce a wide range of GGPP concentrations (**Supplementary Figure S1**).

Although the *DPP1* gene was deleted in our strains to prevent GGPP degradation, the presence of a second phosphatase (LPP1) (28), meant that a fraction of GGPP was converted into geranyl geraniol (GGOH). The two other pathways (ubiquinone biosynthesis and protein prenylation) consume only a small amount of GGPP. We therefore used GGOH as a proxy to verify that the GGPP concentration was indeed modulated *in vivo*, as done previously (23, 29–31). Indeed, while intracellular GGPP measurements require extensive cell quenching and extraction procedures, and liquid chromatography-mass spectrometry methods for accurate quantification of isoprenoid intermediates, extracellular GGOH measurements can rapidly be performed using conventional gas chromatography after simply adding a dodecane layer to the culture. We verified the correlation between GGOH and GGPP concentrations for a subset of the strains used in this study ( $R^2 = 0.97$ , **Figure 2B**). GGOH production varied by a factor of 15 in these strains (**Figure 2C**), and the strains all had similar growth rates ( $0.40\pm0.03$  h<sup>-1</sup>) (**Figure 2C**) indicating that there was no change in cell physiology. As detailed below, we used this set of strains as a base to study a phytoene synthase (https://www.uniprot.org/uniprotkb/P21683/entry, EC 2.5.1.32) and three different lycopene-forming phytoene desaturases (EC 1.3.99.31).



**Figure 2.** Construction of the set of yeast strains with different intracellular GGPP concentrations. **A.** Genomic modification scheme for the yENZ015 strain and intracellular fates of GGPP in *S. cerevisiae*. Overexpressed enzymes are shown in blue, enzyme deletion is shown in red, and enzyme modulation is shown in green. **B.** Correlation between extracellular GGOH and intracellular GGPP concentrations in a subset of the strains. **C.** GGOH concentration (green bars) and specific growth rate of *S. cerevisiae* strains (yellow dots).

#### In vivo enzymatic parameters of phytoene synthase from Pantoea ananas

In a classical enzymatic reaction setup, enzyme concentrations must be carefully controlled to establish an informative set of reaction rate vs substrate concentration data points. Here, we evaluated three enzyme concentrations by expressing *Pantoea ananas crtB* (PaCrtB) under the control of three different constitutive promoters (TEF1pmut2, PGI1p and PDC1p, **Figure 3A**). As described above, we used the GGOH level as a proxy for the GGPP concentration.

In all the strains, expressing *crtB* lowered GGOH concentrations (**Figure 3B**), which is likely due to the decrease in GGPP concentration resulting from its conversion into phytoene. Consistent with this interpretation, the GGOH concentration progressively decreased as the *crtB* promoter strength

increased (TEF1pmut2 < PGI1p < PDC1p), i.e. as PaCrtB expression increased (Figure 3B). Therefore, the GGOH concentration was very low at high PaCrtB level (PDC1 promoter), decreasing the chance of reaching enzyme saturation. In contrast, the phytoene concentration increased as PaCrtB expression increased (Figure 3C). The phytoene concentration was too low for accurate quantification of the phytoene flux at low levels of PaCrtB (TEF1mut2 promoter), but accurate measurements could be made at medium and high PaCrtB levels. Based on these results, we then measured the phytoene production rate and the intracellular GGPP concentration in strains with medium PaCrtB levels (PGI1 promoter) (Figure 3D). A 180-fold variation in GGPP concentration (from 5 to 910 nmol/g DCW) was observed in these strains. Partial saturation was achieved, leading to a nonlinear relationship between the phytoene flux and the GGPP concentration, as observed in classical in vitro experiments. We verified that PaCrtB levels were identical in all strains by western blot and using mass spectrometrybased proteomics (Supplementary Figure S2A and Supplementary Table S2), indicating that PaCrtB activity ( $V_{max}^{cell}$ ) was also similar in all strains. Precise estimates of  $V_{max}^{cell}$  (847±82  $\mu$ M/h, rsd = 10 %) and  $K_{1/2}^{\text{cell}}$  (191±30  $\mu$ M, rsd = 16 %) were obtained by fitting these data using an irreversible Michaelis-Menten model (consistent with previous reports (20) and with the high  $\Delta G^0$  of this reaction) (Supplementary Table S3). The affinity of PaCrtB for GGPP estimated from our in vivo data is significantly lower than measured in vitro (41  $\mu$ M) (20), highlighting the necessity of measuring enzyme parameters directly within the intracellular environment. Meanwhile, the  $k_{cat}^{cell}$  of 6±1 s<sup>-1</sup>, obtained from absolute quantitative proteomics measurements of the enzyme concentration (42 nM) in strains expressing PaCrtB under the control of PGI1p, is comparable to the one obtained in vitro for CrtB in Pantoea agglomerans (14 s<sup>-1</sup>, Supplementary Table S4) (14).

