How to Quantify the Chaperone-Like (Anti-Aggregation) Activity?

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Folding of newly synthesized protein chains can be accompanied by the formation of non-native forms prone to aggregation; stress conditions also produce unfolded protein forms which tend to aggregate [1-5]. Investigators aim for elucidation of the mechanisms of protein aggregation and mechanisms of the protective action of agents possessing chaperone-like (anti-aggregation) activity, namely small heat shock proteins (sHsp) and low-molecular-weight compounds (aminoacids, cyclodextrines, polyamines and so on) [6-8].

The agents possessing chaperone-like (anti-aggregation) activity find application in biotechnology and medicine. To screen agents exhibiting efficient protective action, the quantitative evaluation of the degree of protein aggregation suppression is requisite. Here are discussed the quantitative methods of estimation of the anti-aggregation activity.

Estimation of the initial rate of protein aggregation

Protein aggregates possess higher light scattering capability in comparison with the non-aggregated protein molecules, and therefore the simple way to register protein aggregation is the measurement of the increment of light scattering intensity (I) or apparent optical absorbance begins to increase. Parameter $K_{agg}$ is a measure of the initial rate of aggregation.

$$I - I_0 = K_{agg} (t-t_0)^2 \quad \text{or} \quad A - A_0 = K_{agg} (t-t_0)^2, \quad (t > t_0) \quad (1)$$

Where $I_0$ and $A_0$ are the initial value of the light scattering intensity and apparent optical absorbance, respectively, at $t=0$ and $t_0$ is the duration of lag period on the kinetic curve ($t_0$ is a point in time at which the light scattering intensity or apparent optical absorbance begins to increase). Parameter $K_{agg}$ is a measure of the initial rate of aggregation. The applicability of equation (1) for the description of the initial parts of the kinetic curves of protein aggregation was demonstrated for heat denaturation of glycogen phosphorylase b (Phb) [9-12], creatine kinase [13] and glyceraldehyde-3-phosphate dehydrogenase (GAPDH) from rabbit skeletal muscles [14,15] and dithiothreitol-induced aggregation of a-lactalbumin [16] and insulin [17]. The measurements of the hydrodynamic radius of protein aggregates using dynamic light scattering allow us to conclude that for the proteins under study the point in time $t=t_0$ corresponds to the appearance of start aggregates. Start aggregate contains hundreds of denatured protein molecules. The formation of the start aggregates proceeds on the all-or-none principle. The intermediate states between the non-aggregated protein and start aggregates are not detected in the system [18].

When studying the kinetics of aggregation of polyglutamine peptides [19-21], it was experimentally shown that accumulation of monomers incorporated in the aggregate proceeded according to the time squared law. Theoretical analysis shows that such a law should be valid for nucleation-dependent aggregation [9,22].

The measurement of the $K_{agg}$ value at various initial concentrations of the protein $[P]_0$ allows calculating the order of aggregation with respect to the protein (n):

$$K_{agg} = \text{const} \cdot [P]_0^n \quad (2)$$

In the case of thermal aggregation of Phb [9] and GAPDH [15] the dependence of parameter $K_{agg}$ on $[P]_0$ is linear ($n=1$). This means that the stage of unfolding of a protein molecule proceeds with a substantially lower rate than the following stages of aggregation of the unfolded protein molecules. When unfolding of the protein molecule is a relatively fast process and the stages of aggregation become rate limiting, parameter $n$ exceeds unity. For example, the analysis of the data on thermal aggregation of β-crystallin from bovine lens at 60°C (pH 6.8) [23] and thermal aggregation of yeast alcohol dehydrogenase at 56°C (pH 7.4) [24] shows that parameter $n$ is close to 2.

When analyzing the shape of the kinetic curves of aggregation of Phb denatured by UV radiation [25], we observed that equation (1) is not fulfilled and, to characterize the initial rate of aggregation, we proposed to use the time interval ($t_0$) over which the initial value of the light scattering intensity is doubled. To calculate the $t_0$ value, the initial part of the dependence of the light scattering intensity on time was described by the stretched exponent:

$$I = I_0 \exp \left( \ln(2) \left( \frac{t}{t_0} \right)^m \right) \quad \text{(3)}$$

Where $m$ is a constant. Analysis of the kinetics of aggregation of UV-irradiated Phb shows that the initial parts of the kinetic curves can be described by the equation containing simple exponent:

$$I = I_0 \left[ 1 + K \left( \exp(Kt) - 1 \right) \right], \quad (t > t_0) \quad (4)$$

where $K$ and $K_t$ are constants. This equation can be rearranged in a form containing parameter $t_0$:

$$I = I_0 \left[ 1 + K \exp \left( \frac{t}{t_0} \ln \left( \frac{1+K}{K} \right) - 1 \right) \right] \quad \text{(5)}$$

The reciprocal value of $t_0$, namely $1/t_0$, may be considered as a measure of the initial rate of aggregation. The higher the $1/t_0$ value, the higher is the initial rate of aggregation. It should be noted that the test system based on aggregation of UV-irradiated Phb, in contrast to the test system based on thermal aggregation of proteins, does not contain the stage of unfolding of the protein molecule. Phb denatured by UV irradiation is assembled in primary aggregates with the hydrodynamic radius of 10.4 nm [25].

