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Suppression of thermal denaturation of myosin and salt-induced denaturation of actin by sodium citrate in carp (Cyprinus carpio)

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The effect of sodium citrate (Na-citrate) on myosin and actin denaturation in myofibrils was investigated. Na-citrate significantly suppressed the thermal inactivation of Ca$^{2+}$-ATPase of carp myosin in a concentration dependent manner. The effect was greater than that of sorbitol. A similar effect was observed with myofibrils in which myosin is stabilized by F-actin binding. Na-citrate dissolved myofibrils at lower concentration than NaCl. Nevertheless, Na-citrate at 1 M never denatured F-actin in myofibrils, while 1 M NaCl denatured F-actin almost completely. Na-citrate suppressed the NaCl-induced F-actin denaturation. Sorbitol did not show such protective effect on F-actin denaturation. Moreover, Na-citrate suppressed the freeze denaturation of myofibrils at lower concentration than sorbitol. Thus, Na-citrate was proved to be superior to sorbitol. It was suggested that Na-citrate alone could substitute sorbitol as cryoprotectant in Surimi and NaCl as dissolving reagent of myofibril in thermal gel production.

**Keywords:**
Sodium citrate (Na-citrate)
Myosin
Actin
Denaturation
Sorbitol
1. Introduction

Myosin is the most important component in thermal gel formation from fish and land animal meat (Samejima, Hashimoto, Yasui & Fukazawa, 1969; Kawakami, Morita, Takahashi & Yasui, 1971; Yasui, Ishioroshi & Samejima, 1980; Samejima, Ishioroshi & Yasui, 1981). It is also well accepted that myosin denaturation deteriorates the quality of the final gel products. Thus, there are many attempts to prevent myosin denaturation during the storage of meat or raw material. The satisfactory technology is surimi processing from Alaska Pollock, in which sugar or sugar alcohol was added to washed and dewatered fish meat as cryoprotectant to prevent myosin denaturation during the frozen storage (Arai, Takahashi & Saito, 1970; Kawashima, Arai & Saito, 1973). It is believed that addition of sugar is essential for surimi production because of unstable nature of its Alaska Pollock myosin. However, addition of sugars to surimi gives sweetness although using sorbitol instead of sucrose reduces the sweetness. It is also reported that the other compounds, such as sodium (Na-) glutamate or Na-acetate suppress myosin denaturation (Noguchi & Matsumoto, 1970; 1971; Ooizumi, Yamamura & Arai, 1982; Ooizumi, Nara & Arai, 1984; Matsuura, Ooizumi & Arai, 1984). Strong taste of Na-glutamate and weak suppressive effect of Na-acetate are their demerit. It is assumed that stabilizing effect of Na-acetate relied on carboxyl group in the structure (Ooizumi et al., 1984). Since Na-citrate is practically tasteless and its structure contains three carboxyl groups, its potential to suppress myosin denaturation is highly probable.

There is an accumulation of reports that focused on the suppression of myosin
denaturation by additives. In contrast, there is little report on the protection of F-actin denaturation by additives. This is probably because F-actin in myofibrils is very stable compared to myosin. However, a unique F-actin denaturation upon addition of neutral salt was reported by Wakameda and Arai (1984). Treatment of actomyosin or myofibrils with NaCl or KCl at around 1.5 - 2 M selectively denatures F-actin even at low temperature without accompanying myosin denaturation (Wakameda, Nozawa & Arai, 1983). Since Na-citrate effectively dissolved squid myofibrils at lower concentrations than NaCl (Kuwahara, Osako, Okamoto & Konno, 2006), Na-citrate behaved as strong salt. It is probable that Na-citrate as salt denatures F-actin in myofibrils like NaCl.

In order to apply Na-citrate to Surimi production and thermal gel formation, we investigated how Na-citrate addition affects thermal denaturation of myosin in the detached form or bound form to F-actin and F-actin denaturation induced by heating and salt-treatment of myofibrils

2. Materials and methods

2.1. Materials

Dorsal muscle was harvested from live carp *Cyprinus carpio*. All chemicals were of analytical grade and purchased from Wako Pure Chemical Industries (Osaka, Japan).

