Control of protein phosphorylation with a genetically encoded photocaged amino acid

Edward A Lemke1,3, Daniel Summerer1,3, Bernhard H Geierstanger2, Scott M Brittain2 & Peter G Schultz1,2

We genetically encoded the photocaged amino acid 4,5-dimethoxy-2-nitrobenzylserine (DMNB-Ser, 1) in Saccharomyces cerevisiae in response to the amber nonsense codon TAG. This amino acid was converted to serine in living cells by irradiation with relatively low-energy blue light and was used to noninvasively photoactivate phosphorylation of the transcription factor Pho4, which controls the cellular response to inorganic phosphate.1 When substituted at phosphoserine sites with the endogenous tRNAs or aaRSs of the host cell14,15. This strategy offers the advantage that the modified proteins and peptides are generated inside the cell directly by the translational machinery and are genetically targeted to the desired site. Once established, such systems can be easily generalized to virtually any protein in the cell and allow photocontrol of protein function based on single amino acid modifications. However, so far only amino acids with the photolabile o-nitrobenzyl group14,15, which are photoactivated with UV light (<365 nm, Supplementary Fig. 1 online), have been genetically encoded. The use of UV light restricts application of this technology for cellular studies because high-intensity UV light causes photoreactivities in nucleic acids, destroys disulphide bonds and can cause other cellular damage. To overcome these limitations, we have genetically encoded 1 (Fig. 1a) in S. cerevisiae in response to the amber nonsense mutant TAG. In contrast to the o-nitrobenzyl group, the larger DMNB group has a higher quantum yield for photocleavage. More importantly, DMNB has a substantially red-shifted absorption spectrum, which allows photolysis of living cells with visible blue light from a standard laser scanning microscope (Supplementary Fig. 1)16. More...
Figure 2  In vivo photolysis of the DMNB-Ser Pho4-GFP mutants. (a) Cytoplasmic localization of WT Pho4-GFP grown in high-Pi medium and starved for phosphate. (b) S3→Ala and S2→Ala Pho4-GFP mutants localize to the nucleus in high Pi. (c) The S3→TAG Pho4-GFP mutant localizes in the nucleus even in high Pi medium if grown in the presence of 1. (d) Fluorescent nuclei in the upper part (photolyzed part, above yellow dashed line) of the image for the S3→1 Pho4-GFP show a strong loss in fluorescence in the nucleus. (e) Magnification of two cells before and after photolysis and corresponding line scan. (f) Photolyzed S2→1 Pho4-GFP mutant cells grown in high Pi show a strong loss of fluorescence originating from the nucleus. Signal amplification for images in a and b was reduced about three-fold to compensate for a ~seven-fold increased expression of protein. Scale bars are 10 μm.

over, no photocaged serine derivatives have been genetically encoded so far, and given that serine is critical to the function of many proteins—for example, as a catalytic residue, hydrogen bonding partner or site of post-translational modification—1 should have broad applicability in cellular and biochemical studies.

We used a previously reported orthogonal E. coli amber suppressor tRNA/aaRS (tRNA^Leu^CUA/LeuRS)^12 to incorporate 1 into proteins in yeast and generated a library of ~10^7 LeuRS mutants in which residues Met40, Leu41, Tyr499, Tyr527 and His537 in the leucine binding site were randomized (see ref. 17 for X-ray structure). Next, we used a selection scheme to isolate a LeuRS mutant that selectively uses 1. The selection is based on suppression of amber (TAG) codons in essential or toxic genes, and it has previously been published work, this mutation results in increased hydrolysis of the mutant LeuRS BH5, we mutated residue Thr252 of the editing partner or site of post-translational modification—

1—should have its translocation to the cytoplasm (Fig. 2a). Phosphorylation of Ser223 (termed S6) regulates transcription, whereas phosphorylation of Ser114 (S2) and Ser128 (S3) triggers nuclear export of Pho4 by the receptor Msn5 (ref. 2). Before nuclear export, Pho4 can also be phosphorylated at Ser152 (S4), which in turn prevents reimport of Pho4 into the nucleus. There is a preference for phosphorylation: S6 (transcription) > S4 (import) > S2/S3 (export region) > S1 (ref. 24). Moreover, phosphorylation of Pho4 only at the S6 site can lead to transcription of a subset of phosphate-responsive genes^25. It has also been suggested that differential gene expression could be regulated by phosphorylation at the export region (S2 and S3 site) but no mechanism is known and it is not clear whether differential phosphorylation of S2 and S3 could lead to distinct responses in Pho4 activity.