The theoretical maximum phytoene production flux in these strains is therefore  $847\pm86 \mu$ M/h. The bottleneck for carotenoid biosynthesis in plants is phytoene synthase, whose activity is regulated by a combination of transcriptional, post-transcriptional, and post-translational mechanisms to adjust carotenoid production (32). In our synthetic system however, our results suggest that GGPP biosynthesis is the limiting step since five copies of the GGPP synthase gene are required to achieve just 51 % of the maximum phytoene production flux, due to the lack of phytoene synthase saturation.



**Figure 3.** *In vivo* characterization of phytoene synthase from *Pantoea ananas*. **A.** Scheme for the biosynthesis of phytoene by CrtB. **B.** Extracellular GGOH concentration for the different GGPP producing strains expressing PaCrtB under different promoter strengths. **C.** Intracellular phytoene concentration in the different GGPP producing strains expressing PaCrtB under different promoter strengths. **D.** *In vivo* characterization of CrtB in *S. cerevisiae*: phytoene production as a function of GGPP concentration in strains with medium level of CrtB (PGI1p). Each data point represents an independent biological replicate, red line represents the best fit of a Michaelis-Menten rate law, and shaded area corresponds to 95 % confidence interval on the fit.

# Straightforward extension to analyze downstream enzymes of the carotenoid pathway: phytoene desaturases

The most time-consuming tasks in the proposed *in vivo* enzymology protocol are the molecular cloning and strain engineering steps required to generate strains with different substrate concentrations. However, the strains used to investigate a given enzyme can be recycled to investigate downstream steps in the same metabolic pathway, significantly streamlining experiments, which become a simple plug-and-measure process (**Figure 4A**). Integration of a single copy of Pa*crtB*, controlled by the highexpression promoter PDC1 into our initial set of strains produced strains with a 430-fold variation in phytoene concentration (from 4 to 1641  $\mu$ M). These strains thus form an ideal framework to investigate the *in vivo* kinetics of the next metabolic step with various phytoene desaturases.

To demonstrate this point, we chose three widely used microbial lycopene-forming phytoene desaturases (1.3.99.31) (21, 22, 24, 25): *P. ananas* Crtl (P21685, PaCrtl), which, along with *Rubrivivax gelatinosus* Crtl, is one of only two lycopene-forming phytoene desaturases with published *in vitro* enzymatic parameters (13, 33); and two fungal Crtls widely used in carotenoid production, one from *Phaffia rhodozyma*, formely *Xanthophyllomyces dendrorhous* (A0A0P0KMF9, XdCrtl), and the other from *Blakeslea trispora* (Q67GI0, BtCrtl), with no enzymatic data available to date.

As for the investigation of phytoene synthase, the expression of the phytoene desaturases was controlled by different promoters: TDH3p for PaCrtl, PGK1p for XdCrtl, and PGI1p for BtCrtl. These promoters were chosen based on the relative lycopene producing abilities of the different CrtIs in *S. cerevisiae*. This was especially important for BtCrtl, where high expression in the exponential phase caused cell death due to the formation of lycopene crystals (**Supplementary Figure S3A**). PaCrtl and XdCrtl levels did not vary with the phytoene concentration of the strains, but BtCrtl levels were slightly higher in the strains with the lowest phytoene concentrations (**Supplementary Figure S4A**). The higher levels of PaCrtl compared with those of the fungal phytoene desaturases is explained in large part by the use of a TDH3 promoter. However, lycopene fluxes were 10 times lower with PaCrtl than with XdCrtl or BtCrtl (**Figure 4**). Phytoene levels were high enough to saturate both fungal CrtIs (**Figure 4**) but not PaCrtl, for which lycopene fluxes remained low for all tested phytoene concentrations.

