Review

Reprint of: Current strategies for the use of affinity tags and tag removal for the purification of recombinant proteins

José Arnau *, Conni Lauritzen, Gitte E. Petersen, John Pedersen

Unizyme Laboratories A/S, Dr. Neergaards vej 17, DK-2970 Harsholm, Denmark

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Abstract

Affinity tags are highly efficient tools for protein purification. They allow the purification of virtually any protein without any prior knowledge of its biochemical properties. The use of affinity tags has therefore become widespread in several areas of research e.g., high throughput expression studies aimed at finding a biological function to large numbers of yet uncharacterized proteins. In some cases, the presence of the affinity tag in the recombinant protein is unwanted or may represent a disadvantage for the projected application of the protein, like for clinical use. Therefore, an increasing number of approaches are available at present that are designed for the removal of the affinity tag from the recombinant protein. Most of these methods employ recombinant endoproteases that recognize a specific sequence. These process enzymes can subsequently be removed from the process by affinity purification, since they also include a tag. Here, a survey of the most common affinity tags and the current methods for tag removal is presented, with special emphasis on the removal of N-terminal histidine tags using TAGZyme, a system based on exopeptidase cleavage. In the quest to reduce the significant costs associated with protein purification at large scale, relevant aspects involved in the development of downstream processes for pharmaceutical protein production that incorporate a tag removal step are also discussed. A comparison of the yield of standard vs. affinity purification together with an example of tag removal using TAGZyme is also included.

Keywords: Affinity purification; Affinity tag; His-tag; IMAC; Recombinant protein; Protein purification; Tag removal; Endoproteases; Exopeptidases; Aminopeptidases; Therapeutic protein purification

With the modern advances in genomics, proteomics and bioinformatics, the number of proteins being produced using recombinant techniques is exponentially increasing. High throughput screening approaches are being performed to rapidly identify proteins with a potential application as therapeutic, diagnostic or industrial enzymes [1]. For this purpose, different expression hosts (e.g., Escherichia coli, Saccharomyces cerevisiae, Pichia pastoris, insect and mammalian cell lines) have been developed to express heterologous proteins [2–7]. Additionally, genomic approaches are being pursued to solve the structure of numerous proteins [8,9].

The above-mentioned screening approaches would not be feasible if specific purification procedures were to be developed for each individual protein. Here, the use of affinity tags enables different proteins to be purified using a common method as opposed to highly customized procedures used in conventional chromatographic purification.

When designing a downstream processing strategy for a protein, the inclusion of an affinity tag might be attractive for a number of additional reasons. In many cases, the protein candidate may exist as a version that includes an affinity tag from its early research stages where no biochemical characterization or functional assay is yet available. For structural studies, more than 60% of the proteins produced include a polyhistidine tag (his-tag, [10]). Additionally, the fact that affinity purification normally results in high yields—often over 90%—makes this
Table 1

<table>
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<th>Tag</th>
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<td>[10,38,85,109]</td>
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<td>Calcium-dependent, mAb-based purification</td>
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<td>Modified streptavidin, elution with biotin analog</td>
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<td>Protein aggregation by temperature shift, intein used to remove tag</td>
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<td>Chitin-binding domain</td>
<td>52</td>
<td>Binds only insoluble chitin (see intein, Table 3)</td>
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<td>Thioredoxin</td>
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<td>Affinity purification with modified resin</td>
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<td>Amylase affinity purification</td>
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<td>NusA</td>
<td>495</td>
<td>Increased solubility in E. coli. Affinity tag needed for purification</td>
<td>[19]</td>
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</table>

* Only a few relevant references are included.

An overview of affinity tags and the design of the protein fusion

Recent reports have included several overviews of the currently available affinity tags for protein production and purification [28–32]. Nevertheless, since the choice of affinity tag and the method for tag removal are mutually dependent, an introduction to affinity tags is given herein for clarity.

Affinity tags can be defined as exogenous amino acid (aa) sequences with a high affinity for a specific biological or chemical ligand. A major group of affinity tags consists of a peptide or protein that binds a small ligand linked on a solid support (e.g., his-tags bind to immobilized metals, discussed below). Another group includes tags that bind to an immobilized protein partner such as an antibody or antibody purification using protein A affinity chromatography ([29], Table 1). The protein A-based methodology, used for e.g., purification of monoclonal antibodies (mAb), has been extensively reviewed elsewhere [33,34] and will not be discussed here.

