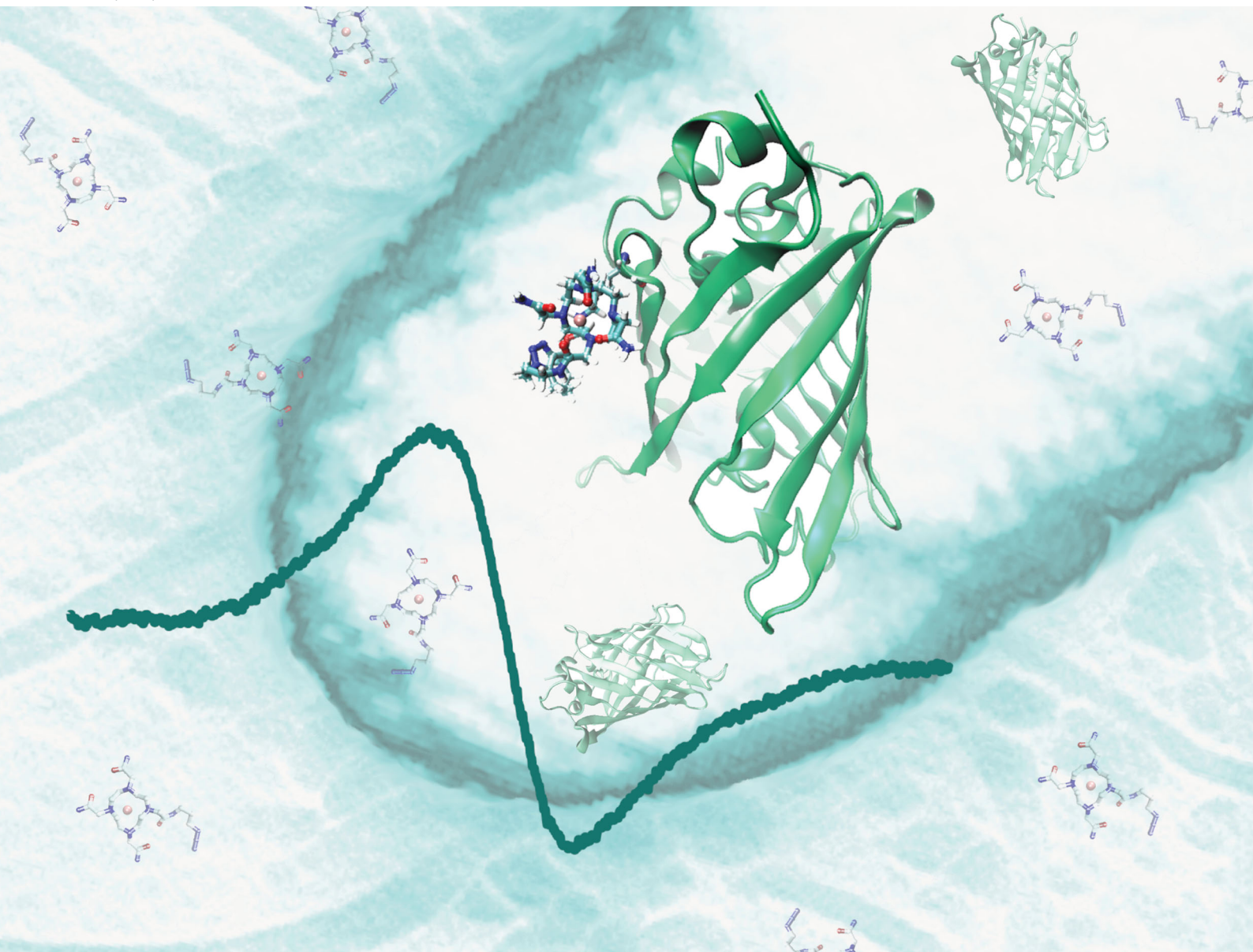


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COMMUNICATION

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In cell Gd³⁺-based site-directed spin labeling and EPR
spectroscopy of eGFP



In cell Gd³⁺-based site-directed spin labeling and EPR spectroscopy of eGFP†

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Label-based functional studies of biomolecules in their native environment require labeling reactions inside living cells. *In cell* spin labeling using alkyne–azide click chemistry with a Gd³⁺-DOTAM-azide complex is shown to provide high spin label stability and narrow EPR lines for EPR spectroscopic detection of a spin labeled protein in living cells at ambient temperatures.

