Neutron scattering: a tool to detect \textit{in vivo} thermal stress effects at the molecular dynamics level in micro-organisms

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\textit{In vivo} molecular dynamics in \textit{Halobacterium salinarum} cells under stress conditions was measured by neutron scattering experiments coupled with microbiological characterization. Molecular dynamics alterations were detected with respect to unstressed cells, reflecting a softening of protein structures consistent with denaturation. The experiments indicated that the neutron scattering method provides a promising tool to study molecular dynamics modifications in the proteome of living cells induced by factors altering protein folds.

1. Introduction

Many studies have addressed the effects of physico-chemical stresses on proteins \textit{in vitro}. They revealed that protein molecular dynamics is much more sensitive to environment than protein structure \cite{1}. A protein can be inactive, for example, in an environment in which its structure is perfectly stable, because of inappropriate dynamics \cite{2}. By the very nature of the forces stabilizing tertiary structure, protein dynamics cannot be considered independently of solvent conditions. Thus, the crowded conditions as well as the presence of extensive molecular interactions that prevail in the cytosol of a living cell are believed to interfere significantly with protein folding and unfolding processes \cite{3}. It follows that the consequences of environmental stress on dynamic state of a protein in the cytosol could be very different from that measured \textit{in vitro} on a homogeneous population of purified protein, under dilute conditions. However, while many studies have addressed the effects of physico-chemical stresses and especially dehydration on protein molecular dynamics \textit{in vitro} \cite{4}, few direct biophysical observations have been performed on whole cells \textit{in vivo} and it is still difficult to appreciate the consequences of environmental stress on the molecular dynamic state of the whole proteome.

Incoherent neutron scattering is a method of choice to probe protein dynamics \cite{5}. Neutrons are scattered by atomic nuclei providing information on the momentum and energy of their motions. The probed length and time scales are defined by the scattering vector (Q) range and energy resolution of the experiment, respectively. By suitably choosing these values, it has been possible to observe internal dynamics of a protein population, \textit{in vivo}, without the requirement for specific protein labelling \cite{6}. Experiments are based on ‘elastic temperature scans’, in which neutrons scattered in a very narrow energy window around the energy of the incident beam are observed as a function of sample temperature. The results of the experiment are atomic mean square displacements (MSD, in Å\textsuperscript{2} units) and an averaged effective force constant (\(k_l\) in Nm\textsuperscript{−1} units), on the length and time scales selected by the experimental

conditions (typically approximately ångström amplitude in picoseconds to nanosecond time scale). These parameters inform on the mean flexibility and resilience of the protein structures, respectively [5]. The resilience has to be considered with respect to the free energy landscape, i.e. it includes both structural rigidity (the enthalpic term owing to internal forces) and conformational sampling (the entropic term). The neutron length–time window is well suited to examine the internal motions in unfolded polypeptides compared with those in folded proteins. Incoherent neutron scattering was used to probe the mean molecular dynamics properties of living cells [6–8]. Because of their relative abundance the macromolecular signal is dominated by the contribution of the proteins. Until now, the approach has been used in the case of adaptation of micro-organisms to temperature. It was found that evolution modifies in depth the molecular dynamics in psychrophile and thermophile microbes, with resilience values being a factor of two smaller or larger, respectively, than values found in mesophiles [7]. In these experiments, we noted a small difference that appeared when *Escherichia coli* cells were exposed to a heat stress. In this work, *Halobacterium salinarum* (*Hs*) was used as a model micro-organism to study, *in vivo* and in greater detail, the effects of temperature stress on molecular dynamics. *Hs* is not a thermophile, but it can tolerate heat stress better than most mesophilic cells [9]. Because of this, it was possible to explore a wider range of stress conditions than in *E. coli* (from 37°C to 65°C). This allowed the identification of a new Q range better suited to assess how a physical stress impacts the mean molecular dynamics properties of a microbial proteome within the cellular context, *in vivo*.