His-tags are the most widely used affinity tags. Purification of his-tagged proteins is based on the use of chelated thioredoxin (Trx). 1

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1 Abbreviations used: Trx, thioredoxin; 6xHis, a stretch containing six consecutive histidine residues; aa, amino acid(s); AAP, Aeromonas proteolytica aminopeptidase; APM, aminopeptidase M; B, Bacillus; CPA, carboxypeptidase A; CBP, carboxypeptidase B; DAPase, recombinant dipeptidyl peptidase, part of TAGZyme; E, Escherichia; ELP, elastin-like polypeptides; FMN, flavin mononucleotide; FP, green fluorescent protein; GST, glutathione S-transferase; His-tag, a polyhistidine tag; IMAC, immobilized metal-ion affinity chromatography; mAb, monoclonal antibodies; MBP, maltose-binding protein; NMR, nuclear magnetic resonance; pGAP, recombinant pyroglutamyl aminopeptidase; pGAPase, an engineered version of recombinant pGAP used in TAGZyme; PHB, polyhydroxybutyrate; Qcycerase, recombinant glutamine cyclotransferase, part of TAGZyme; ScFv, single chain antibodies.
metal ions as affinity ligands. The metal ion is complexed with an immobilized chelating agent (immobilized metal-ion affinity chromatography, IMAC). Protein separation using IMAC occurs on the basis of interactions between certain aa residues, especially histidine and the metal ions within an immobilized metal chelate [35]. The imidazole side chain of histidine shows high affinity for chelated metals [36].

The use of short histidine stretches or his-tags, typically placed as affinity tags at either the N-terminus or C-terminus, enables the purification of the desired protein from the crude extract of the host cells in a single step. Different chromatographic supports and strategies are available for IMAC [35,37]. The most widespread IMAC supports use either nitrilotriacetic acid (NTA) as a ligand for immobilizing metals like nickel in affinity chromatography (Ni-NTA) or different chelating Sepharose matrices. Additionally, a number of alternatives incorporating different metals have been developed [38]. Importantly, the binding specificity enables the purification of proteins under both native and denaturing conditions [35].

Although universally applicable, the use of his-tags and IMAC purification is not recommended for proteins containing metal ions. Similarly, other aa like cysteine and naturally occurring histidine rich regions in host proteins may result in unwanted protein binding during IMAC purification [39].

Since its development, numerous proteins and peptides have been purified using his-tags, and several therapeutic candidates are in clinical studies [40]. His-tags have also been used for purification of proteins using expanded bed adsorption [37,41].

Amongst other widely used tags, a short affinity tag that uses antibody-based purification is FLAG [27,32,42–45]. FLAG is a hydrophilic octapeptide (DYKDDDDK) recognized by the M1 mAb [43]. Recent studies suggested that a shorter FLAG-related peptide (DYKD) could be recognized with similar affinity by M1 [45]. The binding reaction on an M1-chromatographic support is dependent on calcium, so proteins may be eluted from an affinity matrix by EDTA-containing buffer [42]. Similarly, Softag1 and Softag3 are small peptides (sequences SLAELLNAGLGG and TKDPSRVG, respectively; Table 1) recognized with high affinity by polyol-responsive mAb. The use of Softag allows for the elution of pNSroteins from antibodies under mild conditions. SNSsoftags are especially useful for the purification of labile, multisubunit enzyme complexes and to facilitate the study of protein interactions [46–48].

Another example is Streptag II. It consists of a streptavidin-recognizing octapeptide (WSHPQFEK) that was selected for its improved affinity. Proteins containing a Streptag II can be purified by affinity using a matrix with a modified streptavidin and eluted with a biotin analog [29,49–52]. An inherent advantage of Streptag II is its independence from metal ions in the purification, an aspect often of interest when metalloproteins are to be studied or when paramagnetic impurities must be avoided for NMR [51]. Other commonly used affinity tags are illustrated in Table 1.

The choice of a suitable affinity tag depends both on the type of application for the protein of interest and the stage of development of the protein for e.g., a therapeutic drug candidate. Additionally, the costs of the different chromatographic supports and the scalability of the process may be influential. In general, the cost of affinity tags based on mAb binding might be prohibitive for large-scale processes [29].

