Titration of *in-cellula* affinities of protein-protein interactions

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A genetic assay permits simultaneous quantification of two interacting proteins and their bound fraction at the single-cell level using flow cytometry. *In-cellula* affinities of proteinprotein interactions can be extracted from the acquired data through a titration-like analysis. The applicability of this approach is demonstrated on a diverse set of interactions with proteins from different families and organisms and with *in-vitro* dissociation constants ranging from picomolar to micromolar.

The quest for methods that permit rapid and reliable determination of the affinity of 33 34 protein-protein interactions (PPI) is unbroken. In contrast to biochemical *in-vitro* methods such 35 as Isothermal Titration Calorimetry (ITC) and Surface Plasmon Resonance (SPR) that require 36 purified proteins, quantitative genetic assays rely on the expression of the proteins of interest 37 in cells. Many of these assays¹⁻⁵ are inspired by the yeast two-hybrid (Y2H) technique⁶ which 38 is based on the *in-cellula* expression of two proteins, usually named Bait and Prey, fused to an 39 DNA-binding domain (BD) and an activation domain (AD), respectively. Upon physical 40 interaction of the BD-Bait and AD-Prey proteins, a functional transcription factor is 41 reconstituted that drives the expression of a reporter gene. The stronger the interaction, the 42 higher should be the expression level of the reporter.⁷ However, the expression level of the AD-43 Bait and BD-Prey play an important role, too.⁸

We recently introduced a quantitative yeast-two hybrid system (qY2H) that permits for the first time simultaneous quantification of BD-Bait, AD-Prey and the reporter at the singlecell level without the need of any antibodies or purified proteins.⁸ Instead, we take advantage of fluorescent fusion proteins (Fig. 1A) that can be detected by standard flow cytometers. Here we show how this qY2H method can be exploited to perform *in-cellula* affinity titrations by applying the following two important improvements:

50 1) Cellular contents of fluorescent proteins are determined in units of Molecules of
51 Equivalent Soluble Fluorochrome (MESF), so that measured quantities become independent of

the applied apparatus setup. It facilitates the future transferability of the qY2H measurements to other flow cytometers and allows researchers to consistently compare their results. Our reference fluorochrome is the yeast Enhanced Green Fluorescent protein (yEGFP) for which commercial calibration beads exist. The fluorescence intensity of TagRFP is converted into units of MESF of EGFP using independent calibration experiments with a fluorescent tandem protein BD-TagRFP-EGFP (see "Methods").

2) We analyze the data by a titration-like procedure which allows the straightforward extraction of *in-cellula* dissociation constants for Bait:Prey interactions. In a proof of concept, we apply this *in-cellula* titration approach to a diverse set of PPIs with dissociation constants ranging from 117 pM to 17 μ M (Table 1). As in *in-vitro* SPR experiments, each PPI can be measured by Y2H in two different orientations (by exchanging Bait and Prey). Here we study only the orientation that produced the higher reporter level.⁸ This orientation is considered as the molecular configuration with the higher accessibility of the PPI binding interface.⁷

65 In our qYH2 experiments, diploid yeast cells with constitutive expression of BD-Bait 66 and induced expression of AD-Prey are cultured for two hours. Then, their fluorescence 67 intensity is measured by flow cytometer in the three channels corresponding to TagRFP (BD-68 Bait), EGFP (AD-Prey), and TagBFP (reporter). Due to phenotypic variations, BD-Bait and 69 AD-Prey are expressed at different levels among these cells which can be exploited to "prepare 70 samples" for a titration. By gating, we can split the global heterogeneous ensemble of cells into 71 several homogenous subensembles (bins). Each bin contains only cells within two specific, 72 narrow intervals of red and green fluorescence intensity centered at values R and G, 73 respectively. Assuming a linear relationship between fluorescence intensity and molecule 74 numbers, R and G can be considered as measures for the mean cellular content of BD-Bait and 75 AD-Prey in the corresponding bin.

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With the mean value of the blue fluorescence intensity, we can calculate for each bin

77 the normalized reporter level φ . It is obtained by forming the ratio of the expression level for the interaction of interest, E_{interaction} (Fig. 1A) and the level for a covalent BD-AD fusion, 78 $E_{\text{colvalent}}$ (Fig. 1B). This normalization renders φ dimensionless and independent of the 79 80 acquisition apparatus (assuming again a linear relationship between molecule number and 81 fluorescence intensity). Most importantly, we consider that φ reflects the time-averaged 82 fraction of BD-Bait bound by AD-Prey during the reaction (as explained in the caption of Fig. 83 1). Thus, titration curves can be obtained when φ is plotted as a function of G while keeping R 84 fixed (Fig. 1C). The curves can be fitted with the following Langmuir-type equation:

