

## Peptide-size-dependent active transport in the proteasome

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**Abstract.** – We investigate the transport of proteins inside the proteasome and propose an active-transport mechanism based on a spatially asymmetric interaction potential of peptide chains. The transport is driven by fluctuations which are always present in such systems. We compute the peptide-size-dependent transport rate which is essential for the functioning of the proteasome. In agreement with recent experiments, varying temperature changes the transport mechanism qualitatively.

*Introduction.* – Eukaryotic mammalian proteasomes are fascinating molecular machines that digest tagged with ubiquitin proteins [1]. The proteins are cleaved into peptides which can be further used for the synthesis of new proteins. The proteasome's function has been directly linked to a wide range of diseases, such as cancer, neurological diseases, and diseases of the immune defense system. Starting from the discovery of their activity in the late 1970s, presently proteasomes are in the focus of current molecular biology, *e.g.* [2,3]. By now there are several models to explain the proteasome's activity [4], cleavage mechanisms [5–7] and to predict the cleavage results [8]; however, the basic principles of the proteasome operation mechanisms are still poorly understood, mainly due to the lack of experimental results.

In this letter, we focus on the understanding of protein translocation inside the proteasome, leaving the mechanisms of cleavage, targeting, etc., beyond the scope. The main question is: given proteasomes are highly complex pipe-like structures of tenthousands atoms of almost perfect left-right symmetry with respect to the axis of the pipe [9], thus, operating bi-directional. To be cleaved, a protein has to enter the proteasome at one side, pass the active sites (where the cleavage occurs), which makes about 1/3 of the total length of the pipe. Then the cleaved peptides have to pass all the way through the pipe to finally reappear at the other side of the proteasome. Although there is a number of examples where protein transport in cells occurs due to diffusion, *i.e.* Brownian motion [10], diffusive transport may be excluded as the main mechanism for translocation in the proteasome because of the enormous cargo [11]. Also other proposed transport mechanisms, such as the power stroke model of protein translocation, do not suffice to explain translocation [12]. Therefore, the question arises: how is the protein's motion inside the proteasome driven?

Since proteasomes are large multi-subunit structures consisting of proteins, the mechanism of the protein transport is directly related to protein-protein interaction. In [13] it has been noticed that if attachment and detachment rates are specified asymmetrically, the protein-protein binding interaction acts as a ratchet. Following this argumentation, active protein transport, based on the mechanism of a molecular ratchet, has been studied for transport through membranes [14] and has been also discussed as a mechanism for cytosolic destruction by the proteasome [15]. Moreover, maximum-likelihood tests have shown that other models, *e.g.* the power stroke model of protein translocation, do not lead to better agreement with the experiment than the Brownian ratchet model [12]. Noteworthy, in these ratchet effects transport is possible only in a certain temperature interval, and stochasticity, intrinsically present due to fluctuations in any biochemical reaction [16,17], provides the driving mechanism.

Here we propose a model for active protein translocation in the proteasome to explain the peptide size dependence of the transport velocity as well as its temperature dependence which possibly explains the mechanism of temperature reaction or heat shock response, regulating the proteasome activity in the case of some diseases [18–21]. The results describe a system size ratchet effect that is related to similar effects which have been described recently for other noise-induced phenomena [22]. Already the 20S proteasome reveals full functionality (see *in vitro* results in [2, 3]) and the proteasome has a substantial length; thus, for studying the transport we disregard the enzymatic action of the proteasome caps (boundary effects).

*Model.* – Following, *e.g.*, [13] we assume that the protein-proteasome interaction is characterized by a spatially periodic asymmetric potential, motivated by the regular (periodic) structure of unfolded proteins due to the regular backbone structure of amino acid chains. The asymmetry of this potential is caused by the asymmetry of the C-N binding between the amino acids of proteins that enter the proteasome *always* in the  $N \rightarrow C$  direction, *i.e.*, with N head on [6] (see fig. 1, left). The folding structure of the protein is not relevant here, since due to the action of the regulatory complexes the protein enters the proteasome unfold [23]. Moreover, even fold proteins tend to form periodic structures to maximize their amphiphilicity [24].

