The Fh8 tag: A fusion partner for simple and cost-effective protein purification in *Escherichia coli*

Sofia J. Costa, Eduardo Coelho, Lara Franco, André Almeida, António Castro, Lucília Domingues

Abstract

Downstream processing is still a major bottleneck in recombinant protein production representing most of its costs. Hence, there is a continuing demand of novel and cost-effective purification processes aiming at the recovery of pure and active target protein. In this work, a novel purification methodology is presented, using the Fh8 solubility enhancer tag as fusion handle. The binding properties of Fh8 tag to a hydrophobic matrix were first studied via hydrophobic interaction chromatography (HIC). The Fh8 tag was then evaluated as a purification handle by its fusion to green fluorescent protein and superoxide dismutase. The purification efficiency of the Fh8-HIC strategy was compared to the immobilized metal ion affinity chromatography (IMAC) using the His tag. Results showed that the Fh8-HIC binding mechanism is calcium-dependent in a low salt medium, making the purification process highly selective. Both target proteins were biologically active, even when fused to Fh8, and were successfully purified by HIC, achieving efficiencies identical to those of IMAC. Thus, the Fh8 acts as an effective affinity tag that, together with its previously reported solubility enhancer capability, allows for the design of inexpensive and successful recombinant protein production processes in *Escherichia coli*. © 2013 Published by Elsevier Inc.

Introduction

The continuing growth of biotechnology industry demands new strategies for the rapid and economical recombinant protein production through the use of a variety of host organisms and solubility as well as affinity partners [1–5]. The purification of a protein of interest from biological mixtures using rapid, robust and cost-effective methodologies is still a current challenge for academia and industry. Taking into account that the downstream processing comprises up to 80% of the production costs [2], novel solutions that simplify the protein purification process are essential for the biotechnology’s progress. This purification bottleneck has been fairly overcome by the use of affinity fusion partners together with DNA recombinant techniques that allow to clone in frame the peptide or protein affinity tag at the N- or C-terminus end of the target construct [2]. These fusion partners or tags diverge in molecular size and complexity, and can also be used to improve soluble protein production besides facilitating its purification by specific interaction with a known ligand/adsorption matrix [3]. Several affinity tags are commercially available for research or large scale protein production as, for instance, the Glutathione S-Transferase (GST) tag [6], Maltose Binding Protein (MBP) tag [7] and Hexahistidine tag [8] that have affinity to bind immobilized glutathione, maltose and nickel, respectively. Although being widely used, these fusion tags often yield low levels of protein purity due to unspecific and/or weak interactions with the corresponding matrices [9]. New purification tags are constantly emerging, outperforming the existing techniques and advancing the affinity concept or protein detection, as for instance, the Si-tag [10], Tamavadin tag [11], Tab2 tag [12], intein-mediated purifications [13], Heme tag [14], Z-basic tag [15], Dock tag [16] and the HiCaM tag [17]. The ideal purification process should: allow efficient and high yield protein recovery from a biological mixture; be universally applied to any protein without disturbing its function; use a small fusion partner; be compatible with native electrophoresis.

Abbreviations used:

- GST, glutathione S-transferase
- MBP, maltose binding protein
- Trx, thioredoxin
- NusA, N-utilization substance A
- GFP, green fluorescent protein
- SOD, superoxide dismutase
- TEV, tobacco etch virus
- MCS, multiple cloning site
- HIC, hydrophobic interaction chromatography
- IMAC, immobilized metal ion affinity chromatography
- SDS–PAGE, sodium dodecyl sulfate–polyacrylamide gel electrophoresis

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conditions; and it should offer great control of selectivity (binding and elution of the protein of interest) while using inexpensive and high capacity resins [5,18].

In a previous work [21] we have shown that the Fh8 peptide could be used as a solubility enhancer partner, improving protein expression and solubility as the well-known Thioredoxin (Trx), N-utilization substance A (NusA) or MBP fusion partners. Moreover, we have also shown that small fusion tags allowed an easier evaluation of the target protein solubility. Thus, when compared to larger fusion tags, the Fh8 tag was considered an advantageous option for soluble protein production in Escherichia coli due to its low molecular weight and its efficient solubility enhancing effect.