2. Results and discussion

2.1. Thermal stress response of *Halobacterium salinarum* in neutron experimental conditions

Microbiology experiments were performed in order to determine whether neutron experiment could be performed on stressed *Hs* R1 cells (*Hs*). The cells were cultivated at 37°C as described [10] and pelleted at mid-log phase. *Halobacterium* cells can maintain growth up to 50–55°C and stop dividing only at 60°C. The cells were exposed to 55°C and 60°C for 1 h, washed with a hypersaline physiological buffer and sealed in appropriate sample holders for 24 h under anaerobic conditions. The mortality rates were determined before and after heat shock, and after the neutron experiments by using light microscopy counts and live–dead staining protocols. Morphological analyses were also performed by confocal microscopy. In stressed and unstressed samples, the residual cell viability values were found to be over 95 per cent after 24 h of neutron measurements and no morphological changes could be detected (figure 1). *Hs* cells concentrate multi-molar amounts of K⁺ inside its cytosol to counterbalance the difference in osmotic pressure due to the external hypersaline conditions [11]. The process is driven by the membrane potential and is energy-dependent [12]. The biochemical activities and cellular functions are therefore adapted to function optimally in hypersaline conditions [13]. Thus, the measurement of intracellular K⁺ and Na⁺ concentration represent a good indicator of *Hs* cellular integrity. Inductively coupled plasma atomic emission spectroscopy (ICP) measurements performed on *Hs* cells showed that the high cytosol ion content values (3.7 M K⁺ and 1.1 M Na⁺) remained unchanged after 24 h of incubation in the sample holder, even after 1 h of a 60°C heat shock. We also tested the accumulation of the thermosome, a thermal stress protein, within the *Hs* cytosol in the different stress conditions, before and after neutron experiments. The thermosome is a type II chaperonin complex that represents the main protein quality control system in Archaea [14]. The thermosome acts on partially unfolded protein to promote their correct refolding in an energy-dependent manner [15]. Its abundance within *Hs* cells is associated with the accumulation of misfolded proteins substrates in the cytosol. The level of thermosome in the cells was immunodetected in *Hs* cell extracts (figure 2). These experiments showed that, for unstressed cells, the thermosome level remains constant after 24 h of neutron experiment. On the contrary, a significant increase in thermosome protein was observed in heat-shocked cells, before and after neutron experiments. These experiments showed that, in *Hs*, significant neutron measurements could be performed on intact cells, under stress conditions that generate the accumulation of misfolded proteins in the cytosol.

2.2. Neutrons detect protein denaturation *in vivo*

The energy and momentum changes measured in a neutron spectroscopy experiment are related to the time-scale and amplitude of atomic motions, respectively. In order to assess whether the effects of thermal stress on the *in vivo* molecular dynamics state could be detected, we performed neutron scattering experiments on unstressed and stressed *Hs* cells by using the IN13 spectrometer at the Institut Laue Langevin, as described in §3. In a complex system, different motion populations with different MSDs are observed, depending on the length and time scales. Because of this polydispersity, the most reliable analysis is *via* comparison of observations on the same system under different conditions on the same time and length scales. The time scale sampled on IN13 is approximately 0.1 ns, defined by the energy resolution of the instrument (8 μeV). The length scale is defined by the Q (scattering vector or momentum change modulus) range, which is quite broad on IN13: from below 0.3 Å⁻¹ to above 6 Å⁻¹, giving effective access to a length scale extending from a few ångströms at low Q to a fraction of an ångström at the high Q end. The most efficient and arguably most useful neutron spectroscopy method to measure dynamics is by analysis of the temperature-dependent elastic incoherent neutron scattering (EINS), in which the momentum change of elastically scattered neutrons is measured for different temperatures [1]. A Gaussian approximation analysis (see §3) provides the MSD of atomic motions as a function of temperature, in a given time and length scale. An effective force constant for the motions (resilience, (k’)) (N m⁻¹) is calculated from the temperature dependence [5]. The condition for the Gaussian approximation to be valid is that the amplitude of the MSD is well contained within the length–time window defined by the energy resolution and Q range of the spectrometer. In practice, this means that the analysis is limited to the low Q end of the range on IN13, where the length window is sufficiently wide to accommodate MSD of a few ångströms. Hydrogen atoms dominate EINS, because of a scattering cross section that is more than an order of magnitude higher than for other atoms. In the length and time scales
considered, hydrogen atoms in amino acid residues in a protein, for example, move with the groups to which they are bound and reflect well macromolecular internal motions as well as the larger fluctuation amplitudes involved in unfolding processes. In a complex system, hydrogen atoms in different groups have different motion amplitudes, so that the MSD is exactly that: the MSD of all hydrogen atoms. In the case of bacterial cells, the MSD was estimated to be dominated by hydrogen displacements in the proteome [7].

2.3. Effects of temperature stress on the molecular dynamics state of the \textit{Hs} proteome

The Gaussian approximation may be valid in different Q ranges to yield different MSD values, depending on the dominant amplitudes in the corresponding length scales. See, for example, the dynamics analysis of bacteriorhodopsin as a function of hydration in purple membranes of \textit{Hs} [16]. In this study, two motion populations were identified, a hydration-dependent large amplitude population at low Q (attributed to residues in outer loops of bacteriorhodopsin) and a small amplitude population at higher Q (attributed to vibrational internal motions of the residues in the membrane inserted helices). A low Q range (0.5 Å⁻¹ ≤ Q ≤ 1.5 Å⁻¹) was chosen in the current analysis (full double arrow in figure 3). It is sensitive to higher amplitude MSD values that would result, for example, from macromolecular unfolding processes in the stressed cells. Contributions from internal vibrational motions appeared in a higher Q range. Interestingly, no differences in NSD were detected between stressed and unstressed samples, suggesting that vibrational internal motions were less affected by the stress.

