A successful strategy for the recovering of active P21, an insoluble recombinant protein of Trypanosoma cruzi

Marlus Alves dos Santos1,*, Francesco Brugnera Teixeira2,*, Heline Hellen Teixeira Moreira2, Adele Aud Rodrigues1, Fabricio Castro Machado1, Tatiana Mordente Clemente1, Paula Cristina Brigido1, Rebecca Tavares e. Silva1, Cecilio Purcino1, Rafael Goncalves Barbosa Gomes1, Diana Bahia1,4, Renato Arruda Mortara3, Claudia Elisabeth Munte2, Eduardo Horjales2 & Claudio Vieira da Silva1

1Instituto de Ciências Biomédicas, Universidade Federal de Uberlândia, Uberlândia, MG, Brasil, 2Instituto de Física de São Carlos, Universidade de São Paulo, São Carlos, SP, Brasil, 3Departamento de Microbiologia, Imunologia e Parasitologia, Escola Paulista de Medicina, Universidade Federal de São Paulo - EPM/UNIFESP, SP, Brasil, 4Departamento de Biologia Geral, Instituto de Ciências Biológicas, Universidade Federal de Minas Gerais-ICB/UFMG, Minas Gerais, Brasil.

Correspondence and requests for materials should be addressed to C.E.M. (claudia.munte@ifsc.usp.br); E.H. (horjales@ifsc.usp.br) or C.V.D.S. (silva_cv@yahoo.com.br)

* These authors contributed equally to this work.

Structural studies of proteins normally require large quantities of pure material that can only be obtained through heterologous expression systems and recombinant technique. In these procedures, large amounts of expressed protein are often found in the insoluble fraction, making protein purification from the soluble fraction inefficient, laborious, and costly. Usually, protein refolding is avoided due to a lack of experimental assays that can validate correct folding and that can compare the conformational population to that of the soluble fraction. Herein, we propose a validation method using simple and rapid 1D 1H nuclear magnetic resonance (NMR) spectra that can efficiently compare protein samples, including individual information of the environment of each proton in the structure.

Heterologous expression of proteins in Escherichia coli is the most widespread technique used to produce large amounts of protein for structural studies. However, a sufficient quantity of protein is not always obtained using common, native, non-denaturant protocols. Major problems associated with rapid and high-level expression in bacteria include protein aggregation and formation of inclusion bodies1. In some structural genomics projects2,3, approximately 40% of the expressed targets are found in the insoluble fraction. Although these proteins are usually discarded as viable targets, the use of a simple refolding protocol can resolve the problem. Often, the fraction of soluble protein is not sufficient to make the production of large amounts of protein viable, and, in these cases, use of a refolding protocol can be the best solution to increase protein yield. Several techniques have been applied to refold proteins4–6, but the problem of how to solubilize and correctly fold proteins remains a major issue. One of the main difficulties in using refolded proteins is that there are very few experiments that discriminate between different conformational states and guarantee the correct refolding of the protein. Enzymatic7–10 and biological activity11–13 have been widely used to confirm correct protein refolding. However, in general, significant statistical error is associated with these assays, and the structural integrity of the refolded sample cannot be ensured. With respect to direct analysis of protein structure, circular dichroism (CD) is an effective technique only when large changes in secondary structure are present.

High resolution structural studies can be hindered by low yields of expressed protein or lack of an optimized production/purification protocol. For example, obtaining a 3D nuclear magnetic resonance (NMR) spectrum of a 20 kDa protein typically requires 5–10 mg of very pure and high cost isotopically-labeled protein. In x-ray crystallography, large amounts of very concentrated protein are required to test crystallization conditions. In addition, protein purity alone is not always sufficient to ensure successful crystal growth. Even small changes in a protein’s three-dimensional structure can disturb crystal formation14–16. If an alternative conformation or oligomer attaches to the crystal, it may form a crystal surface that prevents attachment of other protein molecules, poisoning its growth17,18. Proteins can exist as multiple conformations in equilibrium due to many factors, such as structural flexibility, enzymatic activity, cooperativity, and oligomerization. For some experiments, conformational purity may not be important. For example, a biological activity assay of a protein with 20% in an inactive conformation is equivalent to another experiment with 20% less protein, but with 100% in an active conforma-
The yield of P21-His₆ was greatly improved when the protein was recovered from inclusion bodies and refolded. In order to compare the yield of soluble to refolded protein, we expressed recombinant P21-His₆ in 8 L 2xYT medium. Expression produced 24.9 g of dry weight bacteria and 8.1 g of insoluble pellet after lysis and centrifugation. Most of the protein was present in the insoluble fraction. sP21-His₆ was purified from the soluble fraction in a single step of Nickel affinity chromatography; however, only 0.68 mg of purified protein was obtained. The refolding procedure recovered 57.76 mg of rP21-His₆ from inclusion bodies, a yield 85 times greater than purification from the soluble fraction.