2.2. Preparation of myofibrils and myosin
Myofibrils were prepared from dorsal muscle of carp according to the method of Katoh, Nozaki, Komatsu and Arai (1979). Muscle mince was homogenized for 30 sec 4 times at cooling with ice-cold water on Polytron homogenizer PT 10-35 (Kinematica, Littau-Lucerne, Switzerland) in chilled buffer consisting of 0.1 M NaCl and 20 mM Tris-HCl (pH 7.5). The homogenate was repeatedly washed by centrifugation at 3000 xg for 5 min at 4 ºC and suspension of the pellet. Finally, the filtrate through two layers of gauze cloth was used as myofibril preparation. Myosin was isolated from thus prepared myofibrils by the method described by Koseki, Kato & Konno (1993). Myofibrils dissolved in 0.5 M NaCl, 20 mM Tris-HCl (pH 7.5) was added with 5mM Mg-ATP to dissociate myosin from actin, then saturated ammonium sulfate was added to give 40 % saturation to remove actin as precipitate. Myosin in the supernatant was collected by raising the saturation to 60 %. Myosin dissolved in and dialyzed against 0.5 M NaCl, 20 mM Tris-HCl (pH 7.5) and the supernatant after centrifugation at 20000 xg for 20 min at 4 ºC was used as myosin preparation.

2. 3. Analysis of Ca$^{2+}$-ATPase activity

Myosin denaturation was assessed by Ca$^{2+}$-ATPase inactivation analysis. Although Na-citrate in protein solution reduced the Ca$^{2+}$-ATPase activity by chelating Ca ion in the reaction mixture, we confirmed that increased CaCl$_2$ concentration to 15 mM removed the problem. We also confirmed that inclusion of Na-citrate up to 0.1 M in the assay medium
unaffected the ATPase inactivation rate measurement. Accordingly, the ATPase was assayed in a medium containing, 0.5 M NaCl, 15 mM CaCl₂, 1 mM ATP, 25 mM Tris-maleate (pH 7.0), and myofibrils at about 0.1 mg/ml. Thermal inactivation rate \((k)\) was calculated by assuming that the inactivation obeys the first order reaction mechanism. To analyze the suppressive effect of compounds quantitatively, thermal inactivation rates in logarithmic values were plotted against the concentrations of the compound added, and the slope of the straight line obtained was defined as E-value as Ooizumi, Hashimoto, Ogura & Arai (1981) proposed.

2. 4. Thermal and freeze denaturation of carp myosin and myofibrils

For thermal denaturation, myosin and myofibrils were heated at 35 and 42 °C, respectively. Myofibrils were frozen stored at -20 °C for 30 days to induce freeze denaturation. Na-citrate or other compounds were added to the above protein solution for studying their suppressive effect. As references, stabilizing effect of Na-glutamate, Na-acetate and sorbitol were also examined (Ooizumi et al., 1981). Myosin denaturation was assessed by Ca²⁺-ATPase inactivation analysis.

2. 5. F-actin denaturation and Salt-induced actin denaturation

F-actin denaturation upon treatment of myofibrils with Na-citrate was studied by analyzing the Ca²⁺-ATPase thermal inactivation profile of the treated myofibrils upon
subsequent heating. Carp myofibrils were mixed with 0.25 or 1.0 M Na-citrate and incubated overnight on ice. Before the heating of 35 °C, Na-citrate concentration was diluted 3 times and NaCl concentration was adjusted to 0.5 M. Their inactivation profiles were assessed by Ca^{2+}-ATPase inactivation. Salt-induced F-actin denaturation upon treatment of myofibrils with Na-citrate was studied by analyzing the Ca^{2+}-ATPase inactivation profile of the treated myofibrils upon subsequent heating. Carp myofibrils were treated with 1.0 M NaCl together with 0, 0.1, 0.25, and 0.4 M Na-citrate in ice cold water overnight. For the analysis of F-actin denaturation upon treatment, the salt concentration of the medium was adjusted to 0.5 M NaCl and 0.25 M Na-citrate by 3 times dilution with 20 mM Tris-HCl (pH 7.5) at which actin denaturation by NaCl was negligible. Thus treated myofibrils were heated at 35 °C and their inactivation profile was compared with that of untreated myofibrils under the same conditions (Yasui, Kawakami & Morita, 1968; Yamashita, Arai & Nishita, 1978; Takahashi et al., 2005).

