Kinetics of Actin Unfolding Induced by Guanidine Hydrochloride†

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ABSTRACT: The kinetics of actin unfolding induced by guanidine hydrochloride has been studied. On the basis of obtained experimental data a new kinetic pathway of actin unfolding was proposed. We have shown that the transition from native to inactivated actin induced by guanidine hydrochloride (GdnHCl) passes through essential unfolding of the protein. This means that inactivated actin should be considered as the off-pathway species rather than an intermediate conformation between native and completely unfolded states of actin, as has been assumed earlier. The rate constants of the transitions that give rise to the inactivated actin were determined. At 1.0−2.0 M GdnHCl the value of the rate constant of the transition from native to essentially unfolded actin exceeds that of the following step of inactivated actin formation. It leads to the accumulation of essentially unfolded macromolecules early in the unfolding process, which in turn causes the minimum in the time dependencies of tryptophan fluorescence intensity, parameter $A$, characterizing the intrinsic fluorescence spectrum position, and tryptophan fluorescence anisotropy.

Protein folding and refolding is often accompanied by the association of the partially folded intermediates ($I−8$). Protein association or aggregation represents an essential problem in biotechnology as overexpression of recombinant proteins is often complicated by the formation of the misfolded protein accumulated in inclusion bodies ($9−13$). The association of specific proteins and deposition of insoluble protein fibrils (amyloid) in various organs result in the development of many disorders such as Alzheimer’s and Parkinson’s diseases, malignant myeloma, cataracts, prion affictions, and many other abnormalities ($7, 16−25$). The crucial importance of protein aggregation originating from partially folded species gave rise of a great number of studies considering this problem (see, e.g., ref 7).

Actin, one of the major proteins of muscle tissue, is a globular protein with a molecular mass of 42 kDa that is known to bind one divalent ion, Ca$^{2+}$, and one molecule of ATP. Actin contains four tryptophan residues (26). Only two of these tryptophan residues, Trp 340 and Trp 356, which are localized in the hydrophobic environment, give essential contribution to actin intrinsic fluorescence (27, 28). At low ionic strength actin exists as a monomer (G-actin), but in the presence of neutral salts it is polymerized into a double-stranded polymer, which is known as the fibrous form of actin, or F-actin.

The release of calcium ion by EDTA or EGTA treatment leads to the transformation of G-actin into the inactivated form (I), in which the protein molecule loses its capability to polymerize (29, 30). Inactivated actin also may be obtained as a result of heat denaturation (29−41), moderate urea or GdnHCl1 concentration (30, 38), dialysis from 8 M urea or 6 M GdnHCl (38, 39), or spontaneously during storage (38). Properties of inactivated actin are invariant to the way of denaturation (30, 42, 43). It also has been shown that inactivated actin is a thermodynamically stable monodisperse associate, consisting of 15 monomers (43). It has a unique structure, with hydrophobic clusters on the surface and polar regions in the interior (30, 42, 43). Thus inactivated actin represents an interesting system for the investigation of different aspects connected with the self-association of partially folded proteins.

In this work we focused on the process of the formation of inactivated actin. To elucidate the mechanism of inactivated actin formation, the kinetics of GdnHCl-induced actin unfolding was studied. It was shown that the essential unfolding of the protein macromolecule precedes the formation of inactivated actin.

MATERIALS AND METHODS

Preparations. Rabbit skeletal muscle actin was purified by the standard procedure (44). G-actin in buffer G (0.2 mM ATP, 0.1 mM CaCl$_2$, 0.4 mM β-mercaptoethanol, 5 mM Tris-HCl, pH 8.2, and 1 mM NaN$_3$) was stored on ice and used within 1 week. Actin was purified by one or two cycles of polymerization−depolymerization using 30 mM KCl for polymerization. Actin samples with the A parameter not