In this work, we investigate the Fh8 solubility enhancer tag as a novel fusion purification handle by combining the calcium binding intrinsic property of the Fh8 molecule with an inexpensive hydrophobic resin (phenyl-Sepharose). The Fh8 (GenBank accession No. AF213970) was first isolated from the excreted/secreted proteins of the Fasciola hepatica parasite and recombinantly produced in E. coli for diagnostic purposes, presenting a molecular weight of 8 kDa [19]. This recombinant protein was previously characterized as a calcium sensor protein that changes its structure upon calcium binding, exposing its hydrophobic residues to interact with its targets or other molecules [20]. We demonstrate here that the solubility enhancer Fh8 tag [21] interacts with the phenyl-Sepharose hydrophobic resin with an identical mechanism as other calcium-binding proteins [22,23]. Furthermore, using the green fluorescent protein (GFP) and the superoxide dismutase (SOD) as target model proteins, we also reveal that the Fh8 tag can be successfully applied as a purification handle for simple, rapid, and low cost recovery of biologically active proteins. By acting simultaneously as a solubility enhancer tag [21] and purification handle at the same time, the Fh8 offers an efficient and economical recombinant protein production in E. coli.

Materials and methods

Cloning of sod and gfp genes into expression vectors

Both gfp and sod genes were first modified and amplified by PCR to be later sub-cloned into the pETM11 (EMBL) and pETMFh8 [21] expression vectors. Both vectors present a N-terminal His6 tag followed by a recognition site for the Tobacco Etch Virus (TEV) protease and the Multiple Cloning Site (MCS). In the case of pETMFh8, the Fh8 tag is located between the His6 tag and the TEV recognition site [21].

The PCRs were conducted using minipreps (GenElute™ Plasmid Miniprep Kit – Sigma) of gfp and sod harboring plasmids as templates and the Phusion High-Fidelity DNA Polymerase (New England Biolabs – NEB), according to the manufacturer’s instructions. Specific primers were designed as follows: Forward primer: 5′-TCTATTCTCATGGATCC-3′ and Reverse primer: 5′-AATAGAATCCGATTTA+21 nt gfp/sod-3′, to introduce the NcoI/BamHI restriction sites (underlined) at the N-terminal of both genes and the XhoI restriction site (underlined) after the stop codon (bold) at the C-terminal of both genes. After DNA purification (QiAquick DNA gel extraction kit – Qiagen), the PCR products and the expression vectors were digested using the NcoI and XhoI restriction enzymes (New England Biolabs) and DNA ligations were carried out using the Rapid DNA Ligation kit (Roche).

E. coli TOP10 competent cells were transformed with the obtained vectors and the resulting clones were verified by colony PCR using the T7 forward and reverse universal primers. The correct insertion of gfp and sod genes into the pETM11 and pETMFh8 expression vectors was additionally confirmed by sequencing with both T7 forward and reverse universal primers.

Expression of Fh8 tag, HisGFP/SOD and Fh8GFP/SOD fusion proteins in E. coli

The expression strains and induction conditions used in this work were selected from a small-scale screening using 10 ml cultures (see Supplementary file 1). GFP recombinant proteins and the Fh8 tag were expressed in 2 L cultures using the E. coli BL21 Codon Plus-RIL strain. SOD recombinant proteins were expressed in 2 L cultures of E. coli Rosetta strain. Recombinant proteins were expressed as follows: pre-cultures were grown overnight (o/n) at 37 °C and a dilution factor of 50 was used for inoculation of all cultures. Cells in a total culture volume of 2 L (eight 1-L flasks of 250 mL of culture media) were grown at 37 °C and 200 rpm to a final OD600nm of 0.4–0.6. E. coli cultures were induced with isopropyl-β-D-thiogalactopyranoside (IPTG) at 0.2 mM, 18 °C, o/n (for SOD and GFP expressions) or at 1 mM, 30 °C, 4 h (for Fh8 tag expression). After induction, three 5 mL samples were taken from each culture for dry weight estimation and the remaining cells were harvested for 25 min, at 4 °C and 4000 rpm. Cell pellets were washed once with phosphate buffer saline 1× (PBS 1×) and collected again by centrifugation. Bacterial pellets were then stored at −20 °C.

Stock solutions of antibiotics for plasmid maintenance and protein expression were prepared, filtered through 0.2 μm and stored at −20 °C to be used in culture media with the following concentrations: kanamycin 50 μg mL⁻¹ and chloramphenicol 10 μg mL⁻¹.