As an abstraction, here we assume that the protein-proteasome interaction potential  $U(x)$  is periodic with the period  $L$ . In reality, there is a basic asymmetry, namely the C-N asym-

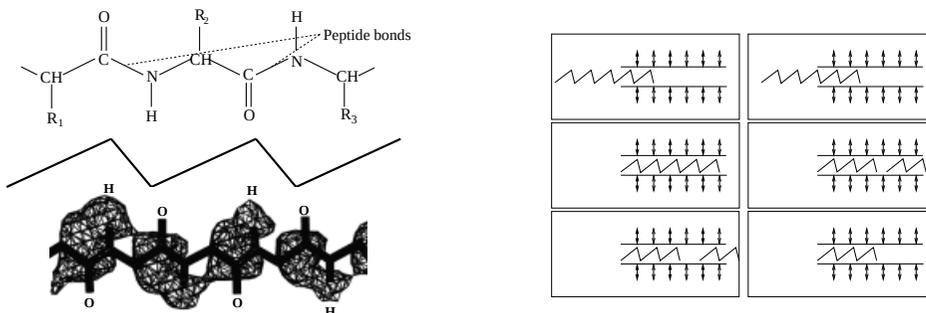


Fig. 1 – Left: The regular part of the protein-proteasome interaction potential is modeled by a spatially asymmetric periodic potential, based on the C-N asymmetry of the peptide backbone. The  $R_i$  mark the amino-acid-specific residues. The lower part shows the equipotential surfaces of a simple peptide which is clearly asymmetric (see also [25]). The H and O atoms are marked for orientation. Right: From top left to bottom right: A protein moves into the proteasome while increasing the number  $N$  of interaction centers. After cleavage  $N$  abruptly decreases for the resulting peptides. When leaving the proteasome,  $N$  decreases as well.

metry of the protein (or peptide) backbone, that is superposed by a nonperiodic (in our sense irregular) part, attributed to the amino-acid-specific residues [25]. The basic structure of a peptide and its model as an asymmetric ratchet potential is sketched in fig. 1 (left). The commensurability of the protein and the proteasome is supported by the fact that both biological macromolecules consist of the same basic structures, namely chains of amino acids. This commensurability has been claimed before, and was the basis of automaton-like models of proteasome digestion processes [6]. The detailed form of the asymmetric periodic interaction potential is less important for this qualitative study. Here, we assume a sawtooth potential, fig. 1. The angles are smoothed, *i.e.*,  $dU/dx$  exists in each point (for details see [26]).

The proteasome acts upon the protein by a certain number of equidistant interaction centers. The dynamics of the protein inside the proteasome is, hence, governed by  $N$  interactions centers, where  $N$  is the number of protein elements (amino acids or multiplicatives of it). There appear the following forces: potential force (protein-proteasome interaction)  $-N\partial U(x)/\partial x$ , fluctuations with collective  $NF(t)$  and individual components  $f_1(t) + \dots + f_N(t)$ , and protein friction forces  $N\beta\dot{x}$  [13], where  $x$  is the coordinate of the protein with respect to the proteasome and  $\beta$  is the coefficient of friction. Due to the small mass of all protein particles, moving in the liquid cytosol, the motion occurs in the overdamped realm [27]; hence we neglect inertia forces. Note that transport is possible only in the case of nonequilibrium fluctuations. In the simplified case, when fluctuations can be represented by a sum of a collective periodic force and individual for every protein residue thermal noise, the model is analytically tractable, predicting the velocity dependence on the peptide size. Numerically, we investigate also different kinds of fluctuations in order to be closer to reality. Normalizing all forces by friction and taking  $\beta = 1$ , the translocation of a protein in the proteasome is governed then by

$$\frac{\partial x}{\partial t} = -\frac{\partial U(x)}{\partial x} + F(t) + \frac{1}{N}(f_1(t) + \dots + f_N(t)). \quad (1)$$

When the protein chain enters the proteasome, the number of interaction centers  $N$  is increased (fig. 1, right). After cleavage,  $N$  is abruptly decreased for the cleavage products. We believe that this crucially changes the transport velocity, providing the cleavage product leaving the proteasome. Equation (1) models the motion for both the initial protein and the peptides after cleavage. We consider three cases of fluctuations, regulating their motion:

*Case 1:* We assume collective oscillations of the peptide elements [28], *e.g.*,  $F(t) = A\cos(\omega t)$ , where  $A$  and  $\omega$  stand for the amplitude and frequency of these oscillations. *In vitro* experiments have shown that the 20S proteasome operates without ATP consumption [2]. So far, the source of energy of the 20S proteasome was not discussed in the literature. We believe that it is supplied by the cleaved peptide itself, as cleavage of peptide chemical bindings is an exothermic reaction. In our model, this energy leads to oscillations of the proteasome and, consequently, to the harmonic noise or periodic force. Additionally, each interaction center undergoes local thermal fluctuations, represented by mutually uncorrelated white noise of intensity  $\sigma^2$ :  $f_i(t) = \xi_i(t)$ , where  $\langle \xi_i(t)\xi_j(t') \rangle = \sigma^2\delta(t-t')\delta_{ij}$ . In this case, the stochastic term in eq. (1) is white noise of intensity  $\sigma^2/N$ . The Fokker-Planck equation for the peptide coordinate probability distribution  $w(x, t)$  associated with eq. (1) is

$$\frac{\partial w}{\partial t} = -\frac{\partial}{\partial x} \left[ \left( F(t) - \frac{\partial U}{\partial x} \right) w(x, t) \right] + \frac{\sigma^2}{2N} \frac{\partial^2 w(x, t)}{\partial x^2},$$

which may be solved in quasi-stationary adiabatic approximation  $\partial w/\partial t = 0$  [29]. We obtain

$$\frac{\sigma^2}{2N} \frac{\partial w(x, F)}{\partial x} - \left( F - \frac{\partial U}{\partial x} \right) w(x, F) = -G(F), \quad (2)$$

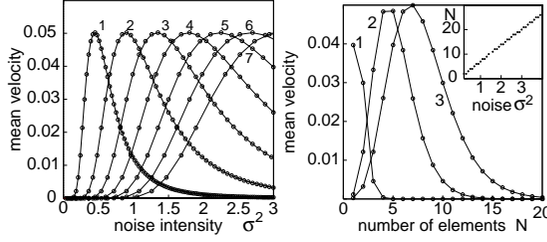


Fig. 2 – Left: mean velocity *vs.* noise intensity for  $N = 1, 2, \dots, 7$ . Right: mean velocity *vs.*  $N$  for different temperatures:  $\sigma^2 = 0.6$  (curve 1), 2, and 3 (curves 2 and 3). The inset plot shows the depth of the protein penetration for velocity 0.0005.

where  $G(F)$  is the probability flux. For any periodic potential  $U(x)$ , the quasi-stationary solution of eq. (2) is

$$w(x, t) = \left[ C(F) - \frac{2G(F)}{\sigma^2/N} \int_0^x \exp \left[ \frac{U(x') - Fx'}{\sigma^2/2N} \right] dx' \right] \exp \left[ -\frac{U(x) - Fx}{\sigma^2/2N} \right], \quad (3)$$

where  $C(F(t))$  and  $G(F(t))$  are unknown functions of  $t$ . Using the periodicity condition  $w(0, t) = w(L, t)$  and the normalization of  $w(x, t)$ , we get  $G(F)$ . If the amplitude  $A$  meets the condition  $LA \ll \sigma^2/N$ , one can expand  $G(F)$  and obtain

$$G(F) \approx G_{01}F + G_{02}F^2 \quad (4)$$

with the expansion coefficients  $G_{01} = L/(I_{10}I_{20})$ ,

$$\begin{aligned} G_{02} &= G_{01} \left[ \frac{I_{11}}{I_{10}} - \frac{I_{21}}{I_{20}} - \frac{NL}{\sigma^2} \left( 1 - \frac{2I_{30}}{I_{10}I_{20}} \right) \right], \\ I_{10} &= \int_0^L e^{U'(x)} dx, \quad I_{20} = \int_0^L e^{-U'(x)} dx, \quad I_{11} = \frac{2N}{\sigma^2} \int_0^L x e^{U'(x)} dx, \\ I_{21} &= \frac{2N}{\sigma^2} \int_0^L x e^{-U'(x)} dx, \quad I_{30} = \int_0^L \int_0^x e^{(U'(x') - U'(x))} dx' dx, \quad U'(x) = \frac{2NU(x)}{\sigma^2}. \end{aligned} \quad (5)$$