Furthermore, the selection of an affinity tag—and particularly if the tag is to be removed in the final product—determines the genetic design of the fusion protein used (Fig. 1).

For affinity purification, the tag may be placed at either end of the protein or in a region with appropriate surface exposure to allow binding or recognition [44]. To enable removal of the tag, a linker region is typically included.

![Fig. 1. Different strategies to incorporate and remove affinity tags.](image)

### Table 1.

<table>
<thead>
<tr>
<th>Tag Type</th>
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</tr>
<tr>
<td>Affinity/solubility tag</td>
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<tr>
<td>Solubility/folding tag</td>
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Fig. 1. Different strategies to incorporate and remove affinity tags. Different fusion proteins including the different elements required for affinity purification and tag removal are depicted. For clarity, only N-terminal fusions are shown. (1) A fusion protein consisting of an affinity tag, a linker region including a specific sequence for endopeptidase cleavage and the native protein; (2) similar to above but the affinity tag (e.g., GST, MBP) increases the solubility of the protein; (3) a fusion protein designed for exopeptidase removal of the tag (only for N-terminal tags, using TAGZymec); (4) a fusion protein where a solubility and folding partner is fused N-terminal to the target protein (e.g., SUMO, sortase A). An affinity tag at the N-terminus is required for purification.
between the tag and the native protein sequence. This linker region may contain a sequence specific for endoprotease cleavage (construction 1 and 2, Fig. 1). The linker contributes to increased accessibility of the affinity tag and is often required for effective endoprotease cleavage. Although the number of different tags available may reflect the different biochemical properties of proteins and the host cells, it is possible to discriminate tags needed for purification (e.g., his-tag) from those developed to aid, e.g., solubility and folding like maltose-binding protein (MBP) [53], glutathione S-transferase (GST) [54] or small ubiquitin modifying protein (SUMO), among others [55,56]. Some tags such as MBP or GST are used for both affinity purification and solubility (construction 2, Fig. 1). Although GST fusions have been widely used in research, the slow binding kinetics of GST to glutathione-Sepharose resin represents a severe drawback since loading of cell extracts to the resin is extremely time consuming for large-scale production [28].

As mentioned above MBP, GST and thioredoxin (Trx, Table 1) have long been used to increase the solubility of complex proteins in E. coli (construction 2, Fig. 1) [16,18,32,53,57–59]. Remarkably, not all fusion proteins containing MBP can be purified on amylase [58]. MBP fusions have also been reported to result in protein degradation [60]. His-tags may be used to aid solubility and folding since both native and denaturing conditions can be used during purification [16]. For example, his-tags may be used as the sole tag for purification and folding when located at the N-terminus of the protein [61], e.g., if exopeptidase cleavage is used for tag removal [62]. Thus, the simplest genetic design includes the affinity tag and the native protein sequence without a linker region (construction 3, Fig. 1).

Other emerging methodologies for affinity purification and tag removal

Novel fusion tags to increase solubility such as a small chaperone, Skp, have successfully been tested for production of otherwise insoluble proteins in E. coli [56]. Other tags used to increase solubility in E. coli include NusA and T7 protein kinase (Table 1) [56,59].

Ubiquitin-based tags have been used to increase expression levels but also to aid in protein folding. One of these technologies (SUMO) has emerged as an alternative for the production, solubility and correct folding of otherwise intractable proteins (construction 4, Fig. 1). When fused to SUMO, insoluble proteins have been observed to fold properly and become soluble [55]. The SUMO tag can be removed using a specific protease (e.g., the yeast SUMO protease-1 Ulp1) that recognizes the conformation of the ubiquitin partner rather than a specific sequence [55]. However, an additional affinity tag is required for initial purification of the fusion protein when using SUMO. The use of SUMO is mostly constrained to E. coli, since highly conserved SUMO proteases are present in eukaryotes that may cleave the fusion protein during fermentation. A split SUMO system has been proposed to overcome this problem [55]. Additionally, the high ratio of SUMO protease to fusion protein required to process fusion proteins remains a challenge for the upscaling of production processes based on this technology.