A clear increase in macromolecular flexibility was detected for the two heat-stressed \textit{Hs} cellular samples: the MSD values for the 55°C and 60°C stressed samples were shifted upwards progressively, compared with the unstressed control (figure 4a). The effective force constant (\(k'\)) expressing macromolecular resilience was also calculated from the slope of the MSD temperature dependence as described in [6] (figure 4b). The (\(k'\)) values indicated less rigid macromolecular states, which is likely to reflect perturbation in the folding state of a significant part of the \textit{Hs} proteome. Unfolded proteins have been measured to display larger fluctuations and lower resilience than folded states [17]. EINS studies on purified enzymes have shown that increased flexibility corresponds to the thermal unfolded state of the proteins. Russo \textit{et al.} [18] measured major losses in rigidity associated with the thermal unfolding of enzymes.

![Figure 1](http://rsif.royalsocietypublishing.org/)

**Figure 1.** Effect of thermal stress on the viability of \textit{Hs}. The cell cultures were heat-shocked at different temperatures (from 40°C to 60°C) for 1 h under agitation. Here, we present the data for the 60°C experiment. After centrifugation, the stressed cell paste was enclosed in the sample holder and the mean molecular dynamic properties of the cell were measured during 24 h by neutron scattering. To assess for the cell mortality rates after the experiment, the pellet was resuspended in the initial volume of cultivation media. Haloarchaeal cells were stained with the LIVE/DEAD BacLight kit. (a) Differential interference contrast image of \textit{Hs}. (b) Green fluorescence corresponds to viable \textit{Archaea} with intact membranes. (c) Red fluorescence indicates non-viable \textit{Archaea}. The viability rates were found to be over 95% after 1 h of stress and after the 24 h neutron experiment.
of a small beta protein and Koutsopoulos et al. [19] showed a clear correlation between the heat denaturation of an endo-
glucanase from *Pyrococcus furiosus* and increased atomic
fluctuation associated with lower resilience values [18,19].

Unfolded proteins in the cell may aggregate. However, on
the time scale of internal motions, it is not unlikely that such
aggregates display a lower rigidity compared with the folded
state, which has a compact core [3]. Still, more experimental
data on isolated proteins or cell extracts would be required to
support this hypothesis. From these studies, it can be inferred

that the increased MSD and lower resilience that we observed
*in vivo* for the temperature-stressed cells correspond to pro-
teome unfolding. The neutron experiments provided
therefore a direct measurement of the thermal stress effect on
the protein dynamics in the cytosolic context of an intact cell.

### 2.4. Conclusions

The observations reported in this paper represent a direct
insight into the folded state of a proteome exposed to tempera-
ture stress conditions. They reveal that neutron spectroscopy
can detect alterations in the mean molecular dynamics state
of the proteome within a living cell in response to environmen-
tal changes. Despite the crowded intracellular environment
and the induction of protein quality control systems, the
dynamic state of a large fraction of the proteome is strongly
perturbed under thermal stress. Interestingly, the cells could
easily recover when replaced in optimal growth conditions.
The experiments demonstrated the sensitivity of the neutron
scattering method to probe *in vivo* the molecular dynamic
alterations related to stress response.

### 3. Material and methods

#### 3.1. *Halobacterium salinarum* cell culture and
growth conditions

*Hs* RI cultures were grown at 37°C, with shaking at 150 r.p.m., in
standard medium containing 4.2 M NaCl [10]. Cells were grown
to mid-log phase (optical density at 600 nm (OD600) = 0.8).
For thermal stress, the cultures were transferred at the desired temperatures and cultivated under agitation for 1 h.

### 3.2. Sample preparation, neutron scattering experiments and analysis

For each sample, 2 ml of *Hs* cultures were centrifuged at 4000 g. The cell pellets were gently suspended with a brush in 500 ml of washing buffer containing 50 mM Tris–HCl pH 7.6; 4.2 M NaCl; 70 mM KCl and 80 mM MgSO4. The operation was repeated twice. A total of 540 mg of cell paste was taken from the pellet with a spatula to fill up completely a 0.3 mm path length gold-plated aluminium sample holders. The device was hermetically closed with an indium seal and mounted on the IN13 back spectrometer at the Institut Laue Langevin (http://www.ill.eu/html/instruments-support/instruments-groups/instruments/in13/).

Experiments on the cells were performed in H2O. The energy resolution and Q range of a neutron spectrometer define a window on a length and time scales. Motions confined within the window will contribute to observable elastic scattering [1]. On IN13, the window corresponds to an MSD of approximately 2 Å2 in 0.100 ns. Internal and unfolding motions are within the energy resolution and Q range. On IN13, the scattering without energy change [1].