Information about the structure and dynamics of biomolecules is a prerequisite to understand their physiological function. The vast majority of structural data present to date has been obtained under highly artificial conditions, *i.e.* in crystals or in highly concentrated solutions and purified from “contaminants”. However, the complex cellular environment can have strong influence on the biomolecules’ behavior,¹ which raises the requirement to develop techniques that allow obtaining structural data *in cell*. Techniques such as X-ray crystallography and cryoEM, which provide atomic resolution, meet their limitations in such complex environments and *in cell* NMR is currently growing, but still a very challenging tool.^{2,3} EPR spectroscopy together with site-directed spin-labeling (SDSL) is an alternative approach, which can give site-specific information on structure and dynamics of complex biomolecular systems independent of their size and homogeneity of the environment.⁴ (Time-resolved) continuous wave (cw) EPR measurements of the spin label dynamics can be performed at physiological temperatures, yielding insights into local structure and dynamics.^{5,6} Moreover, double-electron–electron resonance (DEER) spectroscopy has become a powerful tool for measurements of inter- or intramolecular spin distance distributions within a range of 1.5–10 nm, providing structural data with near-atomic resolution.⁷ Paramagnetic labeling has also

found a broad application in NMR spectroscopy, offering a significant sensitivity enhancement.⁸

First attempts to study paramagnetically labeled biological systems in a cellular context have successfully been made by incorporation of spin labeled proteins into living cells,⁹ using either micro-injection into oocytes or by membrane penetration (*via* electroporation, hypotonic swelling or incubation).^{10–15} However, most of these applications are based on *in vitro* labeling, usually with cysteine specific labels followed by incorporation of the protein mostly into a different cell line and in higher than physiological concentrations. Alternatively, membrane proteins were labeled on the outer cell surface or recombinant proteins were directly expressed with spin-labeled unnatural amino acids (UAA).^{16,17}

In order to label a specific protein at desired positions inside the cell with a spin label of interest, an orthogonal labeling approach is required due to the broad range of chemical functionalities present inside cells. Orthogonal labeling with the genetically encoded *p*-acetyl-L-phenylalanine was applied using a hydroxylamine reagent to generate a nitroxide side chain *in vitro*.¹⁸ However, reaction conditions do not allow *in cell* applications. Click chemistry – here an alkyne–azide cycloaddition (AAC) – is a highly selective labeling approach, which has been applied for labeling of proteins in, *e.g.*, fluorescence studies.^{19,20} Both types of click reactions, the strain-promoted (SPAAC)²¹ and the copper-catalyzed (CuAAC)^{22,23} variant, were shown to be successfully applicable for SDSL with nitroxides *in vitro*.^{24,25} Recently, we demonstrated that the green fluorescent protein eGFP can be spin labeled with nitroxides inside living *E. coli* cells *via* click reaction with genetically encoded UAAs.²⁶ However, *in cell* EPR detection is limited by the fast reduction of nitroxides.²⁷

Gd³⁺ complexes, long known as MRI contrast agents, were introduced for SDSL of proteins and show an outstanding stability in a reducing environment.^{13,14} Successful ligation of a Gd³⁺ tag to *p*-azido-L-phenylalanine introduced into proteins using CuAAC has been shown, but so far only *in vitro*.^{9,28} Moreover, Gd³⁺ labels have been applied for distance measurements using pulse EPR at high magnetic fields.^{29,30}