Fh8 purification by HIC

Three different HIC purifications were tested to evaluate the interaction of Fh8 with the hydrophobic resin in the presence and absence of calcium, following the strategy presented in Supplementary file 2. Cell pellets of 1.5 L culture of the Fh8 tag were thawed and resuspended in a total volume of 3 × 25 ml of lysis buffer (50 mM Tris pH 7.6 and 150 mM NaCl, supplemented with 1× complete free EDTA protease inhibitor (Roche), 5 μg mL⁻¹ DNAse (Sigma) and 1 mg mL⁻¹ lysozyme (Sigma)). The lysis buffer was also supplemented with 5 mM CaCl₂, accordingly. After resuspension, cells were incubated at room temperature for 10 min and then lysed by sonication (Branson) for six cycles of 30 s each, with 30 s intervals. Aliquots of 100 μL of total lysates were taken and stored at 4 °C. Supernatant fractions were collected from the insoluble debris by centrifugation at 10,000 rpm, 30 min, 4 °C and aliquots of 100 μL were stored at 4 °C for Bradford estimation of the total protein content and for sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS–PAGE) analysis. The Fh8 tag purifications were conducted in the Biologic DuoFlow FPLC system (Bio Rad) using a 5 ml pre-packed Phenyl Sepharose 6 Fast Flow High Sub column (GE Healthcare). Supernatant samples were loaded onto the HIC column, using the following buffers: in the HIC-1, the Fh8 tag was purified by HIC using a Tris–NaCl buffer without calcium addition (50 mM Tris pH 7.6 and 150 mM NaCl). In the HIC-2, the Fh8 tag was purified by HIC using the Tris–NaCl buffer supplemented with 5 mM CaCl₂. For both purifications, the elution buffer was used in the same concentration as indicated in Table 1. In the HIC-3, the Fh8 tag was purified by HIC using the Tris–NaCl buffer supplemented with 5 mM CaCl₂ but with two different elution buffers: a first elution buffer with EDTA (50 mM Tris pH 7.6, 150 mM NaCl and 5 mM EDTA) and a second elution buffer identical to the described at Table 1 (50 mM Tris pH 10). Aliquots of all supernatant and flow-through samples, washing steps and eluted samples were stored at 4 °C to be further analyzed.
Protein purification by HIC and by IMAC

Cell pellets of HisSOD, Fh8SOD, HisGFP and Fh8GFP proteins (one pellet per protein, corresponding to 1 L E. coli culture) were thawed and resuspended in a total volume of 2 × 20 mL of lysis buffer (50 mM Tris pH 7.6 and 150 mM NaCl, supplemented with 1 × complete free EDTA protease inhibitor (Roche), 5 μg mL⁻¹ DNase (Sigma) and 1 mg mL⁻¹ lysozyme (Sigma)) with the addition of 20 mM imidazole, for IMAC purifications (1 × 20 mL), or 5 mM CaCl₂, for HIC purifications (1 × 20 mL). Cells were lysed as previously mentioned in the Fh8’s purification and aliquots of total lysates and supernatant samples were stored at 4 °C for Bradford estimation of the total protein content and for SDS–PAGE analysis. SOD and GFP target proteins were purified in parallel by HIC, using the same column as for the Fh8’s purification, and by IMAC, using a 5 mL HisTrap pre-packed column (GE Healthcare). All the proteins were purified following an identical strategy (see Supplementary file 3). The composition of specific buffers used for each purification methodology is described in Table 1. The obtained purified SOD and GFP fusion proteins were dialyzed in 50 mM Tris pH 7.6, 150 mM NaCl buffer, filtered through 0.2 μm and stored at 4 °C and −20 °C until used. Columns regeneration and storage was performed according to the manufacturer’s instructions.

Dual protein purification using HIC/IMAC and IMAC/HIC

Fh8GFP and Fh8SOD fusion proteins were purified by HIC followed by IMAC using the same above-mentioned protocols, with the following modifications: after HIC purification, eluted samples were dialyzed in 50 mM Tris pH 7.6, 150 mM NaCl buffer supplemented with 20 mM imidazole. The IMAC/HIC purification was only conducted for the Fh8SOD protein. Eluted samples from the IMAC purification were dialyzed in 50 mM Tris pH 7.6, 150 mM NaCl buffer supplemented with 5 mM CaCl₂, following an identical protocol as above-mentioned. At the end of the HIC/IMAC or IMAC/HIC purifications, samples were dialyzed in 50 mM Tris pH 7.6, 150 mM NaCl buffer and stored at 4 °C and −20 °C until needed. Aliquots of all purification samples were stored at 4 °C to be further analyzed.