The experimental data from the three enzymes were fitted with an irreversible Michaelis-Menten equation, from which  $V_{max}^{cell}$  and  $K_{1/2}^{cell}$  values were successfully obtained for XdCrtl and BtCrtl but not for PaCrtl, for which only lower limits for  $V_{max}^{cell}$  (13  $\mu$ M·h<sup>-1</sup>) and  $K_{1/2}^{cell}$  (560  $\mu$ M) could be determined **(Supplementary Table S3).** The  $V_{max}^{cell}$  values obtained for the two fungal enzymes were similar, but slightly higher for BtCrtl (191±10  $\mu$ M·h<sup>-1</sup>) than for XdCrtl (168±7  $\mu$ M·h<sup>-1</sup>); however, the  $K_{1/2}^{cell}$  for BtCrtl (17±4  $\mu$ M) was 20 times smaller than XdCrtl's (380±49  $\mu$ M). The  $k_{cat}^{cell}$  obtained by integrating

quantitative proteomics measurements in strains expressing XdCrtI under PGK1p was 0.030±0.006 s<sup>-1</sup> (Supplementary Table S3).

These results highlight the versatility of our approach to compare the *in vivo* behavior of enzymes from different organisms, using the same set of strains, and provide valuable information to understand and optimize natural and synthetic pathways *in vivo*.



**Figure 4.** *In vivo* characterization of three phytoene desaturases. **A.** Scheme for the biosynthesis of lycopene by Crtl. **B–D**. *In vivo* characterization of Crtl from *P. ananas* (B), *X. dendrorus* (C) and *B. trispora* (D). Each data point represents an independent biological replicate, red lines represent the best fits of a Michaelis-Menten rate law, and shaded areas correspond to 95 % confidence intervals on the fits.

# Discussion

Inspired by conventional *in vitro* enzyme assays, we developed an innovative approach to measure enzyme kinetics *in vivo*. We successfully used the proposed method to estimate the *in vivo* equivalents of Michaelis-Menten parameters for a phytoene synthase and two phytoene desaturases in *S. cerevisiae*.

These results highlight the reliability and versatility of our *in vivo* enzymology approach, which hinges on solving two methodological challenges: 1) controlling the substrate pool over a broad concentration range, and 2) measuring the total concentrations of substrate, product and enzyme. The first challenge was addressed using synthetic biology tools to modulate the concentration of a substrate-producing enzyme (here GGPP synthase, CrtE). As mentioned above, our method of using multiple strains to mimic variations in substrate concentrations can be used to target multiple enzymes in the same metabolic pathway, in this case phytoene synthase and phytoene desaturase. The increasing availability of high-throughput platforms for strain engineering should accelerate the entire construction process. The second challenge was overcome thanks to sensitive, quantitative metabolomics and proteomics techniques. In the absence of absolute quantitative proteomics data, the proposed method still allows for the determination of  $V_{max}^{cell}$  and  $K_{1/2}^{cell}$  values, providing sufficient information for most enzymology and metabolic engineering studies.

We demonstrate that substrate concentrations can be varied *in vivo* over a wide range: here, the GGPP concentration was varied by a factor 180 and the phytoene concentration by a factor 430. This variation allowed clear enzyme saturation curves to be observed for the two fungal phytoene desaturases (XdCrtI and BtCrtI). However, while the range of substrate concentrations explored was

