We review new and established methods for the chemical modification of proteins in living cells and highlight recent applications. The review focuses on tag-mediated protein labeling methods, such as the tetracysteine tag and SNAP-tag, and new developments in this field such as intracellular labeling with lipoic acid ligase. Recent promising advances in the incorporation of unnatural amino acids into proteins are also briefly discussed. We describe new tools using tag-mediated labeling methods including the super-resolution microscopy of tagged proteins, the study of the interactions of proteins and protein domains, the subcellular targeting of synthetic ion sensors, and the generation of new semisynthetic metabolite sensors. We conclude with a view on necessary future developments, with one example being the selective labeling of non-tagged, native proteins in complex protein mixtures.

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Introduction
Chemists are becoming increasingly fascinated with derivatizing proteins by genetically non-encodable synthetic molecules. As discussed in this review, such molecules include fluorescent dyes, chemical crosslinkers, pharmacologically active compounds, and synthetic fluorescent probes for ions such as Ca\(^{2+}\). The labeling of proteins with synthetic molecules provides exciting new tools for studying proteins and their function in the cell, and is promising to have a strong impact on drug development. In this review, we will provide a concise overview over established and new methods that provide proteins derivatized with a label, and give illustrative recent examples of their application. For further information, the reader is directed to recently published more exhaustive reviews [1–4]. The focus of our review lies on covalent tag-based labeling methods, as these allow an efficient and irreversible transfer of labels in mammalian cells. However, we will also comment on unnatural amino acid incorporation as a tool of increasing importance for mammalian cell studies. The discussion of methods and applications is preceded by general considerations on tag-mediated labeling. We will conclude with an outlook on future directions and highlight areas that would profit most from new developments.

Tag-mediated labeling
An ideal method for tag-based protein labeling should exhibit the following features: (i) the possibility to introduce any label of choice in one step, (ii) fast and quantitative labeling, (iii) no labeling of non-target proteins, (iv) a small tag to minimize its impact on protein function, (v) the formation of a stable, covalent bond between protein and label, and (vi) no side effects of the reagents used for labeling. Finally, an ideal tag should work in vitro, in complex protein samples, on the cell surface, within the cell and cellular compartments, and in living animals (in vivo), with this order representing an increasing level of difficulty. None of the existing labeling methods fulfills all these requirements. It is notable, however, that the existing methods of tag-mediated labeling can be grouped into three families (Figure 1) that have inherent strengths and weaknesses. Self-labeling tags and self-labeling proteins – such as the tetracysteine-tag and the SNAP-tag, respectively – are able to directly react with the labeling compound. Provided that the labeling molecule is cell membrane permeable, they can therefore be applied to labeling proteins in the cell interior. Compared to self-labeling tags, self-labeling proteins usually provide a higher labeling specificity, but the higher specificity inevitably comes at the price of a larger tag size. Enzyme-mediated labeling of tags (such as Biotin ligase mediated labeling) additionally requires an enzyme to covalently link the labeling compound to the tag. In many cases, this approach combines the small tag size provided by self-labeling tags and the specificity provided by enzymes. However, enzyme-mediated labeling of tags is restricted to labeling cell surface proteins unless the enzyme can be expressed within the cell and intracellular metabolites do not interfere with labeling. Some enzyme-mediated labeling methods are limited with regard to a free choice of the label. This applies to the cases where the label itself plays a role in substrate recognition, as discussed below. By contrast, the label usually has a minor impact on the labeling rate of self-labeling tags and self-labeling proteins, enabling the transfer of – in principle – arbitrary labels.

How important is the size of the tag? The plethora of reported successful fusions of autofluorescent proteins to
various proteins demonstrates that a large tag is usually not problematic. However, while the impact of a tag on the function of the tagged protein is – unfortunately – rarely rigorously checked, in some cases small tags have been reported to perform significantly better than auto-fluorescent protein fusions. As may be expected, this involves proteins that are part of a tightly packed structure or involved in a translocation process. For example, an impact of tag size on protein function has been shown for virion surface proteins [5], beta-tubulin [6], beta-actin with regard to transport into the cell nucleus [31,32], and effector proteins that are translocated by gram-negative bacteria into eukaryotic cells [7]. Further, large tags are usually restricted to N-terminal or C-terminal fusions, while small tags can in principle also be incorporated at internal sites of the protein of interest.