2. 6. Solubilization of muscle protein

Carp myofibrils in 0.1 M NaCl, 20 mM Tris-HCl (pH 7.5) were added by various concentrations of NaCl or Na-citrate in ice cold water for 30 min. The mixture was centrifuged at 20000 xg for 30 min at 4 °C. Protein content recovered in the supernatant relative to one before centrifugation was the index of solubilization (Konno, Yamanodera & Kiuchi, 1997). To measure protein concentration of the supernatant, solubilized proteins was precipitated by centrifugation at 4000 rpm for 30 min at 4 °C with 7.5 % trichloroacetic
acid, and the pellet was dissolved again with 1 M NaOH. Protein concentration was estimated by using Biuret reagent (Gornall, Bardawill & David, 1949).

To confirm the result, we repeated the same set of experiment at least three times by using different preparations, and results were presented only when the same conclusion was made by the separate experiment.

3. Results and discussion

3.1. Stabilizing effect of Na-citrate on carp myosin

First, to estimate the potential stabilizing ability of Na-citrate on myosin, its effect on myosin itself was studied. Myosin in 0.5 M NaCl was heated at 35 °C with various concentrations of Na-citrate and thermal inactivation rates of Ca\(^{2+}\)-ATPase were estimated. Although Na-citrate in myosin solution reduced the activity by chelating Ca\(^{2+}\) in the reaction mixture, we confirmed that increased CaCl\(_2\) concentration to 15 mM removed the problem. We also confirmed that inclusion of Na-citrate up to 0.1 M in the assay medium unaffected the ATPase inactivation rate measurement. For a quantitative analysis of the suppressive effects, thermal inactivation rates in logarithmic values were plotted as a function of the concentrations used according to Ooizumi et al., (1981) (Fig. 1). The plotting for all compounds including Na-citrate gave straight lines with different slopes. The slope of the relationship for Na-citrate was steeper than those obtained with sorbitol or Na-acetate, and similar to that obtained with Na-glutamate. The results indicated that
Na-citrate more effectively suppressed myosin denaturation than sorbitol, most popular and widely used additive. A quantitative comparison of the suppressive effects among compounds was made by using E-value. The E-value for Na-citrate as well as Na-glutamate (1.7) was greater than that for sorbitol (1.1) and Na-acetate (0.7). The E-value of Na-citrate indicated that inclusion of 1 M Na-citrate stabilizes myosin by $10^{1.7}$ times (about 50 times), while the extent achieved by sorbitol is $10^{1.1}$ (about 13 times). Stabilizing effect of Na-citrate was found to be superior to sorbitol. As Na-glutamate with E-value of 1.7 was reported to show the largest stabilizing effect among compounds studied (Ooizumi et al., 1981), stabilizing effect of Na-citrate was comparable to it.

3.2 Stabilizing effect of Na-citrate on myofibrils denaturation

We further studied the suppressive effect of Na-citrate on thermal denaturation of myosin in myofibrils, namely myosin in a stabilized form through F-actin binding. The same set of experiment was conducted by using myofibrils at raised temperature of 42 °C. The rates were estimated at varied concentrations of Na-citrate and their logarithmic values were plotted against the concentration (closed circles in Fig. 1). Again, the plotting gave a straight line. E-value calculated from the slope was 1.4 which was smaller than the ones obtained with myosin alone (1.7). Generally, E-values obtained with myofibrils are smaller than those obtained with myosin for any kind of compounds (Hayashi & Konno, 2006). Probably low concentration of NaCl such as 0.1 M contained in the myofibrils accelerated the inactivation slightly through reduced protection by F-actin. Practically Na-citrate
showed a similar suppressive effect on myofibrils as on myosin alone. The result was somehow different from the case of other organic salt such as Na-glutamate or Na-acetate because their suppressive effect on myofibrils was remarkably low compared with that on myosin subfragment-1 (Hayashi & Konno, 2006).