significantly wider than those used in *in vitro* studies of the same enzymes (Supplementary Table S4) (15, 20), complete saturation was not achieved for PaCrtB or PaCrtI. This may simply be due to a low in vivo affinity (high  $K_{\rm M}$ ) or a low substrate availability around the enzyme. In addition, other explanations merit attention as they also convey general hypotheses on metabolic systems: i) substrate or product toxicity detrimental to cell growth (e.g. the specific growth rate is nearly 50 % lower for lycopene concentrations above 400 µM, Supplementary Figure S3B), ii) overflow mechanisms that relocate some of the substrate to a different cell compartment or even outside the cell (as observed here for GGOH) (34, 35), iii) the presence of alternative pathways that divert additional substrate produced above a certain concentration threshold (36, 37), or iv) intrinsic properties of metabolic systems whereby production fluxes tend to decrease when product concentrations increase (38-40). While these mechanisms contribute to global metabolite homeostasis (41-43) and are essential for cell viability, they limit substrate accumulation and thus prevent complete saturation of the enzyme. Here, the mechanisms involved in the non-saturation of two P. ananas enzymes may be completely different. For PaCrtl, the observed phytoene (substrate) and lycopene (product) concentrations are either equivalent or lower than the values obtained with the two fungal Crtl enzymes. In this case, the absence of saturation points to a true difference in the bacterial and the fungal enzymes' affinity for phytoene. For PaCrtB, given that the GGPP concentration in the cell was extremely high (around 200  $\mu$ M), the lack of saturation is more likely indicative of a heterogeneous distribution of GGPP in the cell, with only a fraction of GGPP in the vicinity of the enzyme.

While classical in vitro enzyme assays involve homogeneous, highly diluted buffers, the intracellular environment is dense, heterogeneous and compartmentalized. In this first in vivo approach, variations in substrate concentrations between compartments was not considered explicitly, though it may have affected the values obtained for the parameters. For example, a higher concentration of phytoene inside lipid droplets compared with the rest of the membrane would give lower-than-expected phytoene concentrations around Crtl enzymes and would thus limit substrate saturation. Preferential localization would also affect measurements as conditions (pH, concentration of ions, etc.) vary between compartments. Iwata-Reuyl et al. found for instance that the measured activity of PaCrtB is 2,000 times lower in the absence of detergents (14), and Fournié et Truan observed that different heterologous CrtI expression systems produced different phytoene saturation patterns (16). Overall, it is crucial to bear in mind that, similarly to in vitro data, enzymatic parameters are only valid under the conditions used to measure them. We therefore argue that enzymatic parameters should not be considered constants since they depend on the microenvironment (pH, ion concentration, temperature, membrane composition, etc.), be that in vitro or in vivo. Still, as mentioned above, these parameters are useful for pathway engineering (44) and an advantage of our in vivo enzymology method is that kinetics parameters are measured under the exact same conditions as the enzyme is expressed.

The final step in our *in vivo* enzymology method involves fitting experimental data with a mathematical model of enzyme kinetics to obtain the corresponding parameters, the same approach as used *in vitro*. Traditional enzymatic models were derived with *in vitro* measurements and assumptions in mind. For instance, the classical Michaelis-Menten relationship applies only at steady-state, a condition clearly met in our experimental setup where all data were collected during exponential growth (i.e. metabolite concentrations and fluxes remain constant over time). However, other assumptions do not hold *in vivo*, in particular regarding the product concentration, which cannot be zero. Nevertheless, for

the enzymes investigated here, where the catalyzed reactions are essentially irreversible, the reverse reaction can be neglected, and the Michaelis-Menten formalism still applies. Another important assumption is that the substrate concentration must be higher than the enzyme concentration. In this study, the *in vivo* substrate/enzyme ratio was between 67 and 12000 for PaCrtB and between 2 and 2500 for XdCrtI (**Supplementary Table S5**). Surprisingly, this criterion was not met for CrtB and CrtI in previous *in vitro* assays (18, 34), with substrate/enzyme ratios between 0.05 and 0.18 for PaCrtB and between 1 and 2.57 for PaCrtI (**Supplementary Table S4**). In our setup, despite some formal assumptions not being fully satisfied, the data were still satisfactorily fit with a Michaelis-Menten equation. This underscores the applicability of our method, from which informative parameters such as the degree of saturation can be inferred to understand *in vivo* enzyme function and to engineer natural and synthetic pathways for biotechnology. The availability of *in vivo* data about enzyme kinetics may also lead to the derivation of specific laws that account for the specificities of *in vivo* studies (e.g., non-negligible product concentrations).