In the following, we introduce established methods and new developments for each of the tag-mediated labeling families. An overview over the self-labeling tags and proteins discussed in this review is shown in Table 1, providing data on the labeling rate and efficiency, and indicating whether the method works for intracellular labeling. An overview over methods used for enzyme-mediated labeling of tags is given in Table 2; as a rough guide to the labeling rate attainable with such methods, we provide typical published labeling protocols for the labeling of tagged cell surface proteins.

### Self-labeling tags

The first tag developed for specific protein labeling is the tetracysteine tag [8], which exploits the large affinity of peptide sequences containing four cysteines – usually...
CCPGCC – towards synthetic molecules containing two arsenic atoms, also called biarsenicals. The detection of labeled proteins is simplified by the fact that the fluorescence of biarsenical dyes such as FlAsH and ReAsH is switched on by binding to the tag. FlAsH and ReAsH are cell permeable, making tetracysteine labeling the best established method for the fluorescent labeling of small tags within cells. The specificity of biarsenical labels for their tag is not perfect; labeling and washing procedures therefore have to be carried out in presence of a competing thiol reagent to reduce background labeling. This labeling procedure – which is also slightly cytotoxic [31**] – may be problematic for tetracysteine tag applicability in vivo.

Recently, a similar self-labeling tag was designed with the aim of eliminating thioly reagents. In this method, dyes derivatized with boronic acids are designed to react with a tetraserine tag [9]. In a proof-of-principle experiment, the high-affinity binding of the label to the tag was demonstrated, but more development is needed to selectivity label tagged proteins within the cell. A series of self-labeling tags was also developed by Hamachi and co-workers [10,11]. They are based on bifunctional molecules containing a recognition element for the protein of interest and a reactive group. The initially noncovalent interaction between the protein of interest and the labeling compound is turned into a covalent one owing to reaction of the reactive group with a proximal amino acid side chain in the protein, as illustrated in Figure 1. The reaction leads to a release of the recognition element. The approach works selectively – but rather slowly – in complex mixtures using the interaction between positively charged Ni2+–containing reagents and the 6-His tag or 10-His tag [10] or the negatively charged FLAG–tag [11]. A new development in self-labeling tags is based on the reaction of N-terminal cysteines with N-cyano-benzothiazole derivatives containing a label [12**]. The reaction is selective owing to the requirement of an amino group in β-position to the cysteine side chain, which is provided by the free N-terminal amine. N-terminal cysteines do not occur naturally in the cell, but can be obtained by enzymatic cleavage of an introduced recognition site for Tobacco Etch Virus (TEV) Protease (ENLYFQ+C, cleavage site marked with an arrow) or from proteins fused to an intein. The authors demonstrate specific, but rather slow labeling in cell lysates and on the surface of living cells; an intracellular labeling has not yet been shown.

**Self-labeling proteins**

Self-labeling proteins react covalently with a substrate that is linked to the label of interest and are mostly derived from enzymes. The most widely used self-labeling proteins are the HaloTag [13] and the SNAP-tag [14], the latter of which was developed in our laboratory. The advantages of these self-labeling proteins are their high speed and specificity (cf. Table 1), the large array of commercially available substrates, and the simple synthetic accessibility of arbitrary custom labels. Owing to the fact that labeling by HaloTag and SNAP-tag is irreversible and quantitative, these tags are well suited for the detection and quantitation of tagged proteins via in-gel fluorescence scanning of SDS-PAGE gels. HaloTag and SNAP-tag work well within cells and also within subcellular compartments when cell-permeable substrates are used. For example, SNAP-tag labeling has been described not only for proteins in the cytosol, but also in the nucleus, endoplasmic reticulum, mitochondria, and golgi apparatus (see, e.g. refs. [63,67]). Even non-permeable substrates that are microinjected do not lead to background staining, as excess dye usually does not remain within the cell [15].