3. 3. Dissolving myofibrils upon addition of Na-citrate

NaCl at 1.5 - 2 M destabilizes myosin as a result of removal of the protection by F-actin (Wakameda et al., 1983; 1984). We found that Na-citrate at 1 M did not accelerate myosin denaturation when heated as myofibrils and rather remarkably stabilized myosin in myofibrils. The fact suggested that Na-citrate did not act as ionic compound, and acted as nonionic compound. To examine the ionic properties of Na-citrate as salt, its dissolving ability of myofibrils was compared with that of NaCl. NaCl at 0.25 M dissolved myofibrillar proteins almost completely. Na-citrate also dissolved myofibrils and the concentration required was as low as 0.05 M (Fig. 2). It was concluded that Na-citrate surely acted as ionic compound. The lower concentration of Na-citrate than NaCl for dissolving indicated a much higher ionic strength at the same molar concentration. This might be reasonable because citrate is a trivalent carboxylic acid giving five times greater ionic strength than NaCl at the same molar concentration by assuming a complete dissociation of hydrogen ion from the three carboxyl groups.

3. 4. Effect of Na-citrate on F-actin denaturation
Na-citrate acted as a strong salt in dissolving myofibril. Nevertheless, there was no trace of F-actin denaturation by Na-citrate addition (Fig. 1). We further examined whether the F-actin denaturation by Na-citrate occurs. The method to detect F-actin denaturation is constructed based on the following facts, 1) selective F-actin denaturation in myofibrils induces myosin detachment from F-actin, 2) myosin that lost the protection by F-actin is very unstable, and 3) inactivation rate of detached myosin and one of F-actin bound myosin is distinguished easily by the inactivation profile of Ca$^{2+}$-ATPase (Murozuka & Arai, 1976; Takahashi, Yamamoto, Kato & Konno, 2005). Carp myofibrils were mixed with 0.25 or 1 M Na-citrate and incubated overnight on ice. F-actin denaturation in the myofibrils was judged by the existence of free myosin detached from denatured F-actin by the treatment. Existence of the detached myosin was detected by studying the Ca$^{2+}$-ATPase inactivation profile of the myofibrils, namely an appearance of quick inactivation phase in the profile is the evidence of the existence of detached myosin. Their inactivation profiles are presented in Fig. 3. Irrespective of the concentrations of Na-citrate used, there was no difference in the inactivation profiles between treated and untreated myofibrils. The profile with untreated myofibrils represented the denaturation profile of F-actin bound myosin. Thus treated myofibrils with Na-citrate contained no free myosin. In other words, the salt did not denature F-actin. A slightly shallow slope for the sample treated with 1 M Na-citrate was due to different Na-citrate concentrations in the heating medium. These are 0.33 M (myofibrils treated with 1 M) and 0.083 M (myofibrils treated with 0.25 M), respectively. The results clearly demonstrated that Na-citrate never caused F-actin denaturation.
3. 5. Protection of F-actin from salt-induced denaturation by Na-citrate

The next attempt was to test the possibility whether Na-citrate suppresses NaCl-induced F-actin denaturation. For the purpose, myofibrils were incubated with 1 M NaCl overnight on ice together with 0, 0.1, 0.25 or 0.4 M Na-citrate. The method to detect F-actin denaturation was the same as used in Fig. 3, namely detecting free myosin in the medium as a consequence of F-actin denaturation. NaCl concentration for the treated myofibrils was dilution to 0.5 M so as to stop the NaCl-induced F-actin denaturation. Furthermore Na-citrate concentration in the heating medium was adjusted to 0.25 M to make the comparison of the inactivation rate easier among samples. Thermal inactivation profiles of the samples at 35 °C are shown in Fig. 4. Myofibrils without NaCl-treatment exhibited a slow inactivation representing the inactivation profile of myosin stabilized by F-actin (open circles). In contrast, myofibrils treated with 1 M NaCl exhibited a very quick inactivation phase implying the existence of very unstable free myosin in the medium (closed circles in Fig. 4). The result obtained was a confirmation that 1 M NaCl denatured F-actin selectively in myofibrils. Na-citrate at 0.1 M practically unaffected the inactivation profile (triangles in Fig. 4). Inclusion of Na-citrate at 0.25 M changed the profile (squares in Fig. 4). The profile showed a breaking point indicating the presence of two species of myosin with different stabilities in the heating medium. Inactivation by unstable myosin contributed mainly in an early phase, while the stable myosin determined the rate in latter phase. Unstable and stable species are myosin detached from F-actin and one bound to F-actin. The phenomenon is
often observed when salt-induced F-actin denaturation is partial. A little slower inactivation in an early phase was due to a partial contribution of the rate of latter phase. Thus, a partial prevention of F-actin denaturation by N-citrate was proved. Amount of F-actin bound myosin could be calculated by extrapolating the line of the latter phase to heating time zero. Relative remaining activity at the point was calculated to be roughly -0.7 in logarithmic scale, which corresponds to 0.5 in ordinary scale. The calculation indicated that roughly a half of myosin in the medium was present in F-actin bound form. When 0.4 M Na-citrate was added, a quick inactivation phase disappeared from the profile and gave the same inactivation profile as untreated myofibrils (diamonds in Fig. 4). The result showed that all of myosin in the medium was in F-actin bound form, or F-actin denaturation by NaCl was completely suppressed by Na-citrate. Na-citrate at 0.6 M gave the same result as 0.4 M (data not shown). Na-citrate at 0.4 M was proved to be high enough to prevent F-actin denaturation completely.