Sixteen years ago, in their review on enzyme function, Dagmar Ringe and Gregory Petsko wrote: "How do enzymes function in a crowded medium of low water activity, where there may be no such thing as a freely diffusing, isolated protein molecule? In vivo enzymology is the logical next step along the road that Phillips, Koshland, and their predecessors and successors have traveled so brilliantly so far." (3). The work presented here, albeit performed in the context of a synthetic metabolic pathway, touches on the difference between kinetic parameters measured *in vitro* and *in vivo* and their interpretation. We notably show how fine-tuning and balancing the expression of the substrate-producing enzyme and the enzyme under study yields datasets from which meaningful and reliable enzymatic parameters can be obtained. Our method is particularly valuable for studying membrane-bound and multimeric enzymes, for which the purification and assay optimization steps of classical in vitro enzymology can be extremely challenging. For membrane-bound enzymes, in vivo enzymology offers a realistic environment devoid of detergents or other interferences, with natural membranes rather than liposomes. By including additional controlled steps this method could be applied to a wide range of variables, such as inhibition and activation parameters (by modulating the pool of regulatory metabolites) and dynamic data with time-course monitoring in response to metabolic or genetic perturbations, and research questions, such as the effects of cell compartments on enzyme efficiency. New formalisms will be required to account for *in vivo* conditions, notably the presence of products, similar enzyme and substrate concentrations, local concentration variations, and molecular fluxes within the cell. We sincerely hope that our work will stimulate further studies delving deeper into how enzymes function in their natural environment.

## **Materials and methods**

#### **Plasmid construction**

Plasmids and primers are listed in **Supplementary Tables S6** and **S7**. Plasmid sequences and annotations are provided in the supplementary data (Supplementary\_data\_Plasmids\_Castano-Cerezo2024.gb). The primers were synthesized by IDT (Leuven, Belgium) and the sequences of PaCrtB, PaCrtI and BtCrtI were codon optimized for yeast and synthetized by Twist Bioscience (San Francisco, California). XdCrtE, XdCrtI and CrtE *from Enterobacter agglomerans* were amplified from pMRI34-CrtE-Gal1-10-HMG1t, YEplac195 YB/I, and pAC-BETA respectively (45–47). Sequences of the mutated versions of the TEF1 promoter TEF1pmut2, TEF1pmut5 and TEF1pmut7 were obtained from Nevoigt et al. (48). Polymerase chain reaction (PCR) was performed using Phusion high fidelity polymerase and Phire Hot start II DNA polymerase (Thermoscientific, Lithuania). DNA fragments were purified using Monarch DNA Gel Extraction Kit from New England Biolabs. DNA fragments were annealed by isothermal assembly using NEBuilder HiFi assembly kit from New Englands Biolabs. Plasmids were cloned and propagated in TOP10 competent *Escherichia coli* cells.

#### **Construction of yeast strains**

All yeast strains used in this study are derived from CEN.PK2-1C and are listed in **Supplementary Table S1**. Yeasts were transformed using Gietz et al.'s high-efficiency transformation protocol (49). Integrative cassettes were obtained by enzyme digestion or PCR and were used without any further purification. Strains were selected using auxotrophy markers or antibiotic resistance at a concentration of 400  $\mu$ g/mL. Antibiotic resistance recycling was performed using vector pSH63 as described in the literature (50). Genome integration was verified by colony PCR using the primers listed in **Supplementary Table S7**. Genomic DNA was extracted using DNA release from ThermoFisher Scientific.

#### Media and culture conditions

All strains were grown in modified synthetic Verduyn media containing glucose (111 mM), NH<sub>4</sub>Cl (75 mM), KH<sub>2</sub>PO<sub>4</sub> (22 mM), MgSO<sub>4</sub> (0.4 mM) and CSM (ForMedium LTD, Hunstaton, England) at pH 5.0 (51). Sterilization was performed by filtration. Fresh colonies from selective plates were precultured in 350  $\mu$ L complete synthetic medium at 28 °C for 8 hours and these cells were used to inoculate cultures with a 1:5 medium:flask proportion to an initial OD<sub>600nm</sub> of 0.002, grown at 200 rpm at 28 °C.