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$$\varphi(G) \cong \beta \frac{G/\alpha}{K'_{\rm d} + R + G/\alpha}$$

where K'_{d} is the *in-cellula* dissociation constant (in units of MESF of EGFP) and α and β are 87 dimensionless parameters that empirically account for the fact that φ is a time-integrated 88 89 property. The parameter α reduces the final cellular content of AD-Prey (measured at the end 90 of the reaction, G) to the time-averaged content (over the entire reaction course, $\langle G \rangle$). Since the induction kinetics under the GAL1-promotor in yeast⁹ displays a quadratic-like time 91 dependence (for short induction times), a reasonable choice for α is $3 [\langle G \rangle =_0 \int_0^1 G t^2 dt = G/3]$. 92 The prefactor β , on the other hand, integrates differences in the expression kinetics of the 93 94 reporter for $E_{\text{interaction}}$ (induced expression) and E_{covalent} (constitutive expression). It can be determined experimentally by monitoring φ for $G \rightarrow \infty$ using a high-affinity couple (such as 95 96 BD-Barstar29F/AD-BarnaseH102A).

Eq. 1

97 We recommend that the titrations are carried out with the lowest possible value of 98 $R=R_{min}$ (as defined by the detection limit of TagRFP by flow cytometery, see "Methods"). It 99 limits overexpression and associated protein burden effects.¹⁰ Furthermore, the auto-activation 100 potential of the BD-Bait fusion is kept at a minimum, too.¹¹ Also, it mimics the condition of *in*- 101 *vitro* affinity titration experiments where the concentration of the titrated species (here BD-102 Bait) is kept fixed and as low as possible to avoid saturation effects. For the titrations with 103 $R=R_{min}$ the parameters $\alpha=3$ and $\beta=1.35$ were used to extract the K'_{d} -values.

Despite substantial differences between our *in-cellula* system and *in-vitro* setups (as previously discussed⁸ in detail), the *in-cellula* affinities strongly correlate with those from *invitro* measurements (R^2 =0.91, Fig. 1D). The slope of the regression line is 0.84. Other *in-cellula* assays usually find lower correlation coefficients (< 0.9) and significantly lower values for the slope of the regression line (0.2-0.6).^{1–5} This is even more remarkable if one considers that the tested set of PPIs in this work is significantly more diverse. It may indicate a higher sensitivity for the qY2H titration approach; more testing will be necessary to confirm this surmise.

111 The presented protocol is robust as witnessed by the small error bars in the titration 112 curves (Fig. 1c). All steps of the protocol have been optimized in liquid phase that can be easily 113 automated for the use of microplates and integrated within robotic pipelines. It sets the stage for high-throughput affinity screenings of PPIs using cross-mating approaches¹² with libraries 114 of yeast clones. As an outlook, affinity-based networks¹³ can be created by attributing weights 115 116 to the PPI edges according to their effective affinities. It contrasts standard Y2H screens that 117 vield networks with only binary information (YES or NO). The topology of edge-weighted 118 networks may help identifying key pathways within the network, and how they change as a 119 function of environmental conditions (stress, metabolism, etc). Thus, we anticipate that high-120 throughput qY2H affinity data would boost the modelling of interactomes and thereby advance 121 significantly systems biology.

122 Methods

123 The qY2H experiments, acquisitions by flow cytometry and analyses were carried out 124 as described in our previous study⁸ with the following particularities. About 10^7 cells were 125 cultured per experiment and interaction (including the covalent BD-AD fusion and the control 126 sample BD-Empty / AD-Empty). To ensure that these cells have been indeed transfected with 127 all three vectors, we selected for the analysis large (=growing) cells with a forward scatter range 128 75 000 < FSC-H < 125 000; "H" indicates signal height. Furthermore, only cells with a red 129 fluorescence intensity of 800 \pm 100 TagRFP-H were analysed. This bin is located just above 130 the 95% threshold of the non-fluorescent cells,⁸ and therefore defines R_{min} .

131 The mean Tag BFP-H value was then calculated for bins of varying *G* values from -500
132 to 25500 yEGFP-H (bin size 1000). For each bin we calculated:

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$$\varphi(G) = \frac{\langle F_{\text{BFP,interaction}} \rangle_G - \langle F_{\text{BFP,CTRL}} \rangle_G}{\langle F_{\text{BFP,covalent}} \rangle_G - \langle F_{\text{BFP,CTRL}} \rangle_G}$$

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135 where $\langle F_{BFP,X} \rangle_G$ is the mean blue fluorescence intensity. The subscripted X refers to the 136 physical interaction, covalent fusion or control couple. The control couple BD-Empty / AD-137 Empty⁸ permits to remove the background of the reporter system.