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There is another way to characterize the initial rate of aggregation. As noted above, the dependences of the light scattering intensity ($I$) or apparent optical absorbance ($A$) on time for aggregation process are usually S-shaped. The analysis of the literature data for the kinetics of protein aggregation shows that at above the inflection point the dependence of $I$ or $A$ on time follows the exponential law [26-30]:

$$I = I_0 + \left[ (I_1 - I_0) \times \exp\left(-k(1-r^t)\right) \right] + \left[ (A_1 - A_0) \times \exp\left(-k(1-r^t)\right) \right]$$

where $I_{\text{lim}}$ and $A_{\text{lim}}$ are the limiting value of $I$ and $A$, respectively, at $t \rightarrow \infty$ and $t^*$ is a length on the $I=I_0$ (or $A=A_0$) horizontal line cut off by the theoretical curve calculated with equation (6).

The slope of a tangent to the theoretical curve passing through the point with coordinates $[t=t^*; I_0=I; A_0=A]$ is equal to the product $k(I_{\text{lim}}-I_0) \div k(A_{\text{lim}}-A_0)$, which is a measure of the initial rate of aggregation. It should be noted that lag period may be lacking on the kinetic curves of aggregation. This renders using equation (1) and (5) impossible. In such a situation the characterization of the initial rate of aggregation through a product $k(I_{\text{lim}}-I_0)$ or $k(A_{\text{lim}}-A_0)$ becomes very reasonable, as it was demonstrated by us for thermal aggregation of tobacco mosaic virus protein coat (50 mM phosphate buffer, pH 8.0, 52°C; [31]). To determine the order of aggregation with respect to protein, the equation analogous to equation (2) can be used:

$$k(I_{\text{lim}} - I_0) = \text{const} \times [P]^n \quad \text{or} \quad k(A_{\text{lim}} - A_0) = \text{const} \times [P]^n.$$

(7)

For thermal aggregation of firefly luciferase [30] and tobacco mosaic virus coat protein [31] parameter $n$ was found to be close to 2. This means that the rate-limiting stage for the overall process of aggregation is the stage of protein aggregation. The $n$ value higher than 2 was obtained for heat-induced aggregation of ovalbumin at 80°C ($n=2.65$ [32]). It is notable that first-order kinetics ($n=1$) was observed for thermal aggregation of creatine kinase from rabbit skeletal muscles [33] and aggregation accompanying renaturation of denatured carbonic anhydrase [32]. Thus, the rate-limiting stage for the overall aggregation process is the monomolecular stage (unfolding of the protein molecule in the case of creatine kinase or the initial stage of unfolding of denatured protein in the case of carbonic anhydrase).

Quantification of anti-aggregation activity of protein chaperones

When analyzing the dependence of the initial rate of aggregation ($v$) on the concentration of protein chaperone, one should take into account two circumstances. First, the binding of a chaperone to a protein substrate is rather firm, and suppression of aggregation is studied under the conditions when the initial concentrations of a chaperone and protein substrate exceed sufficiently the dissociation constant for the complex chaperone–protein substrate. This means that the dependence of $v$ on [chaperone] is a titration curve which allows determining the stoichiometry of chaperone–protein substrate complex. Second, in accordance with equation (2) or (7) the protein concentration $[P]^n$ is proportional to $v^n$. This means that the proportional decrease in the concentration of the protein substrate (for example, as a result of the complexation with a chaperone) should result in the proportional decrease in the $v^n$ value. Thus, to determine the stoichiometry of the chaperone–protein substrate complex, one should construct the $v^n$ versus [chaperone] plot. Besides, the dependence of $v$ on [chaperone] may be nonlinear because of the formation of the chaperone–protein substrate complexes with different stoichiometry.

The dependence of $v$ on [chaperone] consisting of two linear parts was observed for suppression of UV-irradiated Phb by α-crystallin [25]. The dependence of $v^{1/n}$ on [sHsp] of such a type is schematically represented in figure 1A. One can assume that the complicated shape of the $v^{1/n}$ versus [sHsp] plot is due to the dynamic structure of α-crystallin and the initial part of the dependence of $v^{1/n}$ on [sHsp] corresponds to the complexes of the dissociated forms of α-crystallin with the protein substrate. The second linear part corresponds to the formation of the α-crystallin–protein substrate complexes where the adsorption capacity of α-crystallin in respect to the protein substrate becomes decreased. The length cut off on the abscissa axis by the second linear part, [sHsp]*, gives the stoichiometry of the chaperone–protein substrate complex of the second type. The absissa of the point of intersection for two linear parts gives the stoichiometry of the chaperone–protein substrate complex of the first type. It is significant that the titration curves obtained at various concentrations of the protein substrate should retain the same form in the coordinates $[(v/v_t)^{1/n}; [sHsp]/[protein substrate]]$ where $v_t$ is the initial rate of aggregation in the absence of the chaperone. This methodology should be proved by concrete examples.