#### **Carotene quantification**

Samples (10 or 20 mL) of yeast culture were harvested with an OD<sub>600nm</sub> of approximately 5, centrifuged, and washed with 1 mL of MilliQ water. Cell pellets were freeze-dried and stored at -80 °C until extracted.  $\beta$ -apocarotenal solution (40 µL, 50 µM) was added to the dried cells, and carotenes were extracted with glass beads and 500 µL of acetone in three 20 s rounds of agitation at 0.05 m/s with a FastPrep FP120 cell disruptor (ThermoFisher). The acetone phase was transferred to a new tube and the extraction was repeated twice. Acetone extracts were pooled, centrifuged, dried under nitrogen flux, and resuspended in acetone for HPLC analysis. Analyses were carried out on a Thermo Scientific Vanquish Focused UHPLC Plus system with DAD HL. Extract samples (5 µL) were injected into a YMC carotenoid column (100 × 2.0 mm and 3 µm particle size) equipped with a precolumn (100 × 2.0 mm and 3 µm particle size). The mobile phases used to separate and quantify phytoene, lycopene and  $\beta$ -apocarotenal from ergosterol and derivatives were mixtures of (A) methanol/water (95:5) and (B)

dichloromethane. The flow was 0.25 mL/min with the following gradient: 0–0.1 min 5 % B, 0.1–0.5 min 20 % B, 0.5–2 min 60 % B, 2–5 min 80 % B, 5–8 min 80 % B and 8–11 min 5 % B. The absorbance from 210 to 600 nm was measured throughout the run with a data collection rate of 2 Hz and a response time of 2 s. The phytoene concentration was deduced from its absorbance at 286 nm and lycopene and  $\beta$ -apocarotenal concentrations from the absorbance at 478 nm. The reference wavelength (600 nm) was subtracted from each of the wavelengths used for metabolite quantification. Phytoene and lycopene production fluxes were determined by multiplying their concentrations by the cell growth rate.

#### **GGPP** quantification

GGPP was quantified as detailed previously (52). Briefly, 10 mL of broth was filtered through 0.45  $\mu$ m Sartolon polyamide (Sartorius, Goettingen, Germany) and washed with 5 mL of fresh culture medium (without glucose). The filters were rapidly plunged into liquid nitrogen and then stored at -80 °C until extraction. Intracellular GGPP was extracted by incubating filters in closed glass tubes containing 5 mL of an isopropanol/H<sub>2</sub>0 NH<sub>4</sub>HCO<sub>3</sub> 100 mM (50/50) mixture at 70 °C for 10 min. For absolute GGPP quantification, 50  $\mu$ L of <sup>13</sup>C internal standard were added to each extract. Cellular extracts were cooled on ice and sonicated for 1 min. Cell debris was removed by centrifugation (5000 g, 4 °C, 5 min). Supernatants were evaporated overnight (SC110A SpeedVac Plus, ThermoFisher, Waltham, MA, USA), resuspended in 200  $\mu$ L of methanol:NH<sub>4</sub>OH 10 mM (7:3) at pH 9.5 and stored at -80 °C until analysis.

Analyses were carried out on a LC–MS platform composed of a Thermo Scientific Vanquish Focused UHPLC Plus system with DAD HL, coupled to a Thermo Scientific Q Exactive Plus hybrid quadrupole-Orbitrap mass spectrometer (ThermoFisher), as detailed previously (52). Calibration mixtures (prepared at concentrations from 0.08 nM to 10  $\mu$ M) were used to construct calibration curves from which the absolute concentration of GGPP in the samples was determined.

#### **GGOH** quantification

To determine the production of geranylgeraniol in the yeast cultures, strains were cultured in 5 mL synthetic complete media with 500  $\mu$ L of dodecane for 3 days at 28 °C with 200 rpm agitation. Dodecane was then harvested and analyzed by GC and cell biomass was determined. A sample (0.5  $\mu$ L) of the dodecane fraction was directly injected into a GC Shimadzu 2025 equipped with FID detection and an RTX-5 column (15 m, 0.25 mm ID, 0.25  $\mu$ m particle, Restek Corp). The different derivative isoprenoid alcohols were separated using the splitless injection method with a temperature gradient (115 °C for 1 min, 70 °C/min to 250 °C, hold at 300 °C for 5 min). The make-up gas was nitrogen, with a flow of 30 mL/min.