Eq. 2

Finally, *G* values were converted into MESF of EGFP using calibration beads (Ozyme, reference 632594) following the manufactor's protocol. For the conversion of R_{min} , we performed independent calibration measurements with diploid yeast cells expressing the fluorescent tandem fusion protein BD-TagRFP-yEGFP (under the same condition as the qY2H experiments). Cells with a red fluorescence intensity of 800 ± 100 TagRFP-H displayed a mean green fluorescence intensity of 370 000 MESF of EGFP (= R_{min} used in Eq. 1).

Experiments and analyses were performed at least three times for each interaction and averaged titration curves were least-square fitted with Eq. 1.

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149 **Figures & Tables**

150 Table 1: Investigated protein-protein interactions and their *in-vitro* affinities (Kd).

Bait proteins							Prey proteins				
Organism	Family	Name	Mutant	MW	K _d (nM)	Symbol	Organism	Family	Name	Mutant	MW
B. amylo- liquefacien s	RNAse inhibitor	Barstar	WT	10	320 ^a		B. amylo- liquefaciens	RNAse	Barnase	H102A	12
			Y29A	10	420 ^a						
			Y29F	10	117 ^a	•					
			W38F	10	4 000 ^a						
			D35A	10	$25\;000^{\text{ h}}$	0					
			D39A	10	420 000 ^b						
H. sapiens	GTPase	HRas	G12V & C186A	21	122 000 °		H. sapiens	Kinase	CRaf RBD	WT	9
					11 000 ^d	Δ				A85K	9
H. sapiens	Kinase regulatory subunit	CksHs1	WT	10	77 000 ^e		H. sapiens	Kinase	CDK2	WT	34
E.coli	β-Lactamase	TEM	WT	31	$15\ 000\ ^{\rm f}$	•	S. clavuligerus	β-Lactamase inhibitor	BLIP1	WT	21
HIV1	Virulence factor	Nef	LAI	23	11 400 000 ^g		H. sapiens	Kinase	SRC SH3	WT	7
H. sapiens	Adapter	Grb2 SH3	WT	7	17 000 000 ⁱ	\$	M. musculus	Nucleotide exchange factor	Vav1 SH3	WT	8

^a ITC, 50mM Tris/HCl, pH 8 at 25°C.¹⁴

^b Mean values from two studies ^{14,15} with ITC, 24mM Hepes, pH 8, 1 mM DTT at 25°C. ^c Mean values from four studies of Ras G12V (without the membrane anchor): SPR, 50 mM Tris/HCl, pH 7.4, 100mM NaCl, 5 mM

MgCl₂;¹⁶ SPR, 50 mM Tris/HCl, pH 7.4, 100mM NaCl, 5 mM MgCl₂;¹⁷ SPR, 10 mM Hepes, pH 7.4, 150 mM NaCl, 2 mM MgCl₂, and

0.01% Nonidet P-40 25°C;¹⁸ ITC, 50 mM Hepes, pH 7.4, 125 mM NaCl, 5 mM MgCl₂, 25°C.¹⁹ d ITC, 50 mM Hepes, pH 7.4, 125 mM NaCl, 5 mM MgCl₂, 25°C.¹⁹ with HRas WT loaded with a GTP-analogue. The mutant HRas G12V is known to decrease the dissociation constant for the interaction with

CRaf RBD WT by a factor of 11.²⁰ The given value applies the same correction factor. ^e SPR, 10 mM Hepes, 3.4 mM EDTA, 150 mM NaCl, 0.001% surfactant P20, pH 7.4.²¹

^fSPR, 10 mM Hepes, 3.4 mM EDTA, 150 mM NaCl, 0.05% surfactant P20, pH 7.4.²²

^g ITC, 20 mM phosphate buffer, pH 7.5, 150 mM NaCl, 2 mM EGTA, and 5 mM DTT, 25°C.²³

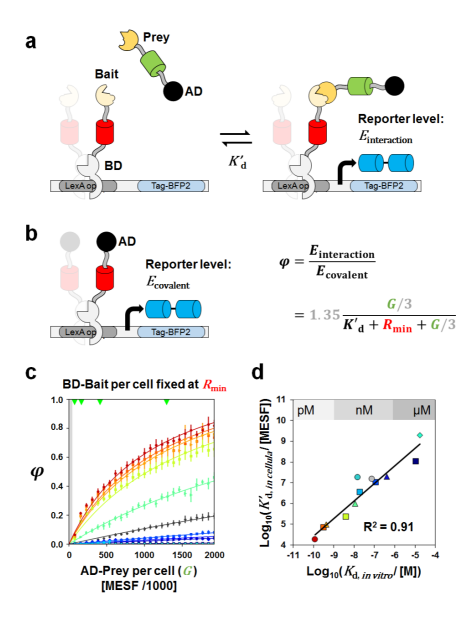
^h SPR, 10 mM Hepes-Na, pH 7.4, 0.15 M NaCl, 3 mM EDTA, and 0.005% (v/v) Tween 20, 25 °C.²⁴ ⁱ SPR, 25°C, 25