Substituting eq. (4) into  $\overline{\langle \dot{x} \rangle} = \int_0^L \overline{G(x, t)} dx$ , where  $\overline{(\cdot)}$  denotes time averaging, yields the average protein transport velocity, as a function of the noise intensity  $\sigma^2$  and the peptide size  $N$ ,

$$\overline{\langle \dot{x} \rangle} \approx \frac{L^2 A^2}{2I_{10}I_{20}} \left[ \frac{I_{11}}{I_{10}} - \frac{I_{21}}{I_{20}} - \frac{NL}{\sigma^2} \left( 1 - \frac{2I_{30}}{I_{10}I_{20}} \right) \right]. \quad (6)$$

The dependence of the transport velocity on the noise intensity, *i.e.* temperature, is shown in fig. 2 (left) for different system size  $N$ , representing typical dependencies of the rocking ratchet [30]. This plot shows that protein transport is possible only in a certain temperature interval. Absence of noise leads to vanishing transport while too high temperature also decreases the velocity. For given temperature, the peptide size  $N$  strongly influences the transport velocity, *i.e.*, with increasing  $N$  the dependence is shifted to the right and becomes wider (curves 1-3). This dependence differs qualitatively for small and large temperature. For small temperature, the velocity decays with increasing  $N$  (curve 1 in fig. 2 (right)). Consequently, when a protein enters the proteasome, initially it moves at large velocity. Then, due to the increase of  $N$  the velocity decreases, and the motion of the protein virtually terminates. At

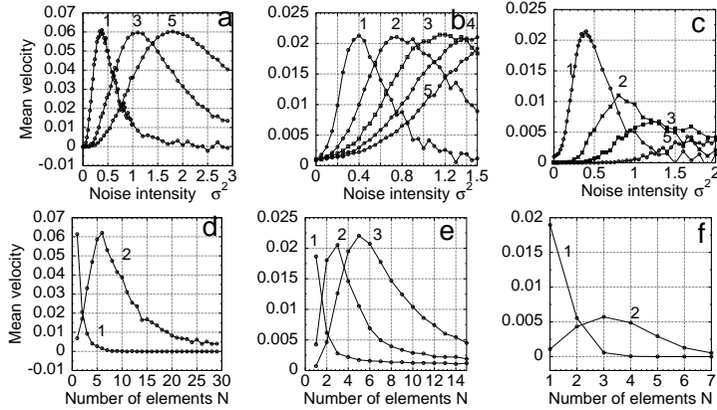


Fig. 3 – a, b, c: mean velocity *vs.* noise intensity for  $N = 1, 2, \dots, 5$ ,  $A = 1.15$ ,  $\omega = 0.1$ . d, e, f: mean velocity *vs.*  $N$  for different temperatures: d)  $\sigma^2 = 0.3$  (1), 2 (2); e)  $\sigma^2 = 0.3$  (1), 1 (2), and 2 (3); f)  $\sigma^2 = 0.5$  (1) and 1.5 (2). a, d: periodic force, b, e: colored noise, c, f: without global fluctuations.

this stage the probability of cleavage is much higher than that of translocation; thus, with large probability the protein will be cleaved there. After cleavage,  $N$  is significantly decreased (fig. 1 (right)), and following the velocity dependence, both fragments move rapidly to leave the proteasome. In agreement with our model, the blocking of uncleaved proteins inside the proteasome has been observed also experimentally [31].

Surprisingly, for large temperature the velocity depends qualitatively different on the peptide size (curves 2, 3 in fig. 2 (right)), *i.e.*, there is an optimal peptide size that corresponds to maximal velocity. In this case, the protein moves accelerated while entering the proteasome until the optimal velocity is achieved. Then the velocity decreases, and again it virtually sticks. After cleavage, the behavior is quite different: if the cleaved peptide is close to the optimal size, it leaves the proteasome rapidly. If the cleaved fragment is too small or too large, it moves with small velocity, thus it blocks the proteasome leading to low efficiency or malfunction of the proteasome. Note that for the protein that enters the proteasome the initial small  $N$  increases, and the velocity will increase, whereas for the cleaved fragment of smaller  $N$  the velocity remains constant. There is experimental evidence [31] that certain proteins cannot be cleaved by the proteasome, *i.e.*, they are inert with respect to its active sites. For such proteins we can determine the penetration depth, defined as the depth  $N$  at which the propagation velocity falls below a predefined very small threshold. The inset of fig. 2 (right) shows that the penetration depth increases linearly with temperature. So far, there are no experimental data to verify this prediction.