#### Western blot

Protein extracts were prepared as described by Zhang et al. (53). Briefly, 1.5  $OD_{600nm}$  of pelleted cells were pre-treated with 100 µL of a 2 M lithium acetate cold solution, and left to stand for 5 min, followed by 5 min centrifugation at 5000 g, 4 °C. The supernatant was discarded and 100 µL of a 0.4 M solution of NaOH was added. After gentle resuspension, and 5 min standing on ice, the samples were centrifuged for 5 min at 4 °C. After discarding supernatants, the pellets were vigorously vortexed with 60 µL of bromophenol blue loading dye solution with 5 % β-mercaptoethanol. After denaturation for 10 min at 99 °C, 5 µL of each sample was loaded onto 10 % SDS page gel. Semi-dry transfer was performed on PVDF membrane (Merck Millipore, Darmstadt, Germany) using a Trans-Blot SD Cell BioRad apparatus (18 V during 20 min), and 5 % bovine milk in TBS as blocking agent. Incubations were

performed with mouse anti-FLAG or mouse anti-V5 (ThermoFisher Scientific), and secondary antimouse IgG coupled with alkaline phosphatase (ThermoFisher Scientific), diluted according to manufacturer instructions. Proteins were detected by incubation in BCIP/NBT AP substrate buffer (Sigma-Aldrich, St. Louis, MO, USA).

#### Proteomics

For cell disruption,  $10^8$  cells were dissolved in 200 µL of lysis buffer (0.1 M NaOH, 2 % SDS, 2 % 2mercaptoethanol, 0.05 M EDTA), heated at 90 °C for 10 min and neutralized with 5 µL of 4 M acetic acid. Glass beads were added and the samples were vortexed at 4 °C for 30 min. Cell debris were pelleted by centrifugation at 3000 g for 10 min and the protein concentration of the supernatant was determined using the Bradford assay. Protein aliquots (400 µg) were cleaned by methanol chloroform precipitation. The protein samples were dissolved in 5 µL of 6 M guanidinium chloride, 5 µL of 0.1 M dithiothreitol (DTT), and 100 µL of 50 mM TEAB and diluted to a protein concentration of 2 µg/µL with 90 µL of water. Absolute protein quantification was performed using heavy isotope labeled tryptic peptides as internal standards. Protein lysate aliquots (20 µg) were spiked with a mixture of ten AQUA peptides, C-terminally labeled with heavy lysine or arginine (ThermoFisher Scientific) with concentrations of 50 fmol/µg, 10 fmol/µg, 5 fmol/µg and 0 fmol/µg. The samples were reduced by adding 2 µL of 0.1 M DTT and incubating at 60 °C for 1 h, alkylated with 2 µL of chloroacetamide 0.5 M at room temperature for 30 min, and digested with 0.5 µg of trypsin overnight. Digestion was stopped by adding 5 µL of 1 % TFA and the samples were cleaned using 100 µL C18 tips. The extracts were lyophilized, reconstituted in 20 µL of eluent A, and transferred to HPLC vials.

Samples were analyzed in triplicate using an UltiMate 3000 UHLC system coupled to a Q Exactive Plus mass spectrometer (ThermoFisher Scientific). Approximately 0.6  $\mu$ g of peptides were loaded on a C18 precolumn (PepMap100, 5  $\mu$ m, 300 Å, ThermoFisher Scientific) and separated on a PepMap RSLC C18 column (50 cm  $\times$  75  $\mu$ m, 2  $\mu$ m, 100 Å, ThermoFisher Scientific) with a 1.5 h gradient with eluent A (Water, 0.05 % FA) and eluent B (80 % ACN, 0.04 % FA) with a flow rate of 0.3  $\mu$ L/min. The peptides were first desalted for 4 min at 5 % B, then separated with a gradient to 50 % B over 65 min, to 90 % B in 3 min, held at 90 % B for 8 min, and then equilibrated for 8 min at 4 % B.