Elastin-like polypeptides (ELP) consist of several-to-numerous repeats of a peptide motif that undergo a reversible transition from soluble to insoluble upon temperature upshift (Table 1) [63–66]. ELP tags have been developed that allow purification of the fusion protein by temperature-induced aggregation, separation by centrifugation and resolubilization. Importantly, the length of the ELP tag sequence has been shown influential on the yield [64]. Thus, no chromatographic support is needed for initial purification of the fusion product. Further refinement of this technology includes a tripartite system that incorporates intein to enable tag removal (see below) in E. coli [66]. The robustness of ELP tags remains to be shown for large-scale processes of commercially relevant proteins. However, comparable protein yields have recently been shown for selected proteins using ELP and his-tag at small scale [65].

Inteins are self-cleavable proteases [67]. The intein-based IMPACT system uses a protein fusion consisting e.g., of an N-terminal chitin-binding domain (affinity tag), a central intein and a C-terminal target protein [68]. Binding to a chitin matrix is followed by on-column cleavage using either a thiol reagent or pH and temperature shift to yield intein cleavage and elution of the target protein [67,69,70].

Yet another system for tag cleavage that allows for one-step, on-column purification employs a protein fusion including the catalytic core of the Staphylococcus aureus sortase A, which recognizes the sequence LPETG and cleaves the TG peptide bond (Table 1). The suggested protein design includes an N-terminal his-tag for IMAC purification, followed by the sortase A domain, the cleavage sequence and the target protein at the C-terminus. This allows for sortase-mediated, on-column tag cleavage in the presence of e.g., calcium [71].

For purposes such as following the fate of the target protein through expression and purification, a number of tags have been developed in recent years [72,73]. Rainbow tags constitute such a tool that can be used to visualize correctly folded proteins and to track the protein during purification [72]. For this purpose, the flavin mononucleotide (FMN)-binding domain of cytochrome P450 reductase (displaying a blue-green or yellow color depending on the oxidation state of the FMN cofactor) and the red colored, heme-binding cytochrome b5 have been used [72,74]. Opposite to other visual tagging system like GFP that require an external energy source, rainbow tags are visible with the naked eye. The use of rainbow tags requires an additional affinity tag for purification.

One remarkable example of more untraditional affinity tags is the use of polyhydroxybutyrate (PHB). The system includes production of PHB granules in E. coli together with a fusion containing (i) a PHB-binding domain, (ii) an intein for tag removal and (iii) the protein of interest [75].
Combining different affinity tags

In recent years, a number of approaches have been published that utilize two different affinity tags fused to the target protein (dual affinity tags, also referred to as tandem affinity purification or TAP) for purification. In this approach, a protein of interest is fused in-frame with an N- or C-terminal tag comprised of two affinity tags surrounding an endoprotease cleavage site [76]. Applying two different affinity purification regimes sequentially, a highly purified protein is obtained [7,32,49,77,78]. These approaches are therefore directed to the production of a highly pure protein for e.g., crystallization [7,49] but are also widely used for the isolation of protein complexes [76,78,79] to study protein interactions. Originally developed in yeast, the method utilizes protein A and calmodulin-binding peptide separated by a specific TEV protease cleavage site and fused in-frame to the target protein. The TAP-tagged protein is expressed at physiological conditions are optimal for di-cation [76]. Once the protein complex has been purified through two affinity chromatography steps as above, it is resolved by SDS-PAGE into its components and protein bands are digested in-gel and identified by mass spectrometry [79]. A number of additional tag combinations have been used in other organisms for TAP and for the study of protein networks [76].

Additionally, dual tagging has been developed to monitor expression (via green fluorescent protein, GFP) and purify proteins via his-tags located within GFP [30].

Comparison of a standard purification process with affinity purification

Numerous reports have been published where different purification procedures are compared [1,7,12,17,29,30,52,67,78,80]. Generally, the yield and efficiency of any specific purification procedure depends on the level of optimization developed for the individual protein and method. It is therefore recommended to use the data presented in different comparisons as indicative rather than definitive since it is unlikely that e.g., identical elution conditions are optimal for different proteins. As an example, suboptimal elution of his-tag proteins has been reported that is likely due to the low imidazole concentration used. Similarly, impurities—that may be accounted for by a certain level of metal leakage from the IMAC support—are observed for the purified his-tag proteins. This is a likely consequence of the use of β-mercaptoethanol during extraction [29]. As mentioned above, unspecific protein binding may occur during IMAC purification and several approaches have been used to reduce this problem, including the use of higher imidazole concentration in the wash buffer [39]. Moreover, the concentration of imidazole used for wash and elution is a powerful tool to fine-tune the purification process for increased purity and specificity. Typically, elution of a ‘detagged’ protein after the his-tag removal step is performed on an (subtractive) IMAC column in the absence of imidazole to facilitate flow through of the protein while the his-tag process enzymes and other unspecific binders are retained in the column.