As the suppressive effect of Na-citrate on NaCl-induced F-actin denaturation was evident, it is reasonable to think that other compounds that suppressed myosin denaturation might suppress the F-actin denaturation similarly. We tested sorbitol. The same set of experiments as done with Na-citrate was conducted by replacing Na-citrate with sorbitol. Sorbitol concentration used was 0.82 M, which corresponds to 0.35 M Na-citrate in stabilizing myosin denaturation (see Fig. 1). Results are presented in Fig. 5. Incubation of myofibrils with 1 M NaCl alone generated a quick inactivation phase (closed circles in Fig. 5). Inclusion of 0.82 M sorbitol unchanged the inactivation profile showing a quick inactivation phase in the profile (triangles in Fig. 5). It was concluded that sorbitol did not
suppress NaCl-induced F-actin denaturation. Electrostatic interaction of Na-citrate to
F-actin seemed important to stabilize actin structure. Stabilizing mechanism of Na-citrate
on F-actin is uncertain at present but practically Na-citrate might be superior additive to
sorbitol by its large E-value on myosin denaturation and protective effect on F-actin
denaturation.

3. 6. Freeze denaturation of myofibrils and its suppression by Na-citrate

Above results clearly demonstrated that Na-citrate strongly protects myosin both in free
form and bound form to F-actin and that it protected salt-induced F-actin denaturation.
Another well known factor to cause myosin denaturation is freezing. It is well established
that sugars and Na-glutamate well suppress freeze denaturation of myosin (Arai et al.,
1970; Kawashima et al., 1973; Ooizumi et al., 1984). Recently, we reported that ionic
compounds such as sulfate ion rather accelerated the freeze denaturation of myosin in spite
of the fact that the ion strongly suppressed the thermal denaturation of myosin (Torigai &
Konno, 1996; Konno, 1998; Hayashi & Konno, 2006). It is also reported that NaCl in
actomyosin solution promotes F-actin denaturation during the frozen storage at around -10
°C (Takatori, Inoue & Shinano, 1992). As Na-citrate behaved as strong ionic compound by
dissolving myofibrils very effectively, it is still possible that Na-citrate causes F-actin
denaturation during the frozen storage of myofibrils, subsequently promotes myosin
denaturation.

Carp myofibrils in 0.1 M NaCl pH 7.5 was mixed with various concentrations of
Na-citrate and were frozen stored at -20 °C. Sorbitol, Na-glutamate and Na-acetate were also used as references (Ooizumi et al, 1984; Matsumoto, Ooizumi & Arai, 1985). The highest concentration was 200 mM because it is reported that freeze denaturation requires lower concentration for protection (Matsumoto et al., 1985). The remaining Ca$^{2+}$-ATPase activities of the myofibrils after storage for a month were measured to assess the freeze denaturation of myosin. Activity dropped to less than 10% that of original value when stored without any additives. With increasing the concentration of Na-citrate added, the remaining activity increased. Na-citrate at 25 mM remarkably suppressed the activity drop and 50 mM was sufficient to keep the activity almost unchanged. The suppressive effect of sorbitol was similar to or less than that of Na-citrate. Sorbitol at 50 mM gave the highest remaining activity. Na-glutamate also effectively suppressed the inactivation at lower concentration range showing the maximal remaining activity at 50 mM. Na-acetate weakly suppressed the inactivation requiring 150 mM to give the maximal remaining activity. Thus, Na-citrate was proved to be an excellent compound to prevent freeze denaturation of myosin. Its effect was similar to or better than that of sorbitol (Fig. 6).