For targeted absolute quantification, full MS spectra measurements were followed by parallel reaction monitoring (PRM) of targeted heavy AQUA peptides and the corresponding light, native peptide of the proteins of interest. The full MS spectra were acquired in profile mode with the following settings: resolution of 70000, mass range of 350–950 m/z, AGC target of 10<sup>6</sup>, and 80 ms maximum injection time. PRM MS2 spectra were acquired with an isolation window of 1.6 m/z, 17500 resolution, AGC target of 10<sup>5</sup>, 80 ms injection time, and an NCE of 27. The inclusion list contained 26 entries covering the light and heavy species of 10 peptides at the most intense charge state (2+ or 3+).

Database searches were performed with Proteome Discoverer (version 3.01.27; ThermoFisher). The raw data were compared with the UniProt protein databases of *S. cerevisiae* strain CEN.PK113-7D (UniProt 02.2023; 5,439 entries), and common contaminants using Sequest HT and Chimerys search algorithms. The Sequest search parameters were a semi-tryptic protease specificity with a maximum of 2 missed cleavage sites. The precursor mass tolerance was 8 ppm and the fragment mass tolerance was 0.02 Da. Oxidation of methionine and acetylation of protein N-termini were allowed as dynamic modifications. Carbamidomethylation of cysteine was set as a static modification. Chimerys database searches were performed with default settings. Percolator q-values were used to restrict the false

discovery rate (FDR) of peptide spectrum matches to 0.01. The FDR of peptide and protein identifications was restricted to 1 % and strict parsimony principles were applied to protein grouping.

A spectral library of the AQUA peptides was generated using Proteome Discoverer with a sample containing only 500 fmol of the ten AQUA peptides, in addition to the PRM analyses of the spiked yeast samples. The PRM data of three concentrations analyzed in triplicate were imported and all transitions were reviewed using Skyline (54). Between four and nine transitions without interference were chosen for each peptide and the ratio of light and heavy peptides of the sum of transitions was calculated for absolute quantification. The list of used peptides and all corresponding transitions are provided in **Supplementary Table S8**.

#### **Calculation of absolute concentrations**

Protein and metabolite concentrations are expressed as absolute intracellular concentrations. Metabolite concentrations initially expressed in  $\mu$ mol/g DCW were converted into absolute intracellular concentrations using a conversion factor of  $6.59 \times 10^{10}$  cells/g DCW (**Supplementary Figure S5**) and a cellular volume of 66  $\mu$ m<sup>3</sup> (55). The intracellular concentrations of PaCrtB and XdCrtI were calculated using a conversion factor of 0.63 g protein/g DCW for yeast grown on ammonium sulfate medium (56). All calculations for data conversion are provided in the Supplementary data.

#### Calculation of enzyme parameters

Enzymatic parameters (V<sub>max</sub> and K<sub>M</sub>) were determined by fitting a Michaelis-Menten equation to the measured relationships between substrate concentrations and reaction rates. Uncertainties on fitted parameters were determined using a Monte-Carlo approach. Briefly, 1000 simulated noisy datasets were generated (where the noise was determined as the standard deviation of residuals obtained for the fit of the experimental dataset), and the mean value, standard deviation, and 95 % confidence intervals of each parameter were determined from the distribution of values obtained for the 1000 datasets. For each enzyme, we verified that all parameters were identifiable based on the covariance matrix and on the results of the Monte-Carlo analysis. The code for parameter estimation and statistical analysis is provided as a Jupyter notebook at <a href="https://github.com/MetaSys-LISBP/in\_vivo\_enzymatic\_parameters.">https://github.com/MetaSys-LISBP/in\_vivo\_enzymatic\_parameters.</a>

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## **Competing interests**

The authors declare no competing interests.

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## Contributions

Funding acquisition: CC, SH, PM, GT; Project administration: CC, SH, GT; Conceptualization: CC, SCC, SH, PM, GT; Investigation: SCC, AC, HK, CT, AT, VG, CC, PM; Methodology: SCC, AC, HB, CT, FB, AT, VG, CC, SH, LGA, PM, GT; Software: PM; Formal Analysis: SCC, AC, HB, CT, AT, LGA, PM, GT; Visualization: SCC, PM, GT; Writing – original draft: SCC, PM, GT; Writing – review & editing: SCC, AC, HB, CT, FB, AT, VG, CC, SH, LGA, PM, GT; Supervision: SCC, VG, PM, GT.

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