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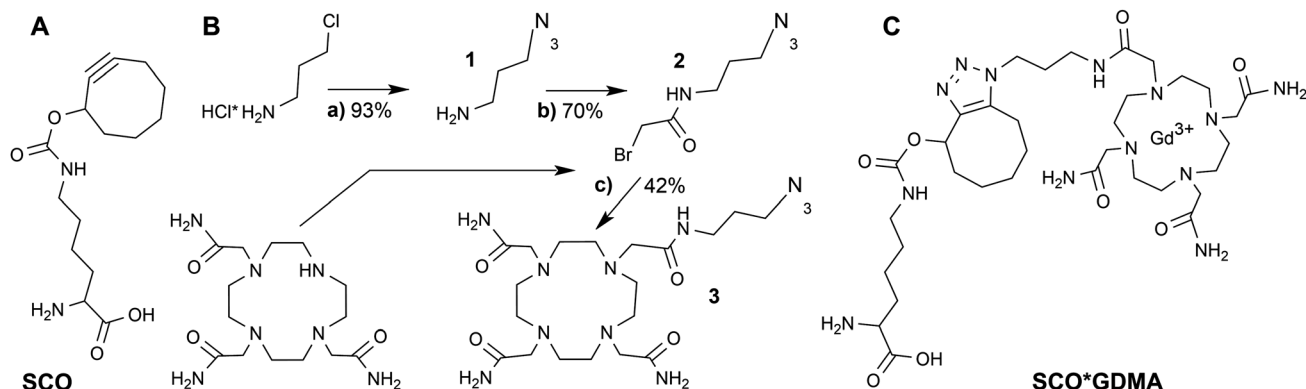
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Scheme 1 (A) Structure of strained-cyclic octanyl (SCO). (B) Synthesis of DOTAM-azide. Reaction conditions: (a) NaN₃, H₂O, 70 °C, 3 days; (b) ClCOCH₂Br, Et₃N, CH₂Cl₂, 0 °C → RT, 15 h; (c) K₂CO₃, MeOH, 55 °C, 24 h. (C) The side chain of SCO after binding with Gd³⁺-DOTAM-azide (GDMA).

In this communication we show the applicability of *in vitro* and *in cell* click labeling using a newly synthesized Gd³⁺-based DOTAM-azide spin label and its direct *in cell* detection with continuous wave EPR at conventional X-band frequency (9–10 GHz) with high sensitivity at room temperature (RT). We use eGFP with an amber mutation at position 39 with strained-cyclic octanyl (SCO, Scheme 1A) because it has successfully been applied for the labeling with azido-functionalized nitroxides and FRET labels and allows precise concentration determination.^{26,31} The incorporation of UAAs containing azide groups for the labeling with alkyne-functionalized markers is possible as well, however, this is less efficient for *in cell* labeling due to azide reduction to amines.^{32–34} However, a lower degree of degradation was reported for mammalian cells than for bacteria making them promising for further applications.³⁵

Gd³⁺ ions are high spin systems ($S = 7/2$) that exhibit large zero-field splitting (ZFS) leading to broad linewidths at conventional X-band frequencies (~ 9 – 10 GHz), which become sharper with increasing frequency. However, the ZFS strongly depends on the chelator structure and symmetry.³⁶ So far one of the smallest ZFS for Gd³⁺ complexes from commercially available chelators was reported for Gd³⁺-DOTAM (GDM) (~ 560 MHz) exhibiting a narrow spectrum with a linewidth Γ of ~ 2 mT at X-band (Fig. 1A) compared to ~ 10 mT of the commonly used Gd³⁺-DOTA complex.^{36–38} This effect was explained by the only structural differences between the DOTA and DOTAM tags – substitution of the Gd-coordinating carboxyl groups by the less

electronegative acetamides.³⁷ The linewidth of protein bound Gd³⁺-DOTA derivatives has been previously shown to be correlated with dynamics at X-band in a similar way to nitroxides.³⁹ However, its cw EPR detection is limited to the sub-millimolar concentration range due to the large ZFS. In contrast, the narrow linewidth of the GDM complex allows for cw EPR detection at X-band in the low μ M concentration range providing a sensitivity comparable to nitroxides, thereby making this complex promising for *in vitro* and *in cell* EPR spectroscopic studies. Due to the lack of DOTAM functionalized as a labeling tag, we synthesized a DOTAM-azide derivative as described in the following, suitable for subsequent labeling of alkyne groups.