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1 Abbreviations: CD, circular dichroism; UV, ultraviolet; GdnHCl, guanidine hydrochloride; N, native; U, unfolded; I, inactivated actin.
lower than 2.56, which corresponds to the content of inactivated actin not higher than 2% (45), were used. Actin concentration was determined with a spectrophotometer (Hitachi, Japan). The molar extinction constant for actin was taken as \(E_{280} = 1.09 \text{ mg mL}^{-1} \text{ cm}^{-1}\) (46). Actin concentration was varied from 0.01 to 0.1 mg/mL. GdnHCl (Nacalai Tesque, Japan) was used without additional purification. The concentration of GdnHCl was determined by the refraction index with an Abbe refractometer (LOMO, Russia). In the experiments of refolding the appropriate concentrations of GdnHCl were obtained by dilution of actin solution in 5 M GdnHCl.

**Fluorescence Measurements.** Fluorescence experiments were carried out using the spectrofluorimeter with steady-state excitation (47). Fluorescence was excited at the long-wave absorption edge where the contribution of tyrosine residues is negligible. The position and form of the fluorescence spectra were characterized by the parameter \(A = (I_{320}/I_{365})_{297}\), where \(I_{320}\) and \(I_{365}\) are fluorescence intensities at \(\lambda_{em} = 320\) and 365 nm, respectively, and \(\lambda_{ex} = 297\) nm (47). In some cases the whole spectrum was recorded. The values of parameter \(A\) and of the fluorescence spectrum were corrected by the instrument sensitivity.

Vertical \((I_{V}^N)\) and horizontal \((I_{H}^N)\) components of fluorescence intensity excited by vertically polarized light were recorded for determination of fluorescence anisotropy \((r)\). The different sensitivity of the registering system for vertical and horizontal components was taken into account:

\[
r = \frac{I_{V}^N - G I_{H}^N}{I_{V}^N + 2G I_{H}^N}
\]

where \(G\) is the ratio of vertical and horizontal components of fluorescence intensity excited by horizontally polarized light \((G = I_{V}^H/I_{H}^V)\).

**Circular Dichroism Measurements.** Far-UV CD spectra were obtained with a dichrograph Mark V (Jobin-Yvon, USA) in a 0.1 cm path length cell. Actin concentration was 0.4 mg/mL. CD spectra of the appropriate buffers were recorded and subtracted from the protein spectra.

**Registration of the Kinetics of Actin Denaturation.** All kinetic experiments were performed in micro cells 101.016-Q5 5 × 5 mm (Hellma, Germany). Unfolding of the protein was initiated by manual mixing of the protein solution with buffer containing the corresponding amount of GdnHCl: 350 \(\mu\)L of the GdnHCl solution of appropriate concentration was injected in the cell with 50 \(\mu\)L of the solution of native protein. The dead time was determined from the control experiments. To this end 350 \(\mu\)L of buffer G was injected in the cell with 50 \(\mu\)L of the solution of native protein, and 350 \(\mu\)L of 4.0 M GdnHCl was injected in 50 \(\mu\)L of the solution of protein in the 4.0 M GdnHCl. These experiments showed that the dead time of kinetic measurements was less than 4 s. The final concentration of protein was 0.1–0.01 mg/mL. The character of the kinetic curves and the calculated values of rate constants \(k_1\), \(k_2\), and \(k_3\) (see below) were invariant of actin concentration within these limits. The spectrofluorimeter was equipped with thermostat that held a constant temperature of 23 °C in the cell and in the special box where the solutions are held before mixing.

The time courses of changes in the parameter \(A\) value were constructed using the experimental kinetic curves of fluorescence intensities successively registered at 320 and 365 nm. The kinetic curves of fluorescence anisotropy \((r)\) were calculated on the basis of the experimental kinetic curves of vertical \((I_{V}^N)\) and horizontal \((I_{H}^N)\) components of fluorescence intensity excited by vertically polarized light and registered successively.