Self-labeling proteins have recently been shown to be employable also for in vivo studies [43*], as discussed in the applications section below. Another recent development in this field is a further self-labeling protein, CLIP-tag [16*]. CLIP-tag allows a free choice of the label and owing to its orthogonality can be used in conjunction with SNAP-tag and HaloTag. An alternative method uses a mutant beta-lactamase that reacts covalently with a beta-lactam derivatized with a quencher and a fluorescent dye [17]. The method, termed BL-tag, is however slower than already existing methods (Table 1) and restricted to the cell surface. Recently, a covalent version of the noncovalent trimethoprim-tag [18] (TMP-tag) has been developed that is based on proximity labeling. It was shown to be applicable to intracellular labeling of proteins tagged with *E. coli* dihydrofolate reductase [19].

**Enzyme-mediated labeling of tags**

In this family of methods, an enzyme is used to attach a label to a usually small recognition sequence in the

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**Table 2**

Overview of enzymes used in enzyme-mediated labeling. Typical labeling protocols were taken from the literature [30,77–79]. *Lipoic acid ligase needs to be engineered to accept different labels, but it seems likely that this is possible for various synthetic molecules.*

<table>
<thead>
<tr>
<th>Enzyme used</th>
<th>Typical labeling protocol for cell surface labeling</th>
<th>Intracellular labeling</th>
<th>Free choice of labels</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sfp-labeling</td>
<td>2 µM Enzyme, 1 µM Label, 30 min</td>
<td>Not shown</td>
<td>Yes</td>
</tr>
<tr>
<td>AcPS-labeling</td>
<td>2 µM Enzyme, 1 µM Label, 20 min</td>
<td>Not shown</td>
<td>Yes</td>
</tr>
<tr>
<td>Biotin Ligase</td>
<td>0.3 µM Enzyme, 10 µM Label, 1-60 min</td>
<td>Yes</td>
<td>No</td>
</tr>
<tr>
<td>Lipoic acid ligase</td>
<td>10 µM Enzyme, 500 µM Label, 5 min</td>
<td>Yes</td>
<td>No*</td>
</tr>
<tr>
<td>Sortase</td>
<td>200 µM Enzyme, 100 µM Label, 10-30 min</td>
<td>Not shown</td>
<td>Yes</td>
</tr>
</tbody>
</table>

*Species: **M** Enzyme, **m** Label, **L** Peptide, **M** Enzyme, **m** Label, **L** Peptide, **M** Enzyme, **m** Label, **L** Peptide, **M** Enzyme, **m** Label, **L** Peptide, **M** Enzyme, **m** Label, **L** Peptide, **M** Enzyme, **m** Label, **L** Peptide, **M** Enzyme, **m** Label, **L** Peptide, **M** Enzyme, **m** Label, **L** Peptide, **M** Enzyme, **m** Label, **L** Peptide, **M** Enzyme, **m** Label, **L** Peptide, **M** Enzyme, **m** Label, **L** Peptide.
PPTase-mediated labeling has been significantly reduced owing to continuous engineering efforts, the tag size for intracellular CoA would interfere with labeling. The method is restricted to labeling cell surface proteins because intracellular CoA would interfere with labeling. Owing to continuous engineering efforts, the tag size for PPTase-mediated labeling has been significantly reduced [22,23]. This makes PPTase-mediated labeling probably the best choice for labeling extracellular small tags with arbitrary substrates. E. coli Biotin Ligase (BirA) is another well established labeling enzyme that can be used for the specific transfer of biotin and of a biotin isostere with a ketone functionality [24]. Recent reports demonstrate that biotin ligases from other species can be used to label proteins with alkyn-functionalized and azide-functionalized biotin [25]. The functionalized biotins can serve as a handle to introduce arbitrary labels in a second step using selective reactive chemistry. BirA can be expressed within mammalian cells, and has been used for the labeling of intracellular proteins (cf., e.g. refs. [26,27]). The limitation of this broadly applicable method is the presence of other biotinylated proteins within cells and the fact that arbitrary labels such as fluorescent dyes cannot be transferred in one step.