DOTAM-azide was synthesized in three steps from commercially available educts according to Scheme 1B. First, 3-chloropropylamine salt was converted into 3-azidopropylamine (**1**) in 93% yield by the exchange reaction with sodium azide as described.⁴⁰ Next, the azide **1** was coupled to bromoacetyl chloride in the presence of Et₃N to give *N*-(3-azidopropyl)-2-bromoacetamide (**2**) in 70% yield. Final condensation of the bromide **2** with 1,4,7,10-tetraazacyclododecane-1,4,7-triacetamide in the presence of K₂CO₃ at elevated temperature resulted in the formation of DOTAM-azide **3** that was isolated in 42% yield. Its structure was confirmed by ¹H and ¹³C NMR spectroscopy and electrospray-ionization mass spectrometry (ESI MS, see Fig. S1, ESI[†]). A detailed description of the synthesis together with spectral data is provided in the S11 (ESI[†]).

The room temperature cw EPR spectrum of DOTAM-azide (**3** in Scheme 1B) after complexation with Gd³⁺ (GDMA) exhibits a peak-to-peak linewidth of 2.8 mT, which is only slightly broader than for the symmetric Gd³⁺-DOTAM (Fig. 1). EPR spectra of both, GDM and GDMA, recorded at different viscosities (see S14, Fig. S3, ESI[†]) show that a decreasing motional averaging of the ZFS anisotropy is reflected in a considerable increase of the linewidth (Fig. 1B) proving its sensitivity to reorientational dynamics.

The EPR spectrum of the spin labeled eGFP-Y39SCO*GDMA (Fig. 1A) reveals a pronounced increase of the linewidth to 4.4 mT, as obtained from single Lorentzian fits, compared to the spectrum of the unbound label (2.8 mT) due to the spin label immobilization. This allows discrimination between

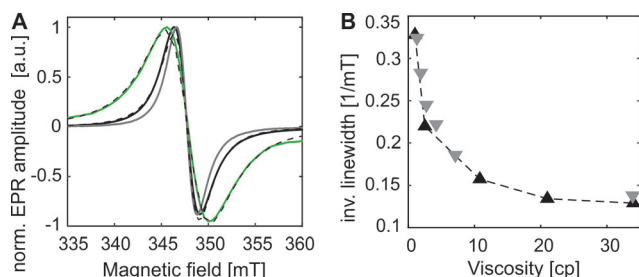


Fig. 1 (A) X-band EPR spectra ($T = 293$ K) of GDM (grey), GDMA free in solution (black) and bound to eGFP-Y39SCO (green) with Lorentzian fits (dotted lines). (B) Viscosity dependence of the inverse linewidth for GDM (grey) and GDMA (black).



bound and unbound spin labels and makes this spin label a promising candidate to probe reorientational motion (Fig. 1A, B and Fig. S3, ESI[†]) at ambient temperatures.

Surprisingly, while SPAAC of eGFP-Y39SCO with GDMA resulted in a high labeling efficiency of $\sim 85\%$ (4 h at 37 °C, see ESI[†]), comparable with the efficiency previously achieved for nitroxides,²⁶ the same did not hold true for CuAAC. CuAAC of eGFP-Y39PrK (PrK is abbreviation for propargyl-L-lysine) with GDMA did not result in fast and sufficient labeling as it has been reported for nitroxides and FRET labels (see SI3, Fig. S2, ESI[†]).^{26,41} Such a retardation of the usually faster click reaction variant might be caused by the interference of the positively charged spin label with the copper catalyst under the present experimental conditions. This makes CuAAC not well suited for *in cell* applications with Gd³⁺ labels and shows SPAAC to be the more efficient strategy even for a strained UAA with a low reaction rate.²⁰

Since distance measurements are of general interest in EPR-aided structural biology, the applicability of the new side chain Y39SCO*GDMA for DEER is addressed in the following. For this purpose, eGFP-Y39SCO*GDMA was additionally labeled *in vitro* with the *S*-(1-oxyl-2,2,5,5-tetramethyl-2,5-dihydro-1H-pyrrol-3-yl)methyl methanesulfonothionate spin label (MTSSL) at the outer cysteine position 48 and at the inner, less accessible, cysteine position 70, yielding side chains C48R1 and C70R1 (see Fig. 2A, for labeling details see SI3, ESI[†]) in a similar way as has been shown before.²⁶