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All measurements were repeated on samples from at least three different cultures. The IN13 spectrometer has an energy resolution of 8 μeV and is sensitive to motions that occur on a time scale up to about 0.1 ns. The EINS signal was measured and analysed over the scattering range 0.5 Å−1 < Q < 1.5 Å−1 (Q = 4π sinθ/λ for elastic scattering, where 2θ is the scattering angle and λ is the incident neutron wavelength). The elastic intensities were corrected for sample holder and buffer scattering, normalized to the scattering of vanadium (purely incoherent scattering) and corrected for sample absorption by using the ILL data reduction program LAMP (information on the program is available on the ILL website at http://www.ill.fr). For each sample, the elastic intensity I(Q) was obtained as a function of temperature, T, rising from 280 to 310 K, and ln(I(Q)) was plotted against Q². The MSD of the scattering nuclei, ⟨u²⟩, was calculated from the slope of the straight-line fit to the experimental data according to the Gaussian approximation that is valid for Q²⟨u²⟩ ~ 2 [5]:

\[
I(Q) = \text{const.} \times \exp(-\frac{1}{6}(u^2)Q^2),
\]

which can be written as

\[
\ln I(Q) = \text{const.} - \frac{1}{6}(u^2)Q^2.
\]

The value of the root MSD quantifies the global flexibility.

The mean resilience, ⟨k’⟩, was extracted from the slope of ⟨u²⟩ as a function of temperature T, using the following relation:

\[
\langle k' \rangle = \frac{0.00276}{\frac{d(u^2)}{dT}},
\]

where ⟨k’⟩ is in N m⁻¹, ⟨u²⟩ is in Å² and T is in degrees K.

The value of ⟨k’⟩ corresponds to a mean effective force constant and defines the average atomic resilience in a free energy potential [5,21].

### 3.3. Viability assays

Serial dilutions were performed on cultures and resuspended cell paste in order to examine the influence of temperature on the *Hs* intracellular dynamics. Viability tests were performed before and after stress exposure and before and after the 24 h neuron experiments. A Neubauer hemocytometer was used for cell count, and the LIVE/DEAD BacLight bacterial viability kit (Invitrogen detection technologies) was used to assess the mortality rate in the different physiological conditions as described [22]. Suspensions of *Hs* (4 × 10⁶ cells ml⁻¹) were pre-treated with LIVE/DEAD kit and pictures were taken with an epifluorescence microscope (IX 81 Olympus) coupled to a QImaging Retiga-SRV CCD digital camera. Fluorescence and contrast images were analysed using Velocity (Perkin Elmer). Survival was calculated as the number of viable cells following treatment divided by the number of viable untreated cells.

### 3.4. ICP measurement of intracellular K⁺ concentration

The K⁺ concentration in *Hs* cells was determined using induced coupled plasma spectrometry (ICP). After stress and neutron experiments, 1 ml of cell suspension was centrifuged. The pellets were washed with K⁺ free isotonic buffers and the cells were lysed by sonication in 10 ml of H2O. Analyses were performed with an ICP-AES Perkin Elmer, Optima 3300 DV plasma-mass spectrometer at the Equipe Géochimie de l’Environnement, LGIT/CNRS in Grenoble, France. A calibration curve was generated from KCl solutions. The total cell number and the average cell volume were determined in each sample by confocal microscopy. These values were used to calculate the intracellular K⁺ concentration. All measurements were performed in triplicate on at least three different experiments.

### 3.5. Protein immunodetection

One millilitre of OD 0.8 *Halobacterium* culture was pelleted and resuspended in 0.2 ml of distilled water. The cells were lysed and homogenized by flushing several times through a needle (diameter 0.6 mm) connected to a syringe. Cell debris was eliminated by centrifugation at 13000 r.p.m. for 10 min. The total protein amount in the supernatant was determined by using a Bradford quantification assay (BioRad). An equal volume of sample buffer was added to the extract. After boiling at 90°C for 5 min, 15 μl of this sample was subjected to 12 per cent SDS-polyacrylamide gels and the resolved proteins were transferred to nitrocellulose membranes (Hybond-P, GE healthcare). The blots were probed with anti-TF55 and anti MalDH antibodies obtained as described in [23]. All polyclona were used at a 1 : 10 000 dilution. Immunoreactive bands were visualized by chemiluminescence according to the supplier’s protocol (ECL detection kit; GE healthcare). The band intensities were measured by using the ImageJ software and were normalized by the total protein amount. The mean intensity values obtained for the 37°C unconditioned condition were arbitrary fixed to ‘1’ (figure 2).

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