A major breakthrough for the intracellular labeling of proteins with various substrates by enzyme-mediated labeling has recently been published, termed lipoic acid ligase mediated labeling. In initial reports, it has been shown that the enzyme efficiently transfers labels containing reactive bromides, azides, and photocrosslinkers to tagged proteins [28,29,30]. In a very recent publication, lipoic acid ligase was engineered to also accept the fluorescent dye coumarin as a substrate, and it has been shown that lipoic acid ligase can be expressed in mammalian cells, enabling intracellular labeling [31]. The utility of this method has been demonstrated for the specific labeling of intracellular tagged proteins in the cytosol and the nucleus. As of now, lipoic acid ligase does not allow a free choice of the label, because the introduced label has to fit into a cavity within the enzyme; new labels therefore require a reengineering of lipoic acid ligase. A further new method uses sortase to mediate specific protein labeling. Sortase is used to specifically join a peptide or protein containing a five-amino acid C-terminal recognition tag (usually LPXTG) with a second peptide or protein containing multiple glycines at its N-terminus. The method has been successfully used for the C-terminal [32,33] and N-terminal labeling of proteins [34,35]. The method works with high specificity on living cells, but is presumably restricted to the cell surface.

Applications of tag-mediated labeling
Study of protein function in living cells
In the past two years, tag-mediated labeling methods have increasingly been applied in live cells to study protein localization, dynamics, and trafficking, as well as the stoichiometry of protein complexes [36,37-41,42]. An illustrating example [40,41] is given by two studies on a protein involved in Hedgehog signaling, which were carried out using SNAP-tag [40] and ACP labeling [41], respectively.

Self-labeling proteins have recently been demonstrated to work in living animals. This allows the introduction of fluorophores that absorb and emit light in the far-red spectral window, which improves imaging in deep tissue compared to autofluorescent proteins. In addition, the labeling time point can be freely chosen, allowing pulse-chase imaging in vivo. Kosaka et al. generated an ovarian cancer cell line stably expressing HaloTag at the cell surface [43]. This cell line was used to generate model tumors in the peritoneal cavity of living mice. The tumors were stained by direct injection of fluorescent HaloTag ligand into the peritoneal cavity and could be imaged noninvasively in the living animals.

Biophysical methods are increasingly being combined with tag-mediated labeling. The chromophore-assisted light inactivation of proteins (CALI) utilizing proteins labeled with a fluorophore via the tetracysteine-tag [44,45] is one such example. This previously established method has again successfully been applied to study the role of the Clathrin light chain in vesicle formation and its impact on neuronal signaling in the neuromuscular junction of drosophila flies [46]. Recently, it has been shown that CALI also works efficiently based on SNAP-tag labeling [47].

Tag-mediated labeling approaches are ideal platforms for the introduction of labels for super-resolution imaging in cells. This has now been demonstrated using HaloTag [48] or SNAP-tag fusion proteins [49] with the stimulated emission depletion (STED) approach, as well as using a SNAP-tag fusion protein with the stochastic optical reconstruction microscopy approach (STORM, Figure 2a) [50]. Super-resolution imaging based on tag-mediated labeling of the proteins of interest combines the advantages of conventional approaches based on antibodies or autofluorescent proteins, namely the possibility to apply super-resolution imaging to living cells and the free choice of the utilized fluorophore [49]. Compared to autofluorescent proteins, synthetic fluorophores offer the advantage of a wider array of available colors, and, depending on the chosen fluorophore, improved photostability. This is important because the attainable resolution depends on the photostability of the utilized dye [80].