The distance distribution obtained from a Q-band DEER trace by a “user-unbiased” Neural network approach in DeerAnalysis2019^{42,43} (Fig. 2B) shows agreement with the distribution obtained by Tikhonov regularization (Fig. S6 and S8, ESI[†]), both revealing a broad distance distribution with two main peaks at 3.5 and 4.3 nm and a smaller contribution at 2.3 nm. To compare the experimental results to the protein structure, we prepared a rotamer library (RL) for the SCO*GDMA side chain. 8192 rotamers were generated by hierarchical clustering of a 250k-membered Monte Carlo ensemble using a forgive factor of 0.8. The library is implemented in MMM⁴⁴ versions 2019.1 and later as the spin label with code GM1 (see SI5 for details, ESI[†]). The distance distribution between C48R1 and Y39SCO*GDMA using the default MTSSL

library of MMM2019.1 and the newly calculated one, respectively, fits well to the middle part of experimental distribution and the main peak at 3.5 nm (Fig. 2B, blue). Only one populated rotamer was obtained for both C48R1 and C70R1 (see Fig. 2A), which is in agreement with the observed low modulation depth in DEER as well as with the low experimental labeling efficiency for C70R1 (see SI3, ESI[†]). Thus, the large distribution width is mainly caused by the conformational flexibility of the long SCO*GDMA side chain. The presence of larger distances found in the experimental distribution can be attributed to the possible dynamics of the outer loop where C48R1 is located. To verify this assumption, MD simulations on eGFP were performed revealing additional possible conformations (see SI5 for details, ESI[†]). This led to additional rotamers for C48R1 resulting in the distance distribution shown in green in Fig. 2B, which covers the experimentally observed long distances. The small contribution at 2.3 nm in the experimental distribution likely arises from a less probable dipolar interaction between C70R1 and Y39SCO*GDMA (see Fig. S6, ESI[†]).²⁶

For application of the SPAAC for labeling of eGFP inside *E. coli* cells we tested the biocompatibility of required reagents. Exposing *E. coli* cells to the Gd³⁺-DOTAM complexes revealed no detectable toxic effects at a wide range of concentrations (see SI7, ESI[†]). Furthermore, Gd³⁺ complexes were able to penetrate the cell membrane of *E. coli* cells during cell growth at 37 °C in a concentration and incubation time dependent manner without the need of applying external stress or additional procedures (see SI7, Fig. S10, ESI[†]). External millimolar spin label concentrations resulted in micromolar *in cell* concentrations. Moreover, no reduction in *E. coli* cell lysate was found for Gd³⁺-DOTAM complex after 10 h of incubation (see SI7, Fig. S11, ESI[†]), which is in line with the previous reports for other chelators.^{13–15}

To test the label specificity, *E. coli* BL21 cells were incubated with GDMA spin label. Afterwards the medium was exchanged to remove extracellular labels and the cells were collected (see SI8 for details, ESI[†]). As shown in Fig. 3A the cells without eGFP overexpression exhibited a broadened signal originating from penetrated GDMA as expected due to the higher viscosity in cells (see Fig. 1B). The lysate of these control cells did not show any EPR signal after washing and concentrating using 10 kDa cutoff concentrators (Fig. 3B), which proves that the EPR signal of the control present in Fig. 3A originates from unbound GDMA only.

For *in cell* labeling, *E. coli* BL21 cells expressing eGFP-Y39SCO were incubated with GDMA for 2 h at 37 °C (see SI8 for details, ESI[†]). The EPR spectrum of the cells shows a line broadening which is increased compared to the control (Fig. 3A). This is the evidence that labeling of eGFP-Y39SCO occurred inside the *E. coli* cells because bound GDMA exhibits a broader line (Fig. 1A) than free GDMA in both, buffer and cells. To further prove *in cell* spin labeling, the cells were lysed and the lysate washed under conditions that restrain further binding reactions (see SI8, ESI[†]). Only for cells that had expressed eGFP, the resulting EPR spectrum was similar to that observed for eGFP-Y39SCO*GDMA obtained *in vitro* (Fig. 3B and 1A). The spin concentration of the bound spin label inside cells was calculated to be $\sim 9 \mu\text{M}$, which is about 50% of the total GDMA concentration