Standard chromatographic methods often include an initial capture step where the proteins present in the crude extract are bound to the matrix followed by gradual retention/elution of the protein of interest. Several subsequent chromatographic steps are needed to obtain a relatively pure protein. This results in time-consuming procedures and a relatively low yield of recovery, typically below 50% of the starting material for optimized processes. Remarkably, the yields obtained in purification of proteins using affinity chromatography are typically over 90% and include a reduced number of steps, adding to the reduction in costs of manufacturing. Importantly, IMAC are chemically stable to the prolonged cleaning-in-place procedures widely used in pharmaceutical production.

To illustrate this, we used a Bacillus amyloliquefaciens pyroglutamyl aminopeptidase (EC 3.4.19.3, pGAP) as a model protein. This protein catalyzes the removal of N-terminal pyroglutamyl residues from peptides and proteins. For comparison purposes, pGAP was produced in E. coli with and without an N-terminal his-tag (HT-pGAP, tag sequence: MEP(H)6L). For untagged pGAP, a purification process was performed that included ammonium sulfate precipitation, two consecutive separation steps using phenyl-Sepharose and a final step using Q Sepharose HP. As shown, many proteins co-purified with pGAP during the phenyl-Sepharose separation steps (Fig. 2A). In spite of an effort for the optimization of these chromatographic steps, especially with regard to the pH used in the final Q Sepharose HP step, a pGAP preparation was obtained where several contaminating bands were apparent, as observed by SDS-PAGE (Fig. 2A). Additionally, activity measurement showed a 40% recovery for this process with significant protein loss at step one and three, respectively (Table 2). For HT-pGAP, a higher amount of protein was observed in the cell extract. The reasons for this difference with the untagged pGAP remain unclear but may result from a higher expression level for this protein [80,81]. However, yield discrepancies due to intrinsic biological variation or culture conditions cannot be excluded. Remarkably, a very pure protein was obtained in a single IMAC step (Fig. 2B) and the yield of the process reached 96% without optimization (Table 2). Moreover, especially for high throughput approaches where no knowledge of the biochemical properties of the proteins is available, affinity tags—and his-tag in particular—may represent a universal platform for purification [1,43,62,82].
Methods for affinity tag removal

In a few specific cases, the affinity tag can be removed by harsh chemical treatment with e.g., cyanogen bromide or hydroxylamine. For this purpose, a unique methionine residue must be placed at the junction between the fusion partner and the protein of interest [83,84]. Chemical methods are rather unspecific and may lead to protein denaturation and side chain modification of amino acids in the target protein [30]. Enzymatic methods are more specific and cleavage is achieved usually under mild conditions.

Most of the available methods for affinity tag removal include enzymatic cleavage of the tag followed by specific removal of the process enzyme(s) by an affinity chromatographic step to yield the detagged protein. Several affinity-tagged process enzymes are therefore used for tag removal.
(Table 3) [30,37,62,67,85–87]. The rationale behind this approach is the need of a step for the efficient removal of the processing enzyme after the tag is removed to ensure that no contaminating protease activity is present in the protein preparation (illustrated for his-tag purification in Fig. 3).

Several endoproteases have been developed for tag removal (Table 3). Enterokinase, thrombin and factor Xa are among the most widely used enzymes for this purpose [37,42,88–93]. Enterokinase and factor Xa are examples of proteases that are able to cleave without requiring a specific aa or sequence at the C-terminus of the cleavage site (Table 3), allowing for the complete removal of the tag and yielding the native protein. Production of a tag-free protein is especially important for the manufacture of therapeutic proteins. Cleavage of the tag using other endoproteases like Thrombin, and virus-derived TEV protease, 3C protease or similar enzymes results in one or two aa residues from the cleavage site left at the N-terminus of the processed protein (Table 3). In these cases, documentation for the effect of the residual exogenous aa left in the therapeutic protein may be required.

Other classes of endoproteases have also been developed for tag removal. Granzyme B is a lysosomal serine protease involved in a number of protein activation processes in higher eukaryotes. An engineered, autoactivable granzyme B has recently been developed for tag removal [94], although its broad substrate specificity (Table 3) may represent a disadvantage if unspecific degradation of the target protein occurs. Caspase-6, a protease involved in protein processing during apoptosis, has also been used for tag processing of GST-fusions [94].