SDS–PAGE of Fh8 and His fusion proteins was conducted according to the Laemmli method [24] using 12–4% gels. SDS–PAGE of the Fh8 tag samples was conducted according to the Schagger and Jagow method [25], using 15–4% gels. Gels were loaded with the PageRuler Unstained Broad Range Protein Ladder (Thermo Scientific). Protein purity and correspondent molecular weights in the SDS–PAGE gels were estimated by densitometry, conducting three independent readings in the Image Lab 2.0 software (Bio Rad), using the Molecular Imager Chemidoc XR+ system (Bio Rad).

The total protein content of supernatant samples and purification samples was estimated by Bradford method [26], using the Bio Rad protein assay dye reagent and bovine serum albumin as standard. Protein quantifications were also conducted by reading the absorbance of eluted samples at 280 nm. All protein quantifications were conducted in triplicates. Purification efficiencies were estimated by the ratio between the protein amount in eluted samples and the protein amount loaded onto the purification column.

GFP fluorescence measurements and SOD activity evaluation

The GFP target protein was considered to be active by emitting green fluorescence. The fluorescence intensity of GFP eluted samples was measured in triplicates using a spectrophotometer with an excitation filter of 475 nm and an emission filter of 505 nm. The resulting fluorescence intensities were normalized by the protein amount (in milligrams). The SOD activity was evaluated according to the method of Marklund and Marklund [27], in three independent assays, by estimating the inhibition of pyrogallol autoxidation promoted by the eluted samples, using a spectrophotometer at 420 nm.

Results

The Fh8 tag interaction with the hydrophobic resin

The Fh8 was expressed in E. coli harboring the pETM8h plasmid with an estimated molecular weight of 12 kDa (see Supplementary file 3). Results of the three Fh8’s HIC purification protocols (Fig. 1 and Table 2) demonstrated the calcium-dependent interaction of Fh8 with the hydrophobic resin, as follows: in the HIC purification without calcium supplementation in the binding step (HIC-1), most of the loaded Fh8 (gel band of 12 kDa) was observed in the flow-through (FT lane, gel band of 12 kDa) and washing (W lane, gel band of 12 kDa) samples, resulting in a purification efficiency of 0%. The eluted sample (using the pH 10, E lane) did not contain the 12-kDa Fh8’s gel band, being majorly composed by two gel bands, one of 23 and other of 54 kDa. In the HIC purifications with calcium addition (HIC-2 and HIC-3), the Fh8 was mostly visible in the eluted samples (E lanes, gel band of 12 kDa) and small leakages were observed in the washing steps. When performing the elution with 50 mM Tris pH 10 (HIC-2), it was possible to observe the Fh8’s recovery together with the other two gel bands of 23 and 54 kDa. On the other hand, when a first elution step with EDTA supplementation (HIC-3) was performed, only the 12 kDa gel band was observed (lane E1). The other high molecular weight gel bands eluted in 50 mM Tris pH 10 buffer (lane E2). The two elution strategies yielded similar purification efficiencies (82 ± 6.2% and 86 ± 4.3%, respectively).

These results showed that the Fh8 tag successfully interacted with the HIC resin by using buffers supplemented with calcium.

SOD and GFP purification by HIC and by IMAC

The Fh8 solubility enhancer tag was evaluated as a purification handle using the HIC in the presence of calcium (purification conditions of HIC-2). Fh8GFP and Fh8SOD recombinant fusion proteins were purified by HIC in parallel with IMAC, and the equivalent

Table 1

<table>
<thead>
<tr>
<th>Purification technique</th>
<th>Buffer Composition</th>
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<tbody>
<tr>
<td>IMAC</td>
<td>Buffer</td>
</tr>
<tr>
<td>Binding</td>
<td>50 mM Tris pH 7.6</td>
</tr>
<tr>
<td></td>
<td>150 mM NaCl</td>
</tr>
<tr>
<td></td>
<td>20 mM imidazole</td>
</tr>
<tr>
<td>Washing</td>
<td>50 mM Tris pH 7.6</td>
</tr>
<tr>
<td></td>
<td>150 mM NaCl</td>
</tr>
<tr>
<td></td>
<td>50 mM imidazole</td>
</tr>
<tr>
<td>Elution</td>
<td>50 mM Tris pH 7.6</td>
</tr>
<tr>
<td></td>
<td>150 mM NaCl</td>
</tr>
<tr>
<td></td>
<td>300 mM imidazole</td>
</tr>
<tr>
<td>HIC</td>
<td>Binding</td>
</tr>
<tr>
<td></td>
<td>150 mM NaCl</td>
</tr>
<tr>
<td></td>
<td>5 mM CaCl₂</td>
</tr>
<tr>
<td>Washing = binding 1:2</td>
<td>25 mM Tris pH 7.6</td>
</tr>
<tr>
<td></td>
<td>75 mM NaCl</td>
</tr>
<tr>
<td></td>
<td>2.5 mM CaCl₂</td>
</tr>
<tr>
<td>Elution</td>
<td>50 mM Tris pH 10</td>
</tr>
</tbody>
</table>
His-fused proteins were employed as a reference control for both HIC and IMAC purifications, being also used to evaluate the specific Fh8's hydrophobic interaction of the Fh8 fusion proteins.