Results from invasion and phagocytosis experiments demonstrated that soluble and refolded P21-His₆ exhibit similar biological properties. In the presence of sP21-His₆, invasion of Vero cells by T. cruzi metacyclic trypomastigotes (TCT) (Figure 1A) was increased approximately 3.6-fold, and invasion by EA (Figure 1B) was increased approximately 2.4-fold. Similar results were obtained for rP21-His₆ (4.4-fold increase for TCT and 2.3-fold increase for EA). The difference between sP21-His₆ and rP21-His₆ was not significant. Phagocytosis of zymosan particles by macrophages was increased 3.5-fold with the addition of 40 μg/ml sP21-His₆ (Figure 1C) and 3.5-fold with the addition of 40 μg/ml rP21-His₆. The similarity of these increases suggests that refolded and soluble protein exhibit the same biological properties.

Finally, CD and NMR experiments validated the correct refolding of P21-His₆. The CD spectra showed equivalent secondary structure composition for both preparations (Figure 2B). Comparison between ¹H NMR spectra of sP21-His₆ and rP21-His₆ showed exactly the same chemical shifts and intensities for every proton in the structure (Figure 2C). In this way, both samples could be compared directly and were found to be identical to one another, with respect to both correct folding and conformational ensemble. To better evaluate the similarities (or differences) between the spectra of both samples, a difference spectrum was calculated and is shown in Figure 2C (green line). The larger differences can be assigned to non-protein components of the sample (residual ethanol from concentrator rinsing, small differences in DSS concentration, low molecular weight contaminants). These differences underscore the sensitivity of the method and the identity of both protein conformations and ensembles. In addition, up field shifts around 0 ppm in both spectra indicate a hydrophobic nucleus in P21-His₆, showing that the structure is folded.

Discussion
In this study, we applied a one-step refolding procedure to P21-His₆, which increased the yield of the protein 85-fold. This refolding method is based on immobilization of the P21-His₆ on Nickel-affinity resin, preventing protein aggregation during the process. Refolding requires approximately 14 hours (which can be performed overnight with no need for human interference) and uses less than 0.5 L of buffer. It is less laborious and quicker than other refolding techniques, such as dialysis.

We used pre-existing biological information about P21 to develop protein-specific assays that could be applied to both refolded and soluble P21-His₆ fractions. Both samples increased parasite invasion and phagocytosis of zymosan particles at similar rates, suggesting that they behaved identically with respect to biological activity. This information would be sufficient in a majority of biological studies, but is not sufficient for structural studies, in which correct folding and conformational homogeneity of samples are essential. As more structure-focused approaches, CD and NMR spectra provided structural comparisons between sP21-His₆ and rP21-His₆. Although CD measurements provide information about protein secondary structure, no information at an atomic resolution can be provided. This method is more useful for identifying incorrectly folded proteins by detecting differences in the spectra, because even if two spectra are identical, we cannot confirm that structures are equal. Simple one-dimensional ¹H NMR overcomes this limitation by providing information about the environment of each ¹H atom of the protein, allowing a comparison of samples obtained by different purification methods at atomic resolution. With this simple ¹D ¹H NMR measurement, we confirmed that sP21-His₆ and rP21-His₆ have exactly the same structures and identical conformational equilibria. In addition, more important information can be obtained from the spectrum: the good dispersion of the peaks over the ppm scale indicates the presence of secondary structure, and the existence of high-field shifted peaks around the reference (DSS at 0 ppm) indicates the presence of a hydrophobic core in the protein. Therefore, the structural and the biological data were in agreement, as expected.
We propose the use of simple and fast 1D ¹H NMR spectra to validate refolding protocols before increasing protein production. This technique is able to detect very small conformational changes in protein structure, as ¹H NMR is sensitive to changes in the environment of each hydrogen atom of a molecule. By searching the REFOLD database, we found that 34 of 1001 refolded proteins used this NMR spectroscopy in some way. In the majority of cases, the studies were driven by NMR, and the refolded protein was produced and isotopically labeled for analysis. Only three proteins from the database used simple 1D ¹H NMR spectra, two of them for visual inspection, and one for comparison with the natural modified protein. In a recent study, the authors compared the 1D ¹H NMR spectra from the refolded protein and the protein purified from soluble fraction. They concluded that the structures were exactly the same because the spectra were identical (no spectral difference was presented). In a recent review focused on tools used to evaluate protein conformations, NMR is briefly mentioned as one of the leading techniques of structural biology. The authors said that NMR "still requires very sophisticated equipment, appreciable amounts of sample, and specialist human resources that keep them away from the pipeline in protein production and control. Newer techniques (or new applications of well-established techniques) are being developed to complement these traditional biophysical assays and to provide information on the success of protein folding." We want to stress that the proposed method uses the detailed information present in NMR spectra in a simple way, to answer a simple question. If NMR (as x-ray crystallography) is analyzed as a tool to determine three-dimensional protein structure, then it presents the difficulties pointed out in the aforementioned review; however, this is not the case for the 1D ¹H spectra needed for conformational comparisons of two samples.