A general drawback of endoprotease-mediated tag cleavage is the need for high ratios of enzyme to protein and long incubation time required for complete tag removal. A further complication is the fact that under the above severe conditions, endoprotease cleavage often leads to undesired, non-specific cleavage of the protein at cryptic sites (e.g., secondary GR sites for factor Xa or other basic aa for enterokinase, Table 3), resulting in protein degradation and lower yields [59,88,89,95–97]. In some cases, cleavage is observed at secondary sites but not at the introduced cleavage site [99].

Generally, processes for large-scale production of therapeutic proteins would encourage the use of on-column affinity purification and tag removal since a clear benefit applies to the reuse of columns and process enzymes for the production of large amounts of protein. Consequently, a number of on-column processes have been developed for tag removal by combining the most commonly used endoproteases with the most efficient affinity tags like his-tag, GST, FLAG, etc. [12,19,30,42,71,85,90,91,97,98]. Using relatively high amounts of endoprotease for tag removal must be accompanied by a process where efficient capture/immobilization of the enzyme is achieved. Several issues related to process validation have to be considered when adopting such a strategy to ensure e.g., no leakage of the endoprotease from the column during purification.

Furthermore, ensuring that no cryptic sites are present in the native sequence of the protein is a pre-requisite when using endoproteases. Contaminating proteases derived from either the host organism or the endoprotease preparation may also contribute to product degradation during extended incubation or incubation with high concentration of protease.

The use of exopeptidases for tag removal: TAGZyme

An alternative strategy to endoprotease cleavage of affinity tags is the use of exopeptidases for tag removal. Several aminopeptidases and carboxypeptidases are available from natural sources. Examples of these are aminopept-
Aprotinin (APM) obtained from porcine kidney and carboxypeptidase A and B (CPA and CPB) from bovine pancreas (Table 3) [99]. APM releases a single aa sequentially from the N-terminus of a protein with P being released slowly. CPA and CPB display different substrate preference, e.g., R is not released by CPA and rapidly processed by CPB. In general, with the growing concern on contamination from animal sources, the potential use of these enzymes for tag removal in pharmaceutical production is somewhat limited. Additionally, since these natural proteins lack an affinity tag, their removal represents an additional challenge for the purification of the target protein after tag cleavage.

Caution must also be used when choosing an amino- or carboxypeptidase that is a member of a metalloprotease family for the removal of his-tags. The metal ion required for peptidase activity may be sequestered by the his-tag in the cleavage step, yielding the peptidase inactive. Aeromonas proteolytica aminopeptidase AAP is a small zinc metalloenzyme that has been used for the removal of a short natural tag in the production of human growth hormone [100,101]. However, no detectable cleavage is observed during the treatment of different his-tag proteins with AAP (unpublished data).

A method to circumvent the above limitations and to exploit the specificity of exopeptidase cleavage is provided by DAPase, a recombinant rat dipeptidyl aminopeptidase I (DPPI) that is part of the TAGZyme system [62]. DPPI is a cysteine peptidase that sequentially removes dipeptides from the N-terminus until a stop position is encountered in the sequence [62,99,102]. Using DAPase alone—for proteins that include a suitable stop position at the N-terminus—or in combination with two accessory enzymes acting on the N-terminal aa, for all other proteins [62,103,104], efficient and precise cleavage of short (≤ 25 aa) N-terminal affinity tags is obtained. This approach enables the purification of any protein and its production with the native N-terminus [62,99,104]. Characteristic for DPPI and DAPase and different from typical proteases and other peptidases, the active site is located on the surface of the protein enabling rapid processing of peptides and proteins [102]. This feature reflects on the low DAPase requirement for tag removal, typically a 1:5000 DAPase to tagged protein ratio (corresponding to 0.2 μg DAPase per mg of protein or 200 ppm). Moreover, complete tag removal is achieved using short incubation times, typically under 1 h [62].