As shown in a recent publication [51], tag-mediated labeling is also a good strategy for investigating protein mobility in living cells by fluorescence recovery after photobleaching or photoactivation.
protein association and protein folding. We have recently
interact can only be labeled if two proteins or protein domains
published that utilize a split tetracysteine motif that
cell interior. A number of examples have been
protein interactions with overexpressed proteins in the
is increased by induced proximity if the two tagged
efficiency of biotinylation of the acceptor peptide by BirA

A further example how tag-mediated labeling can be
exploited for the construction of novel tools is a small-
molecule sensor principle as depicted in Figure 4 [60**].
The method is similar to FRET-based biosensors that
rely on a bacterial binding protein sandwiched between a
FRET pair of autofluorescent proteins; the prototype of
this established design was the maltose sensor developed
by Fehr et al. [61], a recent prominent example are the
glutamate sensors developed by the Tsien laboratory
[62]. In a proof-of-principle experiment, the new sensor
principle was tested with human carbonic anhydrase
(HCA) as the binding protein, and used to build a sensor
for HCA inhibitors. The sensor exhibited a favorable
sensitivity compared to previous approaches. The deci-
vative novelty in this design is the fact that sensors for
metabolites can be generated based on metabolite bind-
ing proteins – such as HCA – that do not undergo a
conformational change upon ligand binding.

New tools for biology

The possibility to generate hybrid proteins containing an
expressed and a synthetic part is increasingly being
exploited to make novel tools for biology. A particularly
pressing problem in biology is the study of protein inter-
actions. A previously introduced approach based on the
tetracycstene tag (Figure 3a) uses a trifunctional com-
ound containing a biarsenical dye to bind to the protein
of interest, a crosslinker triggered by addition of sodium
periodate to induce tethering to binding partners, and
biotin for detection of the interaction partner after SDS-
PAGE and western blotting [52]. This affinity labeling
method has recently been applied to study the interaction
between ubiquitin and the proteasome [53]. A different
method based on E. coli Biotin Ligase works by fusing the
two proteins of interest to biotin ligase and to the biotin
ligase acceptor peptide, respectively (Figure 3b).

With tag-mediated labeling, the targeting of synthetic
sensors to specific subcellular locations and organelles is
possible. This allows combining the subcellular targeting
provided by genetic methods with the sensitive and fast
response provided by synthetic sensors. The SNAP-tag
technology has recently been applied to targeting sensors
for Zn²⁺ [63], Ca²⁺ [64–66], and probes for hydrogen
peroxide [67]. This strategy has also been applied to
pH sensors that can be covalently linked to HaloTag
[68]. A recent example for the nuclear localization of a
highly sensitive BODIPY-based Ca²⁺-sensitive dye is
shown in Figure 2b [66]. Targeted ion sensors allow
the detection of local ion concentration changes in the
cell with a high spatial resolution; targeting the sensors to
locations within the cell that experience a large analyte
concentration change will inherently improve signal to
noise ratios for such measurements. The application of
targetable sensors to complex biological problems will
help in revealing the importance of local concentration
changes on cell function.

Incorporation of unnatural amino acids

The most elegant way to obtain proteins containing non-
natural functionalities and labels is to incorporate these
already during protein synthesis using modified amino
New methods for the detection of protein interactions. (a) Label transfer chemistry based on tetracysteine tag labeling. A trifunctional molecule is used containing the biarsenical molecule FlAsH that binds to the tetracysteine tag, biotin for the detection of interaction partners, and a crosslinker inducible by sodium periodate addition. (b) Proximity biotinylation. (c) Selective crosslinking of interacting proteins (S-CROSS). The trifunctional molecule used for crosslinking (SC-Cy5) contains the fluorescent dye Cy5 and the substrates of SNAP-tag and CLIP-tag. (d) Application example for S-CROSS. The data shown correspond to the rapamycin-dependent interaction of FKBP and FRB [59]. The fluorescent SDS-PAGE gel image shows that crosslinking is dependent on the presence of rapamycin and therefore on the interaction of the two proteins.
chemical biotechnology