**Conclusion**

One-dimensional homonuclear ¹H NMR spectroscopy requires no special sample production and little time for measurement. Typically, a high quality spectrum of a 500 μL protein sample (~100 μM of a small protein) on a 600 MHz spectrometer can be obtained in less than 30 minutes. Most of the time is spent on adjustment of the equipment and definition of experimental parameters. Using spectrometers with a cryoprobe is desirable, but not required. Spectra processing and comparison is very quick and can be performed immediately after acquisition using spectrometer software.

Finally, simple precautions should be taken for the preparation of NMR samples. The NMR buffer should be of neutral to acidic pH and low ionic strength. High pH results in less protonated amides, causing loss of signal. Low salt concentration (up to 300 mM) is required to minimize radio wave reflections during NMR pulses that can cause heating of the sample (tuning and matching of the spectrometer probe head). ¹H NMR is very sensitive to non-water exchangeable hydrogens of small molecules, and, as a consequence, the use of hydrogen-containing buffer agents (like Tris, Acetate, MES, and HEPES) and some other molecules (glycerin, DTT, etc.) should be avoided, or these reagents can be replaced by their deuterated equivalents. Sodium or potassium phosphate buffer is recommended, since the hydrogen atoms of these compounds are exchangeable with water, resulting in a signal that is coincident with the water resonance.
that should be suppressed. Moreover, extensive care with contaminants should be taken, as 1H NMR can detect contaminant signals, even in small concentrations.

**Methods**

**Protein expression and cell lysis.** P21-His was expressed in Escherichia coli BL21 cells transformed with a pET 28a (+) (Novagen) expression vector, which contains an N-terminal His6-tag followed by a thrombin recognition site. Expression was carried out in 8 L of 2xYT medium (16 g tryptone, 10 g yeast extract, 5 g NaCl per 1 L medium) containing 35 µg/mL kanamycin. Bacteria were incubated at 37°C until the optical density at 600 nm (OD600) reached ~0.7, at which point expression was induced by addition of 0.5 mM isopropyl-β-D-1-thiogalactopyranoside (IPTG). Cultures were then incubated for an additional 3 hours at 37°C. Cells were harvested by centrifugation at 14,000 g for 40 minutes at 4°C, and the pellet was resuspended in lysis buffer (50 mM Tris, 300 mM NaCl, 20 mM imidazole, pH 8.0) containing 1 mg/mL lysozyme. The solution was maintained on ice for 30 min before cells were lysed by ultrasound (15 cycles of 40 s at 40 W, 40 s interval). The lysate solution was subjected to centrifugation at 6,000 g for 40 minutes at 4°C and resuspended in lysis buffer (50 mM Tris, 300 mM NaCl, 20 mM imidazole, pH 8.0) containing 1 mg/mL lysozyme. The solution was maintained on ice for 30 min before cells were lysed by ultrasound (15 cycles of 40 s at 40 W, 40 s interval). The lysate solution was subjected to centrifugation at 14,000 g for 40 minutes at 4°C, and both supernatant and pellet were used for native and denaturing purification of the protein, respectively.