The SDS–PAGE comparative analysis (Fig. 2) revealed an identical purification profile between the Fh8-HIC and IMAC methodologies for both Fh8GFP (gel band of 37 kDa, Fig. 2a) and Fh8SOD (gel band of 29 kDa, Fig. 2c) fusion proteins, with low protein amount in the flow-through (FT lanes) and washing (W lanes) samples, and with most of the protein in the elution samples (E lanes). The eluted samples from HIC and IMAC purifications, however, were not completely pure: the Fh8GFP fusion proteins presented a second gel band (at 22 kDa for HIC and at 28 kDa for IMAC), and the Fh8SOD fusion proteins presented other gel bands of different molecular weights.

The purification efficiencies of Fh8GFP and Fh8SOD were also similar between the Fh8-HIC and IMAC methodologies, as shown in Table 3.

Both HisGFP (gel band of 30 kDa, Fig. 2a) and HisSOD (gel band of 21 kDa, Fig. 2c) proteins presented an identical purification pattern: they were successfully purified by IMAC but not so well by HIC, in which they were mostly found in the washing samples (W lanes) rather than in the elution samples (E lanes). The HisGFP purification by IMAC yielded a similar efficiency as the Fh8GFP purifications by IMAC or Fh8-HIC (Table 3). In the HIC purification of the HisSOD protein, other gel band from the soluble extract of E. coli was observed nearby the HisSOD protein gel band. We could confirm that this gel band did not correspond to the HisSOD protein as it was not purified by IMAC (Fig. 2c, lanes FT and W) and it did not present SOD's biological activity. A protein loss of 96 ± 8% was estimated for the HisSOD purification by HIC, taking into account the total protein content and corresponding densitometric analysis in the flow-through and washing samples.

**Table 2**

<table>
<thead>
<tr>
<th>HIC-1</th>
<th>HIC-2</th>
<th>HIC-3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Binding: no calcium</td>
<td>Binding: +5 mM CaCl&lt;sub&gt;2&lt;/sub&gt;</td>
<td>Binding: +5 mM CaCl&lt;sub&gt;2&lt;/sub&gt;</td>
</tr>
<tr>
<td>Elution: Tris pH 10</td>
<td>Elution: Tris pH 10</td>
<td>Elution: TrisNaCl+EDTA (E1) → Tris pH 10</td>
</tr>
<tr>
<td>Loaded (mg)&lt;sup&gt;a&lt;/sup&gt;</td>
<td>37 ± 1.4</td>
<td>22 ± 0.7</td>
</tr>
<tr>
<td>Eluted (mg)&lt;sup&gt;b&lt;/sup&gt;</td>
<td>0 ± 0</td>
<td>18 ± 1.3</td>
</tr>
<tr>
<td>Purification efficiency (%)&lt;sup&gt;c&lt;/sup&gt;</td>
<td>0 ± 0</td>
<td>82 ± 6.2</td>
</tr>
</tbody>
</table>

<sup>a</sup> Values were determined by taking into account the SDS–PAGE densitometric analysis of each target protein and the total protein amount presented in the loading.

<sup>b</sup> Values were determined by taking into account the SDS–PAGE densitometric analysis of each target protein and the total protein amount presented in the elution step.

<sup>c</sup> Efficiency is the ratio between the target protein amount in the elution step and the initially loaded amount of each target protein.