 $\begin{array}{c} 151\\ 152\\ 153\\ 154\\ 155\\ 156\\ 157\\ 158\\ 159\\ 160\\ 161\\ 162\\ 163\\ 164 \end{array}$

h Free-energy calculations.8

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167 **Figure 1**.



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169 a) In our qY2H system, red-fluorescent BD-Bait interacts with green-fluorescent AD-Prey to reconstitute a 170 transcription factor that drives the expression of a blue-fluorescent reporter. Our hypothesis is that the expression 171 level of the reporter, E_{interacting}, reflects the number of BD-molecules bound to the promotor corrected by the 172 fraction of BD-Bait bound to AD-Prey. This fraction is influenced by the affinity between the Bait and the Prey, 173 but also by the expression levels of BD-Bait and AD-Prey. b) When the activation domain is covalently linked to 174 the DNA-binding domain, the expression level E_{covalent} depends only on the number of BD-Bait molecules bound 175 the promotor. Thus, when forming the quantity φ by dividing $E_{\text{interacting}}$ with E_{covalent} , we obtain a measure for the fraction of BD-molecules bound by an AD-Prey molecule. To determine Ecovalent, we constructed a BD-AD fusion 176 177 protein. Unfortunately, the activation domain B42 (as used in a) turned out to be toxic for our yeast strains when 178 used in the BD-AD construct. Instead, we used the activation domain B112. Difference in the activation potential 179 between B42 and B112 are integrated in the parameter β of Eq. 1. c) The quantity φ can be monitored as a function 180 of different levels of EGFP Molecules of Equivalent Soluble Fluorochrome (MESF) corresponding to different 181 cellular levels of AD-Prey. In these titrations, the level of BD-Bait is kept fixed at the lowest possible value (see 182 "Methods"). For the interaction TEM/BLIP1 (cyan line) the titration can be preformed only up to one third of the 183 titrant quantity due to expression problems of AD-BLIP1.8 Green triangles at the top vertical axis indicate the 184 position of used calibration beads. d) When the titration curves are fitted with Eq. 1, we can extract the dissociation 185 constant in units of MESF (K_d '). The estimated K_d '-values show a remarkable correlation with the dissociation 186 constants measured from alternative in-vitro experiments (Table 1).

- 189 1. Dutta, S., Koide, A. & Koide, S. High-throughput Analysis of the Protein Sequence-
- 190 Stability Landscape using a Quantitative Yeast Surface Two-hybrid System and Fragment
- 191 Reconstitution. J. Mol. Biol. 382, 721–733 (2008).
- 192 2. Jeong, K. J., Seo, M. J., Iverson, B. L. & Georgiou, G. APEx 2-hybrid, a quantitative
- 193 protein-protein interaction assay for antibody discovery and engineering. *Proc. Natl.*
- 194 *Acad. Sci.* **104**, 8247–8252 (2007).
- Colas, P., Cohen, B., Ferrigno, P. K., Silver, P. A. & Brent, R. Targeted modification and
 transportation of cellular proteins. *Proc Natl Acad Sci USA* 97, 13720–13725 (2000).
- 197 4. Hu, X., Kang, S., Chen, X., Shoemaker, C. B. & Jin, M. M. Yeast Surface Two-hybrid for
- 198 Quantitative *in Vivo* Detection of Protein-Protein Interactions via the Secretory Pathway.
- 199 *J. Biol. Chem.* **284**, 16369–16376 (2009).
- 200 5. Younger, D., Berger, S., Baker, D. & Klavins, E. High-throughput characterization of
- 201 protein–protein interactions by reprogramming yeast mating. *Proc. Natl. Acad. Sci.* **114**,

202 12166–12171 (2017).