The selective introduction of 1 at specific sites in the export region should allow one to dissect the effects of specific phosphorylation of each of these sites on the dynamics of protein trafficking between the nucleus and cytoplasm. To this end, we first substituted a nonsense amber mutation for the S3 codon (S3→TAG) in a wild-type (WT) Pho4-GFP (green fluorescent protein) fusion construct and co-expressed the mutated gene with the mutant suppressor tRNA^Leu^CUA/LeuRS pair in S. cerevisiae in presence of 1 to generate the S3→1 Pho4-GFP mutant. Because the phosphorylation site S3 is blocked, the S3→1 Pho4-GFP mutant accumulates in the nucleus even at high Pi concentrations, which is consistent with the published effect of the S3→Ala mutation on localization of Pho4-GFP fusion constructs (Fig. 2)^23. This demonstrates that the S3→1 Pho4-GFP is still recognized by the import receptor Pse1 (ref. 26) and localized to the nucleus when the S3 phosphorylation site is blocked (additionally, the photocaged Pho4-GFP can also be transcriptionally active—see Supplementary Fig. 4 online). Cells grown in high Pi, in the absence of 1 showed no significant fluorescence (other than autofluorescence, Fig. 2c), which indicates that there is little background translational read-through to the C-terminal GFP.
Figure 3  Real-time analysis of in vivo photolysis of DMNB-Ser Pho4-GFP mutants. (a) The normalized average fluorescence intensity is plotted as a function of time for S3 → 1 Pho4-GFP (gray; photolyzed, filled squares, number of nuclei n = 71; nonphotolyzed, open circles, n = 67) and S2 → 1 Pho4-GFP (black; photolyzed, filled squares, n = 101; nonphotolyzed, open circles, n = 101). The solid lines depict monoeponential fits with time constants of $t_{S3} = 28.4$ ± 2.4 s (gray) and $t_{S2} = 47.0$ ± 5.1 s (black).

(b,c) Nuclear fluorescence is still visible after photolysis of the S2 → 1 S3 → Ala Pho4-GFP (b) and S3 → 1 S2 → Ala Pho4-GFP (c) mutants. Analysis for the S3 → 1 S2 → Ala Pho4-GFP mutant cells in high P$_i$ (red; photolyzed, filled squares connected by red line, n = 109) and no P$_i$ (orange; photolyzed, filled squares n = 156), and for the S2 → 1 S3 → Ala Pho4-GFP mutants in high P$_i$ (purple; photolyzed, filled squares n = 80; nonphotolyzed, open circles n = 60, for better clarity only this baseline is shown) and no P$_i$ (light blue; photolyzed, filled squares n = 123). Error bars show s.e.m.; **P < 0.01.

Laser photolysis of the photocaged S3 → 1 Pho4-GFP mutant should result in photodeprotection of 1 to afford WT Pho4-GFP, which can then be phosphorylated by the cyclin-CDK complex and exported from the nucleus into the cytoplasm. To test this notion, we photolyzed cells expressing S3 → 1 Pho4-GFP in the presence of high P$_i$ by scanning only the upper part of the microscope field with a visible blue light laser ($\lambda = 405$ nm, Fig. 2d). The lower part of each field was not irradiated and therefore served as an internal control in all experiments. We acquired a time series before and after photolysis using excitation at $\lambda = 488$ nm, which is beyond the absorption wavelength of the caging group (Supplementary Fig. 1). Fluorescence from nuclei in the photolyzed area of the image disappeared over time (Fig. 2d). In addition, Figure 2c shows two exemplary cells before and after photolysis and a corresponding line scan that shows the decrease in nuclear fluorescence and a small increase in fluorescence in the larger cytoplasmic volume after photolysis. These results demonstrate that the function of WT Pho4-GFP can be restored by removal of the photolabile protecting group with a short visible light trigger, thus resulting in export from the nucleus by Msn5 in the presence of extracellular P$_i$.