**Dual purification system: HIC/IMAC and IMAC/HIC**

Depending on the protein's application, it is often necessary to perform two or more purification steps to increase its purity level. The purity of eluted Fh8GFP and Fh8SOD proteins from the Fh8-HIC purification was further evaluated by a second purification with IMAC. A second HIC purification was also conducted with the Fh8SOD protein after IMAC purification. As observed in Fig. 2b and d, the HIC/IMAC or IMAC/HIC purifications improved both Fh8GFP and Fh8SOD proteins purity, achieving purity levels between 85% and 92% of the target protein, evaluated by densitometric analysis of SDS–PAGE. Moreover, Fh8GFP and Fh8SOD were purified by these dual strategies with similar efficiencies as the single purifications (Table 3): the Fh8GFP yielded a purification efficiency of 98 ± 7.0% after HIC/IMAC, and the Fh8SOD achieved a purification efficiency of 70 ± 28% or 78 ± 4.4% after HIC/IMAC or IMAC/HIC, respectively.

**Functional assays of purified fusion proteins**

The Fh8-HIC purification methodology was further compared to the IMAC purification regarding the biological activity of the purified proteins (Fig. 3). In general, the Fh8GFP purified by IMAC resulted in higher relative fluorescence units (RFU) per mg of protein than the Fh8GFP purified by HIC. The Fh8 tag did not interfere with the GFP fluorescence, and the Fh8GFP protein was, thus, biologically active after being purified by the Fh8–HIC methodology. The Fh8GFP purified protein from HIC/IMAC presented also high RFU values. The several GFP target proteins did also exhibit green light under natural daylight. Photos of all purified GFP proteins are available in the Supplementary file 4.
The Fh8SOD fusion protein purified by the Fh8-HIC strategy presented similar biological activity as the HisSOD protein purified by IMAC, and higher biological activity than the Fh8SOD purified by IMAC. In good agreement with the results observed for the GFP activity, the Fh8 tag did not also affect the SOD activity as fusion protein. The eluted sample from HisSOD purification by HIC did not present SOD activity, resulting in similar values as the Fh8GFP negative control protein. Taking into account this low activity and the SDS–PAGE analysis of Fig. 2, we confirmed that this eluted protein did not correspond to the HisSOD.

The Fh8SOD fusion protein purified by the Fh8-HIC strategy presented similar biological activity as the HisSOD protein purified by IMAC, and higher biological activity than the Fh8SOD purified by IMAC. In good agreement with the results observed for the GFP activity, the Fh8 tag did not also affect the SOD activity as fusion protein. The eluted sample from HisSOD purification by HIC did not present SOD activity, resulting in similar values as the Fh8GFP negative control protein. Taking into account this low activity and the SDS–PAGE analysis of Fig. 2, we confirmed that this eluted protein did not correspond to the HisSOD.

**Discussion**

In this study, a novel methodology for protein purification using the Fh8 solubility enhancer tag [21] was investigated. The chromatographic properties of the Fh8 tag were firstly demonstrated through simple proof-of-principle experiments and the usefulness of the Fh8 as purification tag was then evaluated by its fusion to two different model proteins: GFP and SOD. The purification efficiency of Fh8-fused proteins by HIC was also compared to the His tag technology.

Taking into account the calcium binding property of Fh8 molecule [20] and its efficiency as solubility enhancer tag [21], the Fh8 was explored as a purification handle using a simple methodology: the hydrophobic interaction chromatography. Experiments using the Fh8 tag by itself demonstrated the calcium-dependent specificity of the binding mechanism and its affinity to the phenyl-Sepharose hydrophobic resin. In fact, the Fh8 could only bind the hydrophobic matrix in the presence of calcium in the mobile phase. Without calcium, the Fh8 tag presented low affinity for the matrix. The Fh8 tag was also able to bind to the phenyl-Sepharose resin.
under low salt concentration in the mobile phase, corroborating the results obtained for other calcium binding proteins [22,23]. The salt concentration in the mobile phase has significant contribution for the HIC performance: when using anti-chaotropic salts as sodium chloride, in a medium of high salt concentration, the bound form of the protein is thermodynamically more favorable than the unbound state. These salts bind water molecules strongly, excluding them from the protein and ligand surfaces, which start to interact hydrophobically (salting-out effect) [28–30]. Therefore, when using low salt concentrations in the mobile phase, the binding of proteins to the HIC matrix is not favored. Considering that the purification buffers used in this work have low salt concentrations, most of the contaminant proteins in the E. coli extract are excluded in the binding step, promoting selectivity towards the purification of the desired fusion protein. The Fh8 tag eluted from the phenyl-Sepharose column either with EDTA, or with pH 10. The later was selected to perform the purification of Fh8 fusion proteins as it allowed a single-step and rapid elution of all bound proteins. The use of a calcium chelating agent has already been demonstrated to be effective for the elution of other calcium binding proteins [17,22,23]. In fact, this elution mechanism proved to be highly selective towards Fh8 tag since other proteins of superior molecular weight (between 20–25 and 50–75 kDa) were only observed in the elution with pH 10, after the first elution with EDTA. Overall, the Fh8 tag interacts with the HIC resin as a calcium binding protein and it has the properties required for protein purification by HIC, offering the possibility to control the binding, selectivity and elution steps with the exclusion of major E. coli contaminants.