Many human proteins do include a suitable stop position for DAPase cleavage at the N-terminus, such as a P at position two or three of the mature protein. Examples of these include several interleukins (IL-2, IL-3, IL-5, IL-10 and IL-13) and growth hormone. R or K at position 1 represents also an effective stop position for DAPase cleavage [62]. Human lactalbumin and lysozyme are examples of this class of proteins. For the above human proteins, a short, even-numbered affinity tag such as a 6×his-tag maybe fused directly to the sequence of the mature protein. Subsequently, tag removal can be effectively performed using DAPase alone [62].

For all other proteins without a stop position at the N-terminus, an additional Q is added as the last residue of the affinity tag (placed at an uneven position) adjacent to the desired N-terminal aa of the mature protein. Upon DAPase cleavage in the presence of Qcyclase, the occurrence of an N-terminal Q leads to the effective formation of a N-terminal pyroglutamyl that prevents further DAPase cleavage. Subsequent removal of DAPase and Qcyclase by IMAC and treatment with pGAPase (e.g., on-column) removes the pyroglutamyl residue yielding the protein with the native N-terminus [62]. Typical yields for tag removal using TAGZyme are 90–98% for optimized processes.
Process design for affinity purification and N-terminal his-tag removal

As mentioned above, most pharmaceutical applications necessitate that recombinant proteins do not contain any exogenous aa residue. A combination of the high yields obtained using for e.g., IMAC purification, the need for efficient tag removal and the fact that the latest generation of process proteases are typically tagged to enable their removal in the purification process has to be considered when designing a production and purification process.

Evidence for better yield using N-terminal tag compared to a C-terminal has been reported [22] although this is protein-dependent to some extent. Placing the tag at the N-terminus may also aid to increase expression levels at least in a bacterial host. Moreover, tag sequence optimization can be performed to match the host preferences. A simple process for purification and tag removal of a his-tag protein (N-terminal) includes a first IMAC step, a tag removal step followed by a second subtractive IMAC where the native protein is eluted at high purity. One example of this class of processes is illustrated below.

A process integrating production of his-tag Trx in E. coli and tag removal

Trx was chosen as a well-characterized model protein to attempt the study of the molecular interaction of metal with histidine in a minimal sequence [105]. A process was therefore designed for the production of his-tag Trx (HT-Trx) in E. coli. The genetic design included a 10-aa his-tag (MKHQHQHQHQH) for initial IMAC purification, followed by a short HHP sequence adjacent to the first residue (S) of the native Trx (HHP-Trx, Fig. 4). The occurrence of a P in this sequence allows the removal of the 10-aa tag using DAPase alone and protects from further degradation (Fig. 4). The presence of HH at the N-terminus of the processed, detagged protein was decided to enable a detailed study of nickel interaction by paramagnetic NMR relaxation [105].

The fact that the desired mature protein sequence includes two H residues represented a challenge for IMAC purification where effective elution of HHP-Trx and retention of DAPase (containing a C-terminal his-tag) [62] is needed. In this and similar cases, the subtractive IMAC step is performed using a buffer containing a low concentration imidazole that allows for binding of the tag removal enzyme(s) to the IMAC while precluding binding of the detagged protein (Fig. 4).

The 10 aa his-tag sequence allows high expression of the recombinant protein in E. coli [62]. Additionally, the MK motif serves a double purpose. First, the presence of N-terminal MK results in a low frequency of methionine excision in E. coli [106,107]. Second, the presence of K at position 2 represents a quality control for DAPase processing. Thus, the fraction of HT-Trx molecules where M excision occurs is not cleaved by DAPase (an N-terminal K is a natural stop position for DAPase) [62]. Consequently, these molecules retain a functional his-tag and are effectively removed in the subtractive IMAC step.
HT-Trx was initially purified on an IMAC (Fig. 4). After subsequent removal of the tag using DAPase and desalting, approx. 443 mg protein was obtained from approx. 2.5 L E. coli culture. Subsequent DAPase removal was achieved by a subtractive IMAC step. The final HHP-Trx preparation (354 mg) was obtained after a last desalting step (Fig. 4). No unspecific cleavage of HHP-Trx was observed, emphasizing the lack of endoprotease activity and the efficient protection from further exopeptidase cleavage of the tag provided by P (Fig. 4).

The use of the highly pure HHP-Trx in NMR studies demonstrated that HH forms a stable dimer complex with nickel, where each metal ion binds two HHP-Trx molecules with different orientation [105].