**Native P21-His purification from the soluble fraction (sP21-His6).** The soluble fraction was added to 2 mL of Ni-NTA resin (Qiagen) and extensively washed with lysis buffer. To remove low-affinity bound proteins, a second wash with wash buffer (50 mM Tris, 300 mM NaCl, 50 mM imidazole, pH 8.0) was performed. sP21-His6 was eluted with elution buffer I (50 mM Tris, 300 mM NaCl, 50 mM imidazole, pH 8.0) was added to 2 mL of Ni-NTA resin (Qiagen) and extensively washed with lysis buffer. To remove low-affinity bound proteins, a second wash with wash buffer (50 mM Tris, 300 mM NaCl, 50 mM imidazole, pH 8.0) was performed. sP21-His6 was eluted with elution buffer I (50 mM Tris, 300 mM NaCl, 50 mM imidazole, pH 8.0) containing 1 mg/mL lysozyme. The solution was maintained on ice for 30 min before cells were lysed by ultrasound (15 cycles of 40 s at 40 W, 40 s interval). The lysate solution was subjected to centrifugation at 6,000 g for 40 minutes at 4°C and resuspended in lysis buffer (50 mM Tris, 300 mM NaCl, 20 mM imidazole, pH 8.0) containing 1 mg/mL lysozyme. The solution was maintained on ice for 30 min before cells were lysed by ultrasound (15 cycles of 40 s at 40 W, 40 s interval). The lysate solution was subjected to centrifugation at 14,000 g for 40 minutes at 4°C, and both supernatant and pellet were used for native and denaturing purification of the protein, respectively.

**Denaturing P21-His6 purification and refolding from the insoluble fraction (rP21-His6).** In order to extract the expressed protein from inclusion bodies, the lysate pellet was resuspended in urea buffer (50 mM Tris, 300 mM NaCl, 5 mM imidazole, 6 M urea, 5% (v/v) glycerol, pH 8.0) and stirred overnight at room temperature. After centrifugation (14,000 × g for 40 min at 4°C), the supernatant, containing P21-His6 extracted from the inclusion bodies, was immobilized in a column containing 5 mL Ni-NTA resin fixed on an Akta Purifier system. After extensive washing with urea buffer, the urea concentration was slowly decreased with a linear buffer gradient (~7 mM/min of urea, 0.5 mL/min)3. To remove glycerol and additional impurities, the column was washed with wash buffer until no protein signal was detected. rP21-His6 was eluted with elution buffer II (50 mM Tris, 300 mM NaCl, 220 mM imidazole, pH 8.0), concentrated, and dialyzed against buffer II (see above).

**Cell lines and cultures.** Vero cells were cultured in Dulbecco’s Modified Eagle Medium (DMEM) supplemented with 10% fetal bovine serum (FBS, Cultilab), 10 mg/mL streptomycin (Sigma), 100 U/mL penicillin (Sigma), and 40 mg/mL gentamycin (Sigma) at 37°C in a humidified atmosphere containing 5% CO2. Peritoneal macrophages from BALB/C mice were harvested from the peritoneal cavity in 5 mL DMEM. Mice were stimulated with 1 mL 3% thioglycollate medium 3 days prior to harvest.

**Parasites.** To generate tissue-cultured trypomastigotes (TCT) from the CL strain, confluent monolayers of Vero cells were infected with metacyclic trypomastigotes and maintained for at least two weeks to establish the intracellular cycle. Extracellular amastigotes (EA) were derived from trypomastigotes by axenic differentiation. Infected Vero cell cultures were subjected to centrifugation (2500 × g, 5 minutes), and the pellet was resuspended in liver infusion tryptose (LIT) medium. Finally, the resuspension was incubated for 18 hours at 37°C to generate at least 95% pure extracellular amastigotes.

**Animals and ethics.** Male or female BALB/c mice were six to eight weeks old and were maintained under standard conditions on a 12 hour light, 12 hour dark cycle in a temperature controlled room (25 ± 2°C) with food and water ad libitum. All experimental protocols were approved by the Ethics Committee from Universidade Federal de Uberlândia. Moreover, maintenance and care of these animals complied with the guidelines of the Laboratory Animal Ethics Committee from the Institution. Animal euthanasia was performed in accordance with international welfare grounds,
according to the American Veterinary Medical Association Guidelines on Euthanasia.

Host cell invasion assay. A volume of 500 μL vero cell suspension (1.5 × 10^5 cells) was added to multiwell plates containing sterile glass coverslips and incubated overnight. The invasion assay was performed by adding EA (20 parasites per cell) or TCT (10 parasites per cell) to plates in the presence or absence of 40 μg/mL sP21-His6 or 40 μg/mL rP21-His6. The plates were incubated for 2 hours at 37°C in a humidified incubator containing 5% CO2. After incubation, the cells were gently washed three times with PBS, fixed with Bouin’s solution, and stained with Giemsa.