In order to investigate if the Fh8 tag could be used as a purification handle, namely, if chromatographic properties of Fh8 were preserved after fusion to target proteins, we selected two model target proteins with different characteristics, GFP and SOD. These proteins were fused to the Fh8 tag and to the His6 tag, and a Fh8-HIC purification protocol with mild conditions that did not interfere with target biological activity of target proteins was developed. Results from this work demonstrated that both Fh8GFP and Fh8SOD fusion proteins were soluble expressed in E. coli, and their purification efficiencies and biological activities after Fh8-HIC purification were comparable to those of Fh8-fused and His-fused proteins after IMAC purification. The higher biological activity observed for the Fh8SOD fusion protein purified by HIC or by HIC/IMAC may be explained by the calcium addition during the HIC purification protocol and its effect on SOD activity itself. Actually, the addition of calcium has a positive effect on SOD activity [31]. In order to corroborate this effect, we also compared the biological activity of Fh8SOD protein purified by IMAC with or without the addition of CaCl2 5 mM, and an increase in SOD activity per mg of protein was observed (data not shown).

The HIC purification of both GFP and SOD proteins was only possible when fused to Fh8 tag as shown by the weak interaction of the His6-tagged proteins with the HIC matrix. This unfavorable interaction of HisGFP and HisSOD to the HIC matrix can be
explained by the previously-mentioned low salt concentration used in the binding buffer. The difference between the pH value used in both binding and washing buffers (pH 7.6) and the pI values of target proteins (5.8 for HisGFP and 5.9 for HisSOD) is probably causing a net charge shield around these proteins that will also decrease the chance for hydrophobic interactions [30,32].

The use of dual tags for expression and purification of recombinant proteins has become an increasingly popular method that simplifies purification and yields homogeneous preparations of the protein of interest [33]. Our results showed that the dual Fh8-HIC and IMAC purification system is an added-value strategy that can be used sequentially, complementing each other, to obtain an active and more purified protein when desired. Moreover, the use of two consecutive purification steps and the distinct nature of HIC and IMAC methodologies allows for the efficient removal of contaminating proteins [17].

The data presented here have proven the feasibility of the Fh8-HIC purification strategy as a rapid, easy and cost-effective affinity methodology for protein recovery from E. coli extracts, even without an optimized purification protocol. Naturally, the Fh8-HIC methodology can be further optimized, as for instance, by testing other buffer solutions, hydrophobic matrices and EDTA/EGTA elution gradients, in order to use the Fh8 purification tag in a broader range of conditions.

The novel Fh8 purification tag presents attractive features when compared to other commonly affinity fusion technologies (see Table 4), being its efficient dual solubility enhancer and purification handle functionality the most relevant one. In fact, the Fh8 tag is one of the few existing fusion tags to combine both protein solubility enhancer and purification effects, along with the MBP and GST tags (Table 4). The MBP tag is a well-known solubility enhancer partner, but it presents some problems in protein interaction with the resin when used as a purification handle [34]. The GST tag is often described as a poor solubility enhancer [21,35]. Moreover, both fusion tags are large-sized tags, and require specific and expensive resins for protein purification. The Fh8 tag differs from MBP and GST traditional purification tags as it makes use of intrinsic calcium-binding properties for an economical hydrophobic interaction chromatography under mild conditions, instead of expensive resins, harsh buffers and additional compounds and ligands. The Fh8 tag goes forward the existing fusion tags by accumulating an efficient solubility enhancer activity and simple and cost-effective purification in a low molecular weight peptide.

The Fh8-HIC purification strategy is calcium-dependent and, consequently, chelating agents must be avoided during protein binding and washing steps. This could be considered a limitation of the process but it is important to mention that this limitation can also be found in the IMAC technology. Due to the calcium-dependent mechanism for protein purification using the Fh8 tag, this Fh8-HIC strategy can be potentially applied for several different target proteins without requiring the development and optimization of a new system for each target protein.

**Table 4**

Comparison of Fh8 fusion partner to other commonly used affinity partners.