To study the kinetics of Pho4 translocation, we analyzed the acquired image series of the S3 → 1 Pho4-GFP mutant as a function of time before and after photolysis (Fig. 3a). Because of the low contrast in the cytoplasmic region, only fluorescence changes in the nucleus were measured in every image and analyzed over time. We used a fully automated procedure for data analysis that is based on a wavelet transformation to detect fluorescent nuclei (Supplementary Fig. 5 online). The photolyzed nuclei showed a fluorescence decrease that followed a time constant of $t_{\text{S3 TAG}} = 28.4$ ± 2.4 s (errors for time constants depict jackknife estimates, see Supplementary Fig. 5 for details on data processing and data fitting). This time constant is consistent with previous experiments that found the translocation process to be complete within 3–6 min after addition of phosphate to P$_i$-starved cells.

To identify possible differences between phosphorylation at the S2 and S3 sites, we generated three additional mutants, including single-point DMNB-Ser mutants and mutants in which the other phosphorylation site was converted to alanine. We repeated the uncaging experiments with a S2 → 1 Pho4-GFP mutant (Fig. 2f) shows a fluorescence loss similar to that shown in Fig. 2d) and measured slower export kinetics ($t_{\text{S3 TAG}} = 47.0$ ± 5.1 s, Fig. 3a), which indicates that the S2 → 1 and S3 → 1 Pho4-GFP mutants have differential phosphorylation or export kinetics.

We then analyzed the S2 → 1 S3 → Ala Pho4-GFP and S3 → 1 S2 → Ala Pho4-GFP double mutants (Fig. 3b,c). Unlike the single mutants studied above, these mutants can only be phosphorylated at one site of the export region. In agreement with the previously reported effect of S3 → Ala Pho4 mutants, we found that nuclear export remained blocked after photolysis ($t_{\text{S3 TAG S3 Ala}} > 100$ s). In contrast, there was still a slow kinetic decay for the S3 → 1 S2 → Ala Pho4-GFP mutant ($t_{\text{S3 TAG S3 Ala}} > 0$, Fig. 3d). This decay is absent if cells are additionally starved for P$_i$, which indicates that it arises from residual export activity (Fig. 3d). Previous measurements of S3 → Ala Pho4-GFP localization under steady state conditions did not reveal this activity, possibly because of competing import activity. Taken together, our data suggest that differential phosphorylation at sites S2 and S3 leads to distinct export kinetics for these isoforms of Pho4. It is possible that this differential activity is part of the mechanism regulating differential gene expression, by controlling the kinetic localization of Pho4 in the nucleus.

In summary, we described a method to nondestructively control protein function in vivo with high spatial and temporal resolution. We used this methodology to photoinitiate the phosphorylation of Pho4 by the cyclin-CDK complex and to monitor the kinetics of protein trafficking in real time. This general method should be applicable to other proteins in which serine is involved in a specific binding or catalytic activity, and it is currently being extended to additional residues and to mammalian cells.

**METHODS**

Evolution of tRNA synthetases for incorporation of 1. To evolve a LeuRS specific for 1, a previously published protocol was used. Briefly, positive selection was carried out by suppression of two amber (TAG) codons in the gene encoding the transcriptional activator GAL4. Production of functional full-length GAL4 drives expression of a genomic URA3 reporter gene and complements uracil auxotrophy, thus allowing clones harboring active synthetase mutants to be selected on synthetic dropout medium lacking uracil (SD
medium containing dextrose) supplemented with 1 mM l (stock solutions of l are 0.1 M in 0.1 M HCl). Negative selection of synthetases that accept endogenous amino acids was carried out by growth on medium lacking l but containing 0.1% 5-fluoroorotic acid, which is converted into a toxic product by URA3 protein. Based on its growth rate in the presence and absence of l, clone BHS emerged after five alternating rounds of positive and negative selection. The T2S2A mutation was introduced into the plasmid pescTrpLeuRSBH5 isolated from the selected clone BHS as described previously18 to afford plasmid pescTrpLeuRSBH5T2S2A.