**RESULTS**

To clarify the process of GdnHCl-induced formation of inactivated actin, kinetic curves reflecting the denaturant-induced changes in the intensity of intrinsic fluorescence were studied (see Figure 1A). The dependence of fluorescence intensity on the GdnHCl content after 24 h of incubation under conditions of desired denaturant concentration (Figure 1B) and complete spectra for native (N), inactivated (I), and unfolded (U) actin were registered (Figure 1B insert). Comparison of the fluorescence spectra in native, inactivated, and unfolded states shows that the greatest differences in fluorescence intensity are in the vicinity of 320 nm. That is why the wavelength 320 nm was used for the registration of the kinetic curves. Figure 1A shows that the shapes of the kinetic curves registered at various final GdnHCl concentrations differ significantly.

At low GdnHCl concentrations (below 1.0 M) the fluorescence intensity decreases monotonically with time, slowly approaching its equilibrium values (cf. Figure 1). At high GdnHCl concentrations (above 3.0 M) the intensity of actin intrinsic fluorescence also decreases monotonically and rapidly approaching the value typical of the completely unfolded protein (Figure 1A). The most intriguing results were obtained for moderate GdnHCl concentrations from 1.0 to 2.0 M. Within this concentration range, the fluorescence intensity first decreases over time but then increases slowly to the equilibrium value (Figure 1B).

The GdnHCl-induced change in the position and shape of the fluorescence spectrum over time was also studied. For this purpose, the changes in ratio of the intensities on two slopes of the spectrum, the so-called spectral parameter \(A = I_{320}/I_{365}\), were used (47). Figure 2 shows the time course of parameter \(A\) measured at 1.2 M GdnHCl. The insert to Figure 2 represents the dependence of the parameter \(A\) value on GdnHCl concentration determined after 24 h incubation of the protein at the desired denaturant concentration. It can be seen that early in the unfolding process the position and shape of the fluorescence spectrum tend to be similar to those typical of the completely unfolded protein. Then parameter \(A\) slowly increases and finally reaches the equilibrium value observed after 24 h incubation of the protein in the presence of 1.2 M GdnHCl.

Figure 3A shows the time courses of the fluorescence anisotropy changes, accompanying the unfolding of actin at 1.2 and 4.0 M final GdnHCl concentration, while the equilibrium fluorescence anisotropy values are shown in Figure 3B. Kinetic curves were calculated from the experimentally registered time courses of vertical and horizontal components of intrinsic fluorescence (Figure 3A insert).

Registering far-UV CD spectra also monitored the kinetics of actin structural changes induced by GdnHCl. Figure 4 shows the far-UV CD spectra of actin in 1.8 M GdnHCl.
measured just after mixing native actin with GdnHCl solution and overnight.

DISCUSSION

In all studies devoted to actin unfolding it has been assumed that actin successively transfers from native to inactivated and then to the completely unfolded state (29–41):

\[ \text{N} \rightarrow \text{I} \rightleftharpoons \text{U} \]  

(2)
i.e., inactivated actin (I) was considered as an on-pathway intermediate between the native (N) and completely unfolded (U) states. All equilibrium experiments appeared to support this model. In fact, data presented in Figures 1B and 3B and in the insert to Figure 2 suggest that GdnHCl induces two successive conformational transitions in actin. The transition from native to inactivated actin takes place at low GdnHCl concentrations (0.0–0.8 M), whereas the transformation of the inactivated actin into completely unfolded protein occurs between 1.8 and 4.0 M GdnHCl. It also suggests that in the relatively wide range of GdnHCl concentrations, between 0.8 and 1.8 M, actin exist in its inactivated form.
Pathway of Actin Unfolding

The minimum seen in the kinetic curves at 1.0, 1.2, 1.5, and 1.8 M GdnHCl suggests that the transition from the native to the inactivated state occurs via some intermediate state in which the fluorescence intensity is lower than in the native and the inactivated state (Figure 1A). It is known that actin in the unfolded state but whose secondary structure is much more ordered. The transition from U* to I is most likely to be irreversible. Nonetheless, we regard the more general situation taking into account $k_3$. At GdnHCl concentration greater than 1.8 M the scheme must take into account the reversible transition between I and U. In this work this transition is not regarded. In the frame of the kinetic scheme (eq 3) the time dependencies of the protein fraction in native $\alpha_N(t)$, inactivated $\alpha_U(t)$, and essentially unfolded $\alpha_{U*}(t)$ states are