Phagocytosis assay. Zymosan particles (10 mg/mL, Sigma) were resuspended in sterile PBS, sonicated for 15 minutes, and centrifuged. To determine particle concentration, a Neubauer chamber was used for counting. Resident peritoneal macrophages were seeded in cell culture wells (5 × 10^5 cells/well) containing 13 mm round glass coverslips and were then incubated overnight in a humidified atmosphere containing 5% CO2 at 37°C to allow the cells to adhere. Non-adherent cells were removed by washing with PBS before adding DMEM containing 10% FBS. To perform the phagocytosis assay, zymosan particles were added to wells (20 particles/each macrophage fixed) in the presence or absence of 40 μg/mL sP21-His6. Particles were incubated with macrophages for 2 hours. The cells were then washed with PBS, fixed in Bouin’s solution, and stained with Giemsa. Results were expressed as the number of internalized particles/100 cells. These results were compared to those previously determined for rP21-His6, using the same protocol.

Circular dichroism spectropolarimetry (CD). Spectra were recorded on a Jasco 815 CD spectrometer using 0.1 cm thick quartz cells. Samples containing 0.15 mg/mL sP21-His6 or rP21-His6, in 10 mM sodium phosphate buffer (pH 7.4) were analyzed at 20°C to assess structural similarity.

Nuclear magnetic resonance spectroscopy (NMR). Samples of sP21-His6 and rP21-His6 were concentrated to the same concentration (1.3 mg/mL, 70 μM) in a final volume of 500 μL and then simultaneously dialyzed against 2 L NMR buffer (50 mM potassium phosphate, 150 mM KCl, pH 6.0) at 4°C. The NMR buffer was changed twice to ensure that the samples were in exactly the same buffer conditions for NMR measurements, as “H signals are very sensitive to changes in their environment. DSS (100 μM, for spectrum referencing) and D2O (5% v/v), for tuning and matching the spectrometer probe head) were added to both samples. 1D 1H spectra were recorded at 303 K in a high-resolution, small bore 600 MHz Bruker Avance III spectrometer. Since the spectrometer was equipped with a TCI cryoprobe, a 4× enhancement in the signal intensity was achieved. The intensive water signal around 4.8 ppm, typical for an aqueous protein sample, was suppressed using a standard watergate W5 pulse sequence. To ensure spectra with a good signal-to-noise ratio were obtained for the diluted samples, each 1D experiment was acquired with a total of 256 scans, resulting in an experimental time of approximately 17 minutes per spectrum. Proton chemical shifts were referenced (0 ppm) to the “H resonance frequency of the sharp and intense signal of the methyl groups in DSS.

Acknowledgments
This study was supported by grants and fellowships from FAPEMIG, CAPES, FAPESP (grants 2010/51867-6, 2012/21153-7), CNPq and INBEQMeDI. CVS was supported by grants from Fundação de Amparo à Pesquisa do Estado de Minas Gerais (FAPESP - Processos numbers APQ-00621-11, APQ-00305-12); Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (CAPES) (Process number 23038.005295/2011-40).

Author contributions
M.A.S. performed re-folding purification, the biological experiments, acquired, analyzed and interpreted the data and drafted the manuscript; F.B.T. performed in-column re-folding purification and NMR experiments, acquired, analyzed and interpreted the data and drafted the manuscript; H.H.T.M. performed solution fraction purification, CD and biological experiments, acquired, analyzed and interpreted the data and drafted the manuscript; A.A.R., F.C.M., T.M.C., P.C.B., R.T.S., C.P. & R.G.B.G. performed biological experiments; D.B. & R.A.M. critically revised the manuscript and contributed to its intellectual content; E.H. designed soluble fraction purification and CD experiments, drafted the manuscript and critically revised the manuscript and contributed to its intellectual content; A.A.R., F.C.M., T.M.C., P.C.B., R.T.S., C.P. & R.G.B.G. performed biological experiments; D.B. & R.A.M. critically revised the manuscript and contributed to its intellectual content; E.H. designed soluble fraction purification and CD experiments, drafted the manuscript and critically revised the manuscript and contributed to its intellectual content. All the authors approved the final version to be submitted.
Additional information

Competing financial interests: The authors declare no competing financial interests.

How to cite this article: Santos, M.A.d. et al. A successful strategy for the recovering of active P21, an insoluble recombinant protein of Trypanosoma cruzi. Sci. Rep. 4, 4259; DOI:10.1038/srep04259 (2014).

This work is licensed under a Creative Commons Attribution-NonCommercial-NoDerivs 3.0 Unported license. To view a copy of this license, visit http://creativecommons.org/licenses/by-nc-nd/3.0