- 203 6. Fields, S. & Song, O. A novel genetic system to detect protein protein interactions.
 204 *Nature* 340, 245–246 (1989).
- 205 7. Estojak, J., Brent, R. & Golemis, E. A. Correlation of two-hybrid affinity data with in
 206 vitro measurements. *Mol. Cell. Biol.* 15, 5820–5829 (1995).
- 207 8. Cluet, D. et al. A quantitative tri-fluorescent yeast two-hybrid system: from flow
- 208 cytometry to in-cellula affinities. *Mol. Cell. Proteomics* (2020)
- 209 doi:10.1074/mcp.TIR119.001692.
- 210 9. Li, J. et al. Green fluorescent protein in Saccharomyces cerevisiae: Real-time studies of
- 211 the GAL1 promoter. *Biotechnol. Bioeng.* **70**, 187–196 (2000).
- 212 10. Bolognesi, B. & Lehner, B. Reaching the limit. *eLife* 7,.

¹⁸⁸ References

- 213 11. Rajagopala, S. V. & Uetz, P. Analysis of Protein–Protein Interactions Using High-
- 214 Throughput Yeast Two-Hybrid Screens. in *Network Biology* (eds. Cagney, G. & Emili,
- A.) vol. 781 1–29 (Humana Press, 2011).
- 216 12. Kolonin, M. G., Zhong, J. & Finley, R. L. Interaction mating methods in two-hybrid
- 217 systems. *Methods Enzymol.* **328**, 26–46 (2000).
- 218 13. Gromiha, K. Y. and M. M. Analysis of protein-protein interaction networks based on
- 219 binding affinity. *Current Protein & Peptide Science*
- 220 http://www.eurekaselect.com/135165/article (2016).
- 14. Schreiber, G. & Fersht, A. R. Energetics of protein-protein interactions: Analysis of the
- Barnase-Barstar interface by single mutations and double mutant cycles. J. Mol. Biol.
- **223 248**, 478–486 (1995).
- 15. Frisch, C., Schreiber, G., Johnson, C. M. & Fersht, A. R. Thermodynamics of the
- interaction of barnase and barstar: changes in free energy versus changes in enthalpy on

226 mutation 1 1Edited by J. Karn. J. Mol. Biol. 267, 696–706 (1997).

- 16. Herrmann, C., Horn, G., Spaargaren, M. & Wittinghofer, A. Differential Interaction of the
- 228 Ras Family GTP-binding Proteins H-Ras, Rap1A, and R-Ras with the Putative Effector
- 229 Molecules Raf Kinase and Ral-Guanine Nucleotide Exchange Factor. J. Biol. Chem. 271,
- 230 6794–6800 (1996).
- 231 17. Block, C., Janknecht, R., Herrmann, C., Nassar, N. & Wittinghofer, A. Quantitative
- structure-activity analysis correlating Ras/Raf interaction in vitro to Raf activation in
- 233 vivo. Nat. Struct. Mol. Biol. 3, 244–251 (1996).
- 18. Fischer, A. *et al.* B- and C-RAF display essential differences in their binding to Ras: the
 isotype-specific N terminus of B-RAF facilitates Ras binding. *J. Biol. Chem.* 282, 26503–
- 236 26516 (2007).

- 237 19. Kiel, C. et al. Improved Binding of Raf to Ras-GDP Is Correlated with Biological
- 238 Activity. J. Biol. Chem. 284, 31893–31902 (2009).
- 239 20. Kiel, C. Untersuchung von Ras/Effektor-Komplexen mit gezielt veränderten
- 240 elektrostatischen Eigenschaftens. (Dissertation, Ruhr-Universität Bochum, Fachbereich
- 241 Biochemie, 2003).
- 242 21. Bourne, Y. et al. Crystal Structure and Mutational Analysis of the Human CDK2 Kinase
- 243 Complex with Cell Cycle–Regulatory Protein CksHs1. *Cell* **84**, 863–874 (1996).
- 244 22. Albeck, S. & Schreiber, G. Biophysical Characterization of the Interaction of the β -
- 245 Lactamase TEM-1 with Its Protein Inhibitor BLIP[†]. *Biochemistry (Mosc.)* **38**, 11–21
- 246 (1999).
- 247 23. Arold, S. *et al.* RT Loop Flexibility Enhances the Specificity of Src Family SH3 Domains
 248 for HIV-1 Nef^{†,‡}. *Biochemistry (Mosc.)* **37**, 14683–14691 (1998).
- 249 24. Sato, Y. et al. Characterization of the Interaction between Recombinant Human Peroxin
- 250 Pex3p and Pex19p: IDENTIFICATION OF TRP-104 IN Pex3p AS A CRITICAL
- 251 *RESIDUE FOR THE INTERACTION. J. Biol. Chem.* **283**, 6136–6144 (2008).
- 252 25. Nishida, M. Novel recognition mode between Vav and Grb2 SH3 domains. *EMBO J.* 20,
 253 2995–3007 (2001).