**Emphasis on the design of the tag is essential**

In the example above, a tag was designed to enable a process for the purification and removal of a his-tag to yield a pure protein. In this case, the tag design enabled removal by cleavage with DAPase alone. As described above, for proteins which do not include a suitably placed DAPase stop position, a his-tag maybe used that includes Q at the C-terminus of the tag and at an odd position. Virtually, any protein can therefore be produced using this approach. However, a critical concern is the design of the tag and the sequence used to ensure that tag removal is effective.

One important issue is the compatibility between affinity tag sequence and the specificity and sequence constraints of the enzyme chosen for tag removal. An example is the FLAG sequence that includes a cleavage site for enterokinase (DYKDDDDK, cleavage site underlined) [42,43]. Recent studies suggest that a shorter peptide, DYKD, can also be recognized with almost the same affinity by the M1 mAb used for the octapeptide FLAG version [43]. A tag sequence that includes DYKD can be used for protein purification and tag removal using TAGZyme only if the K residue is located at an even position in the fusion protein, e.g., by adding an uneven number of aa at the N-terminus and ensuring a stop position is placed at the appropriate location. Thus, a recommended FLAG tag designed for production in E. coli and effective removal using TAGZyme could be e.g., MK AD YK DQ Q (added residues underlined). MK is added to avoid processing of protein molecules where methionine excision has occurred and to facilitate high expression in E. coli. A is added for effective DAPase cleavage and to avoid K at an uneven position. Finally, QQ is added for effective DAPase cleavage (of the DQ dipeptide) and to introduce a stop position that can be removed using pGAPase after Qeyclase conversion of the second Q to pyroglutamyl.

To design an effective process for tag removal using endoprotease cleavage, the linker region and the cleavage sequence need to be carefully considered together with the sequence of the protein to be produced. This is required to avoid unspecified cleavage at cryptic/secondary sites that is often encountered [88,89].

**Conclusion**

The market for protein drugs is expected to increase significantly from 2004 ($34 bn) to 2010 ($52 bn). New protein drugs currently under development will be introduced and yet novel protein drugs will be developed [108]. Generic protein drugs are also a competitive market where the use of more effective purification processes will add a significant advantage. In a competitive pharmaceutical market where production costs are high, the development of more economically sound production and purification processes may be a trend to follow.

Genome-based projects directed to the elucidation of protein structure or function and also strategies that make use of directed evolution of existing proteins will also benefit from a universal platform for production and purification. In this scenario, the use of affinity tags may represent a valid alternative to costly standard chromatographic methods for purification.

A variety of affinity tags and methods for tag removal have been developed. Most of these methods are suitable for laboratory scale applications and their use in pharmaceutical production is therefore not straightforward. In this scenario, an integrated view of the production and purification process that includes choice and design of the affinity tag together with the choice of tag removal strategy is required.

Although several tags have been extensively used over the years, his-tags are the unquestionably preferred affinity tag for protein purification. Other tags are also generally used for enhanced solubility and folding. The major advantages of using a his-tag compared to standard purification were illustrated with a model protein, B. amyloliquefaciens pGAP. A significant increase in yield and a reduction of chromatographic steps are achieved using a his-tag. Moreover, the impurities observed in the pGAP preparation obtained using standard chromatography were effectively avoided during IMAC purification of HT-pGAP.

For therapeutic applications, the tag has to be removed from the fusion protein. Here, the choice of affinity tag has to be considered as an integral process that includes a method for tag removal. Methods for tag removal are also abundant for research applications. Most of these use endoproteases and a cleavage site in the fusion protein. Major concerns associated with these approaches for pharmaceutical applications are the relatively high amounts used and the need for subsequent effective removal and unspecific cleavage of the target protein. Also, residual aa after cleavage may be unacceptable for therapeutic use in humans. An alternative to circumvent these limitations is the use of exopeptidases for removal of N-terminal tags. The use of a purpose-designed his-tag in E. coli was illustrated for the production of HHP-Trx. The N-terminal tag was effectively removed using DAPase, and enzyme removal using subtractive IMAC resulted in a pure HHP-Trx.
For affinity purification and tag removal, a careful consideration for the design of the tag is necessary to ensure an effective process. Finally, the potential application of TAG-Zyme technology is also illustrated for FLAG.

Using his-tag purification and tag removal is bound to become the method of choice for pharmaceutical production in the years ahead as well as a universal platform for protein structural studies and genomic projects. The different methods available discussed here should assist in designing a more effective production process.

References


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