Consequently, small temperature leads to preferably small size of the cleaved fragments, whereas larger temperature leads to domination of some specified peptide size in the proteasome output. As a hypothesis, we believe that the qualitatively different proteasome behavior with respect to temperature may be responsible for temperature reaction or heat shock response [18, 19, 21], providing immune defense in the case of some diseases. For large temperature (fig. 2 (right) curve 2, 3), the protein is accelerated while entering the proteasome, hence leading to more efficient operation of the proteasome.

To confirm these results we have performed computer simulations [32] whose results agree with our analytical findings (fig. 3). Some quantitative deviations result from taking into account only a limited number of terms in the expansion eq. (4).

*Case 2:* Since collective fluctuations of the protein or peptide elements, represented by a periodic-in-time force, can be used only under certain assumptions, next we consider the system in the absence of deterministic periodic forces, but with colored noise  $F(t) = \zeta(t)$ , where  $\zeta(t)$  is harmonic noise with some dominating frequency (due to the molecular spectrum and oscillations of the peptide chain [28]). We obtain this noise as the solution of

$$\ddot{\zeta} + 2\delta\dot{\zeta} + \omega_0^2\zeta = \xi', \quad (7)$$

where  $\xi'$  is the Gaussian noise,  $\langle \xi'(t)\xi'(t') \rangle = \sigma_a^2\delta(t-t')$ , and the parameters  $\delta = 0.01$ ,  $\omega_0 = 0.1$ ,  $\sigma_a^2 = 10^{-4}$  regulating the width, the dominating frequency of the power spectrum, and the intensity of  $\zeta(t)$ . The results agree with the case of the periodic force (fig. 3 (b, e)). Hence, if exclusively noisy fluctuations are present in the system, the generic behavior described above persists and leads to the same temperature and peptide size dependences.

*Case 3:* Next we consider the case of no collective fluctuations,  $F(t) = 0$ , *i.e.*, there is no eigen-collective mode in the peptide oscillations. The local noise of each interaction center is presented now by the sum of uncorrelated colored- and white-noise components:  $f_i(t) = \zeta_i(t) + \xi_i(t)$ . The colored noise generates nonequilibrium fluctuations, that is a necessary condition for directed transport. The motion of the protein or peptide is described by

$$\dot{x} = -\frac{\partial U(x)}{\partial x} + \frac{1}{N}(\zeta_1 + \dots + \zeta_N + \xi_1 + \dots + \xi_N), \quad (8)$$

with  $\zeta_i(t)$  and  $\xi_i(t)$  as above. Also in this case the behavior is similar, but an increase of the peptide size leads not only to a shift of the velocity dependence towards large noise intensity, but also to a decrease of its maximal values (fig. 3(c)), since nonequilibrium fluctuations originating from different interaction centers act occasionally in converse direction. For small temperature (fig. 3(f), curve 1) the transport decreases with increasing peptide size  $N$ , whereas for large temperature the rate depends nonmonotonously on  $N$  (curve 2). The optimal velocity in the latter case is smaller, as for the case of collective fluctuations.

*Conclusion.* – A model for active protein transport inside the proteasome has been investigated analytically and numerically. The peptide size as well as temperature influence the transport rate significantly in agreement with experimental results. Following our predictions, protein transport occurs only in a certain interval of temperature. The size dependence of the transport velocity leads to a preferred fragment (peptide) size, since for such sizes the probability of cleavage exceeds the probability of further translocation significantly. Under certain conditions, uncleaved proteins may get stuck inside the proteasome, thus leading to decrease of its efficiency, which is in good agreement with experimental observations [31]. Calculating the average velocity, we have considered only the periodic constituent of the protein-proteasome potential. Slight inhomogeneity (nonperiodicity) of this potential will lead to fluctuations of the velocity around its average, leaving the qualitative behavior of dependencies on the peptide size intact.

Varying the temperature, the model predicts qualitatively different transport regimes, thus possibly explaining the mechanism of temperature reaction or *heat shock response* in the case of certain diseases as well as in *in vitro* results that can be obtained in a wider range of temperature. In agreement with experiments, there is a preferred fragment size, that dominates the proteasome output. We emphasize that our model is based only on two assumptions: spatial asymmetry of the protein chain and presence of nonequilibrium fluctuations, which are certainly fulfilled in proteasomes.

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