$$\alpha_N(t) = \frac{[N(t)]}{[N(0)]} = e^{-k_1 t}$$

$$\alpha_U(t) = \frac{[U^*(t)]}{[N(0)]} = \left[(k_2 + k_3)k_1 + k_3 \right] e^{-k_1 t} - k_3 k_2 + k_3 k_1 - k_3^2 + k_2 k_3 e^{-(k_2 + k_3) t} \right] e^{(k_1 + k_2) t} (k_1 - k_2 - k_3)$$

$$\alpha_{U*}(t) = \frac{[I(t)]}{[N(0)]} = -k_2 k_3 e^{-(k_2 + k_3) t} - (k_1 + k_2) e^{-k_1 t} - k_1 + k_2 + k_3) (k_2 + k_3) (k_1 - k_2 - k_3)$$

The fitting routine to analyze the kinetic curves of the relative fluorescent intensity and to determine the rate constants $k_1$ and the fraction of the protein in the native, inactivated, and essentially unfolded states was based on the nonlinear least-squares method.

Analysis of the Actin Unfolding Kinetics. The time dependence of fluorescent intensity of the system containing actin in the native, inactivated, and essentially unfolded state is determined by the equation:

$$I(t) = \alpha_N(t)I_N + \alpha_I(t)I_I + \alpha_{U*}(t)I_{U*}$$

where $\alpha_N(t)$, $\alpha_I(t)$, and $\alpha_{U*}(t)$ are the fraction of the protein in the native, inactivated, and essentially unfolded kinetic intermediate states; $I_N$, $I_I$, and $I_{U*}$ are fluorescence intensities of actin in these states, respectively; and $\alpha_N(t) + \alpha_I(t) + \alpha_{U*}(t)$...
of 1.2 M. The best fit of the experimental data was achieved by the concrete kinetic scheme that fits the unfolding process (see above). The values of rate constants \( k_i \) and the calculated best fit of the fluorescence intensity that correspond to the values \( k_1 = 2.1 \times 10^{-2}, k_2 = 7.2 \times 10^{-3}, \) and \( k_3 = 7.6 \times 10^{-4} \) s\(^{-1}\). This figure represents also the changes in the populations of native (\( \alpha_0 \)), inactivated (\( \alpha_i \)), and essentially unfolded (\( \alpha_{\text{U}^*} \)) actin molecules, calculated according to eq 4. Figure 5B shows that the residuals are of statistical character, though there is a small nonlinearity for short times. The proximity of chi square \( \chi^2 = \Phi(n - p) \), where \( n \) is the number of experimental points and \( p \) is the number of examined parameters] to the unity \( \chi^2 = 1.12 \) and the randomness of the residuals prove the validity of the chosen kinetic model.

GdnHCl Dependence of the Rate Constants \( k_i \). The GdnHCl dependencies of the rate constants \( k_1, k_2, \) and \( k_3 \) were determined (Figure 6). As one would expect, \( k_3 \) is practically equal to zero. At small denaturant concentration \( k_2 \) is considerably greater than \( k_1 \). Therefore, the limiting stage of the reaction under these conditions is protein unfolding, whereas all unfolded molecules will immediately refold to form the inactivated actin. With the increase in GdnHCl concentration the rate constant \( k_1 \) increased and the rate constant \( k_2 \) decreased. This leads to the accumulation of essentially unfolded molecules early in the unfolding process (in the range of 1.0–2.0 M GdnHCl) and, consequently, to the appearance of the characteristic minimum in the kinetic curves (Figures 1–3).

Thus, our data show that inactivated actin does not represent an intermediate state in the pathway of this protein unfolding from the native to the unfolded state, as has been suggested earlier. On the contrary, this conformation should be considered as an off-pathway species, which appears via the primary unfolding of the protein. This result may also be essential for the search of the pathways of actin folding in vitro.

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REFERENCES


![Figure 5: Analysis of the kinetic curves of actin denaturation monitored by the change of intrinsic fluorescence intensity.](image-url)