Expression, purification and characterization of hSOD-His, harboring 1.

Protein expression was performed with strain SCY4 (MATa, ade2-101 ura3-1 leu2-3, 112 trpl his-31 ACT2a:kan [circl]) transformed with the hSOD expression plasmid pCISOD-3TAG-His, ref. (18) and the trRNALeu/LeuRS-encoding plasmid (pescTrpLeuRS derivatives). Cultures were grown to saturation at 30 °C in SD medium lacking leucine and tryptophan and diluted 50-fold into SD medium (containing dextrose) lacking leucine and tryptophan and containing 4 mM l. This culture was incubated for 16 h at 30 °C and cells were harvested. Lysis and purification was performed as described previously, except that washing buffer contained 50 mM instead of 25 mM imidazole18. SDS-PAGE analysis was performed using GelCode Blue Stain (Pierce). Yields were routinely quantified using a BCA assay (Pierce).

Cloning and expression of DMNB-Ser Phot4-GFP mutants. Phot4-GFP (ref. 1) was a gift (see Acknowledgments). Mutants were generated using the Quick-Change method to introduce either TAG or alanine mutations at Ser114 (S2) and/or Ser128 (S3). The respective Phot4-GFP mutant genes were inserted into the pesc-His plasmid (Stratagene) using NotI andSac sites and expressed under control of the Gal1 inducible promoter. The plasmid was cotransformed into an INVSC1 strain (Invitrogen) harboring plasmid pescTrpLeuRSBH5T2S2A. The strain was grown in SD medium lacking galactose and l until mid-log phase, pelleted and resuspended in medium containing 2% galactose and 2% raffinose at a similar optical density; 4 mM l was then added to this medium. Cells were allowed to grow in the dark for an additional 7–8 h, washed and resuspended in yeast nitrogen base (Formmouds plus 2% glucose) and 20 mM KH2PO4 (for high P, measurement). For measurements in the absence of P, cells were allowed to grow for 7–8 h, washed and resuspended in two volumes of yeast nitrogen base plus 2% glucose plus 20 mM KCl. Cells were grown for an additional 2–3 h before the imaging experiment.

Real-time imaging and analysis. Experiments were performed on an inverted Bio-Rad (Zeiss) Radiance 2100 Rainbow laser scanning confocal microscope with a 1.4 numerical aperture oil objective and an argon laser (l = 488 nm). For photolysis a visible blue light laser (l = 405 nm) was used. The scan speed was adjusted to 166 lines per second, resulting in a pixel dwell time of about 10.4 μs. The image size was 512 by 512 pixels (82 μm by 82 μm), with a pixel size of 0.16 μm by 0.16 μm. Thus, for a typical nucleus with a size of 10 by 10 pixels, the total laser light exposure time was approximately 1 ms. The image depth was 8 bit, and displayed images were filtered with a 3 by 3 median filter. The laser power measured at the nose piece of the objective was approximately 65 μW at λ = 488 nm and 410 μW at λ = 405 nm in all experiments. For quantitative analysis of the real-time data, the automated procedure described in Supplementary Figure 5 was used.

Statistical analysis. Statistical analysis in Figure 3d was done using a one-way ANOVA to perform a Dunnett’s test between photolyzed cells harboring S3TAG→S2→Ala Phot4-GFP grown in the presence of high P and the other data shown in this panel for each time point. Significance levels were assigned to P values according to *P < 0.05 and **P < 0.01. All analyses were performed using Igor Pro (WaveMetrics). Error bars are given as s.e.m.

Note: Supplementary information and chemical compound information is available on the Nature Chemical Biology website.