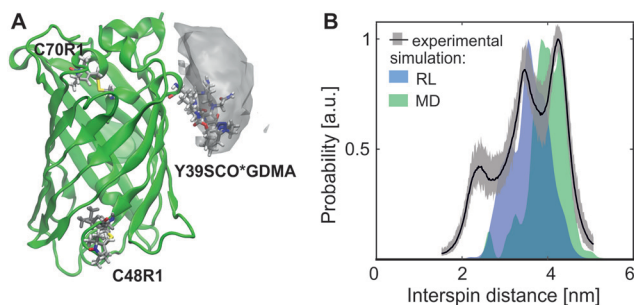


Fig. 2 eGFP-Y39SCO*GDMA-C48R1-C70R1 (A) Structure (PDB: 4EUL) with calculated rotamers of GDMA, position 39 (99.5% probability envelope in gray), and of R1, positions 48 and 70. (B) Q-band DEER data: experimental (black, grey) and calculated (RL, blue and MD, green) distance distributions for Y39SCO*GDMA-C48R1 sites.



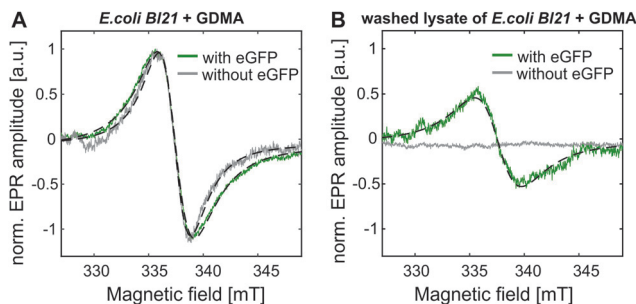


Fig. 3 (A) Amplitude-normalized EPR spectra of *E. coli* BL21 cells after incubation with GDMA at 37 °C without (grey) and with (green) over-expressed eGFP-Y39SCO. (B) The lysates of the samples shown in A were washed to remove free GDMA. Lorentzian fits are shown in dotted lines.

(~18 μM) determined for intact cells before purification. This means that half of the GDMA present in the cells reacted with eGFP-Y39SCO. This is in reasonable agreement with the result of a fitting of two Lorentzians with fixed linewidths for free and bound GDMA to the EPR spectrum of *E. coli* cells with eGFP-Y39SCO incubated with GDMA, yielding an amplitude ratio for free : bound of 1.9 : 1.

Prolongation of the incubation time from 2 to 6 hours increased the labeling efficiency from 5 to ~22% limited by the GDMA to protein ratio of 1 : 5 inside cells. For native protein concentrations below 20 μM the GDMA concentration present in the cell is not a limiting factor, and the achievable labeling efficiencies are expected to be higher. The observed reaction yields agree with previous reports for SCO as a strained UAA with a low reaction rate in SPAAC, whereas future utilization of other variants can accelerate the reaction rate by several orders of magnitudes.²⁰

In summary, the SPAAC with a Gd^{3+} -DOTAM-azide spin label is shown to proceed in living *E. coli* cells, thus demonstrating the feasibility of this reaction for *in cell* labeling of UAA-modified proteins using Gd^{3+} -based spin labels. The newly synthesized Gd^{3+} spin label allowed *in cell* EPR detection, overcoming previous limits for *in cell* cw EPR due to reduction of nitroxides by the high chemical stability of the new Gd^{3+} spin label and its small line width. The observed increase in the (ambient temperature) cw EPR line width upon binding of the label to the protein opens the possibility for studying protein dynamics *in cell*. The method is shown to be applicable for (low temperature) inter spin distance measurements *in vitro*. Utilization of artificial amino acids functionalized with strained alkenes providing higher click reaction rates and short chain lengths will further enhance this applicability.

Conflicts of interest

There are no conflicts to declare.

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