The roles of both tRNA and aminoacyl-tRNA synthetase in protein synthesis are central to UAA. Each tRNA is loaded with its respective amino acid by a specific aminoacyl-tRNA synthetase. Upon recognition of a codon in the coding mRNA by means of a specific anticodon, the tRNA transfers its amino acid to a nascent protein chain. There are three general ways to exploit these parts of the protein translation machinery to achieve UAA incorporation into proteins: (i) Aminoacyl-tRNA synthetases can also load their cognate tRNAs with a close analog of the respective canonical amino acid. When bacterial strains auxotrophic for the canonical amino acid are grown in the presence of the amino acid analog, the analog will replace the canonical amino acid in all proteins. A recent application of this method is the generation of intrinsically colored proteins using fluorescent tryptophan analogs [69*]. (ii) In vivo, tRNA can be loaded with an unnatural amino acid that does not correspond to its original amino acid. If protein synthesis takes place in the presence of this mischarged tRNA, the unnatural amino acid is incorporated into the nascent protein upon translation of the codon corresponding to the utilized tRNA. To allow an orthogonal introduction of the unnatural amino acid, a tRNA corresponding to a codon not used for natural amino acid encoding has to be used, also referred to as a free codon. This is accomplished by tRNA interacting with a stop codon or a four-base codon. This methodology is usually used in cell-free expression systems, but can also be applied to Xenopus oocytes by injection of the mischarged tRNA. The advantage of this approach is the possibility to use unnatural amino acids that are quite large, with a recent example being the incorporation of the fluorescent dye BODIPYFL into nicotinic acetylcholine receptors in Xenopus oocytes; the labeled receptors could subsequently be imaged as single molecules at the surface of intact oocytes [70*]. (iii) The most elegant and currently most actively explored approach is the generation of pairs of tRNA and aminoacyl-tRNA synthetase that recognize an unnatural amino acid and that are orthogonal to the host cell’s protein production machinery. With this approach, proteins incorporating an unnatural amino acid encoded by a free codon can simply be produced by growing cells equipped with the tRNA/aminoacyl-tRNA synthetase pair in the presence of the unnatural amino acid. Using this method, it is now possible to produce proteins containing various unnatural amino acids at defined positions in milligram quantities from bacterial expression systems [4]. Importantly, the approach has now been successfully implemented in mammalian cells [4], promising various previously impossible applications. Nevertheless, the use of UAA incorporation in mammalian cells could profit from further technology development to make it more accessible to non-expert users. Furthermore, the high concentrations of non-incorporated, free unnatural amino acid might be problematic, for example, if fluorescent amino acids are used for live cell imaging. At the moment, the most promising use of this method in cells therefore lies in applications where the free unnatural amino acid does not interfere with downstream applications, such as the photocaging of protein functions, as has recently been demonstrated by photocaging nuclear localization signals in proteins using a lysine bearing a photocleavable protecting group [71**].

Conclusions and future developments

The utility of protein labeling methods to study protein and cell function has now been widely recognized in the scientific community and is increasingly being exploited. The specific introduction of diverse functional synthetic molecules into proteins by tag-mediated labeling expands the utility of methods such as super-resolution imaging and chromophore-assisted light inactivation (CALI) of proteins, and has led to new developments such as subcellular targeting of synthetic sensors, the generation of semisynthetic small metabolite sensors, and methods for the study of protein interactions like affinity labeling and S-CROSS. Regarding new labeling methods, the introduction of lipoic acid ligase is a major breakthrough for the intracellular labeling of proteins fused to small tags, as the approach promises to also be applicable in vivo. The further engineering of lipoic acid ligase should lead, in the...
future, to a broader available palette of transferable labels. With the self-labeling proteins SNAP-tag and HaloTag and further developments of lipoic acid ligase mediated labeling, the toolbox for fluorescent labeling of tagged proteins from \textit{in vitro} to \textit{in vivo} will probably soon be essentially complete.