In works [32,34,35] the $kA_{\text{lim}}$ product was used for the quantitative characterization of the chaperone-like activity of whole casein and β-casein. It is of interest that the $kA_{\text{lim}}$ value versus [whole casein] plot for suppression of aggregation accompanying renaturation of denatured carbonic anhydrase by whole casein looks like a curve consisting of two linear parts [32].

Experimental data on suppression of thermal aggregation of the catalytic subunit of protein kinase CK2 (CK2α) at 40°C by C-terminal domain of glucose-regulated protein (grp94-CCKT) obtained by Roher et al. [36] were analyzed by Kurganov [27]. To calculate the initial rate of aggregation, the $kA_{\text{lim}}$ product was calculated. The linear character of the dependence of the $kA_{\text{lim}}$ value on the molar ratio [grp94-CCKT]/[CK2α] was demonstrated. The following stoichiometry of the grp94-CCKT–CK2α complex was calculated from the length cut off on the abscissa axis by the straight line: 4 molecules of grp94-CCKT per 1 molecule of CK2α.

Quantification of anti-aggregation activity of chemical chaperones

Protective effect of chemical chaperones is revealed as a diminishing of the initial rate of aggregation in the presence chemical chaperone (Figure 1B). When studying the protective action of chemical chaperones (2-hydroxylpropyl-β-cyclodextrin [15] or proline [25]), we showed that the dependence of the initial rate of aggregation $v$ expressed by parameter $K_v$ or $1/t_v$, on the concentration of chemical chaperone (I) followed the Hill equation [37]:

$$v = \frac{v_t}{1 + [I]/[I]_{1/2}}^{1/n},$$

(8)

Where $v_t$ is the initial rate of aggregation in the absence of a chaperone, $[I]_{1/2}$ is the concentration of semisaturation, i.e., the concentration of the chaperone at which $v/v_t=0.5$, and $h$ is the Hill coefficient. The value of $h$ was found to be 1.8 for 2-hydroxylpropyl-β-cyclodextrin and 1.6 for proline. The values of $h$ exceeding unity are indicative of the existence of positive cooperative interactions between chaperone-binding sites in the protein substrate molecule [37]. Parameter $[I]_{1/2}$ may be considered as a measure of the affinity of the chaperone to the protein substrate. The lower the $[I]_{1/2}$ value, the higher is affinity of the chaperone to the protein substrate. It is significant that the shape of the dependence of the initial rate of aggregation on the chaperone concentration remains unchangeable at variation of the protein substrate concentration as was shown, for example, for suppression of protein aggregation by proline [25].
It should be noted that in both cases the UV-irradiated protein (GAPDH [15] or Phb [25]) was used as a protein substrate. It has been just such protein substrates which allowed us to register the effect of chemical chaperones directly on the stage of aggregation of denatured proteins. In other test-systems, for example, in the test-systems based on heat-induced aggregation of proteins, the general effect of chemical chaperone involves the action of chaperone on the aggregation stage and the action of chaperone on the stages preceding the aggregation stage [10,38-40].

Combined action of chaperones

The protective activity of sHsp can be modulated by the chemical chaperones. For example, it was demonstrated that arginine enhanced aggregation stage and the action of chaperone on the stages preceding the initial rate of aggregation, parameter 1/v0 for inhibitor 1, inhibitor 2 and inhibitor 1 + inhibitor 2, respectively. When the action of one inhibitor is not dependent on the presence of the other, parameter j is equal to unity. The case j > 1 corresponds to synergism and the case j < 1 corresponds to antagonism in the combined action of two inhibitors.

We used parameter j for analysis of combined action of α-crystallin and proline on aggregation of UV-irradiated Phb [25]. To characterize the initial rate of aggregation, parameter 1/i1/2 was calculated. When concentration of proline was equal to 0.15 M, a slight antagonism was observed between α-crystallin and proline (the values of parameter j fall in the interval 0.81–0.91). However, at higher proline concentration (0.5 M) each inhibitor (α-crystallin or proline) acts independently of one another.

Parameter j may be used also for analysis of combined action of protein chaperones. For example, Yousef and Jalili [44] studied the combined action of α-crystallin and β-casein on dithiothreitol-induced aggregation of bovine pancreatic insulin. The kA/kim product was used for characterization of the initial rate of aggregation. The analysis of the data presented in this work shows that parameter j is close to unity. Thus, two chaperones act independently of each other.

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