<table>
<thead>
<tr>
<th>Tag</th>
<th>Size (aa)</th>
<th>Function</th>
<th>Purification protocol</th>
<th>Advantages</th>
<th>Disadvantages</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fh8 ([Costa et al. [21], and this work)]</td>
<td>69 PH</td>
<td>Calcium dependent hydrophobic interaction; selective elution with EDTA or pH manipulation</td>
<td>Efficiently enhances protein solubility in E. coli; small-sized tag; simple and cost-effective purification using gentle and mild conditions</td>
<td>Large tag; expensive process; problems in protein interaction with the resin Poor solubility enhancer; large tag; expensive process; problems in protein interaction with the resin Purification efficiency can be reduced; expensive process Requires detergents to minimize adsorption of host proteins to the particles; high salt concentration for elution; large size Expensive process; difficult elution*</td>
<td></td>
</tr>
<tr>
<td>MBP ([Di Guan et al. [7])</td>
<td>396 SE PH</td>
<td>Affinity towards amylose; elution with maltose</td>
<td>Simple purification process</td>
<td></td>
<td></td>
</tr>
<tr>
<td>GST ([Smith [6])</td>
<td>211 SE PH</td>
<td>Affinity towards glutathione, elution with free glutathione</td>
<td>Simple purification process</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Histidine ([Hochuli et al. [8])</td>
<td>6–10 PH</td>
<td>Affinity towards Ni²⁺; elution with imidazole</td>
<td>Small-sized tag; simple purification process</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Si-tag ([Ikeda et al. [10])</td>
<td>273 PH</td>
<td>Affinity towards silica particles; elution with high concentration of MgCl₂ or CaCl₂</td>
<td>Inexpensive commercial silica particles; simple purification process</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tamavidin ([Takakura et al. [11])</td>
<td>140 PH</td>
<td>Affinity towards biotinylated magnetic microbeads</td>
<td>High affinity protein immobilization</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Tab2 tag ([Crusius et al. [12])</td>
<td>7 PH</td>
<td>Antibody-mediated protein identification and purification; elution at acidic pH (2.3)</td>
<td>Specificity of binding</td>
<td>Expensive process; harz conditions for elution</td>
<td></td>
</tr>
<tr>
<td>Z-basic ([Hedhammar and Hober [15])</td>
<td>58 PH SE</td>
<td>Cation-exchange chromatography</td>
<td>Simple and cost-effective purification process; suitable for processes under denaturing conditions</td>
<td>Poor solubility enhancer</td>
<td></td>
</tr>
<tr>
<td>Dock tag ([Kamezaki et al. [16])</td>
<td>22 PH</td>
<td>Dockeer-cohesin calcium dependent affinity</td>
<td>Highly selective</td>
<td>Expensive process</td>
<td></td>
</tr>
<tr>
<td>HiCaM ([McCuskey et al. [17])</td>
<td>162 PH</td>
<td>Tandem IMAC-HIC purification</td>
<td>High purity level</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Elution was carried by heating beads at 99 °C.

**Conclusion**

The Fh8 solubility tag was successfully established in this work as an efficient purification handle, thus becoming one of the few existent fusion partners to combine an effective solubility enhancer activity and protein isolation in a single tag. Moreover, the Fh8 tag offers several benefits over other solubility and purification tags: a low molecular weight tag that may not disturb the biological activity of target proteins (as demonstrated in this study with two model proteins), a highly soluble and easy target protein production in E. coli, besides simplicity and economy of the entire protein’s production process. Namely, it promotes target protein solubility directly into the E. coli cytoplasm and it does not require specialized buffers and substrates for protein purification, also making use of inexpensive and high-capacity matrices.

We foresee the novel Fh8 tag as a robust fusion partner of most utility for the rapid and cost-effective large scale soluble production and purification of several proteins. When desired, the Fh8 tag can also be used in a two-step purification procedure together...
with IMAC methodology, as well as other purification strategy, to further improve the protein purity level.

Conflicts of interest

The Fh8 tag utilization for the improvement of protein soluble expression in E. coli is covered by a worldwide patent (WO 2010082097) licensed to Hitag Biotechnology, Lda. The authors S.C., A.A. and A.C. are co-owners of the patent and are associated with Hitag Biotechnology, Lda.

Contributors

SC and EC participated in the study design and process development, carried out the experimental work and drafted the manuscript. LF participated in the experimental work and AA participated in results discussion and helped to draft the manuscript. LD and AC conceived and supervised the study, participated in its coordination and helped to draft the manuscript. All authors read and approved the final manuscript.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.pep.2013.09.013.

References