What is missing? Firstly, unnatural amino acid incorporation by genetic methods is, in principle, the least invasive method to obtain recombinant labeled proteins. The further development of this method in mammalian cells is eagerly awaited. Secondly, a more rigorous investigation of some methodological aspects of labeling seems a worthwhile endeavor. One area that merits a systematic investigation in this respect is the impact of tag size on protein function. In addition, getting a labeling compound of interest across the cell membrane is a recurring problem for intracellular labeling. New general methods that would allow the delivery of arbitrary compounds into the cell interior are therefore urgently required. Alternatively, an improved set of rules that allows the prediction of the membrane permeability of labeling compounds and therefore a better design of the compounds before their actual synthesis would be very valuable. Thirdly, one thing is still missing in the toolbox for protein labeling: general approaches that work on native, non-tagged proteins. While current methods are restricted to recombinant proteins, native protein labeling approaches would allow the study of proteins in native tissue, in their physiological context and at physiological protein concentrations. Thus, they would allow studies that can currently only be carried out with antibody techniques, but provide further possibilities by tagging with a reporter that is covalently bound and much smaller than an antibody.

We currently see three routes towards this aim. One route has become known as activity based protein profiling [72], and relies on the specific covalent interaction of small molecules that react with the active site of enzymes. However, this labeling procedure is invasive as it leads to inactivation of the enzyme of interest. A less invasive route has been described by Hamachi and co-workers and is based – similar to their self-labeling tag approach (see above) – on turning an initially noncovalent interaction into a covalent one. Here, the noncovalent interaction is provided by a molecule that specifically binds to the native protein of interest, for example, an enzyme inhibitor [73*]. The reactive group used is the tosyl group, which is why this approach has been termed ligand-directed tosyl chemistry (LDT). The molecule mediating the interaction is released when the covalent coupling of the label to the protein takes place, which reduces its impact on protein function compared to previous approaches. This method has been shown to allow selective native protein labeling in the cell interior and even in blood cells of living animals [73*]. The method is remarkably specific, but the general reactivity of the tosyl group with nucleophiles may inherently limit the attainable selectivity towards proteins of low abundance. A third potential route may be provided by enzyme-mediated labeling using engineered enzymes that are specific for native, non-tagged proteins. With ubiquitylation, Nature provides an illustrating general example for protein labeling that works specifically and with the same chemistry on hundreds of proteins within the cell. Unfortunately, the ubiquitylation process is rather complex and involves a number of enzymes that work in conjunction. A different enzyme that may be easier to control is transglutaminase, which catalyzes the formation of a covalent bond between the carboxamide side chain of glutamine and primary alkylamines. The enzyme has been shown to have a relaxed substrate specificity and is capable of reacting with various dissimilar peptides [74,75]. We believe that transglutaminases hold great potential for native protein labeling if it should prove possible to increase their specificity towards target proteins of interest by protein engineering.

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Papers of particular interest, published within the annual period of review, have been highlighted as:
- of special interest
- of special interest


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with a resolution down to 40 nm in the living cells. Using the STED super-resolution imaging approach, the labeled overexpressed in mammalian cells and labeled with tetramethylrhodamine SNAP-tag fusion proteins of cytoskeletal and membrane proteins are localized pulse-chase studies of tagged proteins in living cells. In this study, the proteins of interest are fused to SNAP-tag and/or CLIP-tag, respectively, and incubated with a trifunctional photoconvertible fluorescent probes for protein labeling. This publication presents a novel metabolite sensor design based on the human carbonic anhydrase and the self-labeling proteins SNAP-tag and CLIP-tag. The tags allow the introduction of synthetic fluorophores and an intramolecular inhibitor for human carbonic anhydrase. The sensor is sensitive to inhibitors of carbonic anhydrase, and ratiometric fluorescence changes of up to 100% are achieved using various fluorophore FRET pairs. The proof-of-principle with human carbonic anhydrase shows that the sensor design works without the requirement of a conformational change in the binding protein.

How to obtain labeled proteins and what to do with them


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