Na-citrate functions as a very strong salt in dissolving myofibrils. Nevertheless, it never caused salt-induced F-actin denaturation. Being different from inorganic ion, organic salts including Na-citrate, Na-glutamate, and Na-acetate were proved to suppress freeze denaturation of myosin as well as its thermal denaturation (Hayashi, Azuma, Koseki, & Konno 2007).

In thermal gel formation by surimi, it is essential to dissolve myofibrillar protein prior to heating, termed as salting process. As Na-citrate well dissolved carp myofibrils like NaCl,
the salting with Na-citrate could be possible without using NaCl. As Na-citrate is practically tasteless, thermal gel without salty taste can be produced by replacing NaCl with Na-citrate. Frozen surimi as a raw material for gel production usually contains sugar or sugar alcohol as cryoprotectant. As Na-citrate very strongly suppressed myosin denaturation in myofibrils during the frozen storage, protection of myosin denaturation by replacing sugar with Na-citrate as cryoprotectant could be possible. The use of Na-citrate as a cryoprotectant makes it possible to produce surimi without sweet taste. By using the surimi, it becomes possible to produce the thermal gel without salty or sweet taste. Such surimi will expand the application area in food industry.

Supposing that Na-citrate at 50 mM was added to leached fish meat, spontaneous dissolving of muscle protein is expected (Fig. 2), namely surimi would be transformed to sol (dissolved surimi). Such surimi would be categorized into salt-added surimi. In Japan, NaCl-added surimi in small quantity is produced by the special request of manufacturers. However, the surimi is not distributed widely because of its short shelf-life due to unstable nature of myosin in the surimi as a consequence of loss of protection by F-actin. As Na-citrate well protected myosin from freeze denaturation, a new type of frozen surimi, “salt-added surimi” would become practical. Study on the practical application of Na-citrate in frozen surimi production is now on going.

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Figure captions:

Fig. 1. Suppressive effect of Na-citrate on thermal denaturation of myosin and on myofibrils. Carp myosin (open symbols) and myofibrils (closed symbols and dashed line) were heated at 35 ºC and 42 ºC, respectively in the presence of various concentrations of Na-citrate (circles), Na-glutamate (squares), Na-acetate (diamonds), and sorbitol (triangles).

Fig. 2. Solubilization of carp myofibrils with Na-citrate. Carp myofibrils (0.1 M NaCl) were incubated with additional concentrations of NaCl (cross marks) or Na-citrate (circles). Relative protein content in the supernatant obtained by centrifugation of the mixture to whole protein used was expressed as percent dissolved protein.

Fig. 3. Effect of Na-citrate on the F-actin denaturation in carp myofibrils. Carp myofibrils (0.1 M NaCl) were incubated with 0.25 M (circles) or 1 M (squares) Na-citrate on ice overnight. ATPase inactivation profile at 35 ºC for the treated myofibrils (closed symbols) after adjusting the NaCl concentration to 0.5 M was compared with those without treatment (open symbols).

Fig. 4. Suppressive effect of Na-citrate on the NaCl-induced F-actin denaturation in carp myofibrils. Carp myofibrils were incubated with 1 M NaCl together with 0 M (circles), 0.1 M (triangels), 0.25 M (squares), and 0.4 M (diamonds) Na-citrate on ice overnight. ATPase inactivation profiles for thus treated myofibrils (closed symbols) after reducing NaCl
concentration to 0.5 M were analyzed. The sample without treatment was also shown (open circles with dashed line).

Fig. 5. Effect of sorbitol on the salt-induced F-actin denaturation in carp myofibrils. Carp myofibrils were treated with 1 M NaCl together with either 0 M (circles) or 0.82 M sorbitol (triangles). Other procedures were the same as in Fig. 4. Treated myofibrils (closed symbols) and untreated myofibrils (open symbols) were compared.

Fig. 6. Suppressive effect of Na-citrate on the freeze denaturation of carp myofibrils. Carp myofibrils (0.1 M NaCl, 20 mM Tris-HCl (pH 7.5)) containing various compounds at various concentrations were frozen stored at -20 °C for 30 days. The compounds used were Na-citrate (closed circles), Na-glutamate (squares), sorbitol (open circles), and Na-acetate (triangles).
Fig. 1

Log $k$ (relative) vs. Addivites (M)
Fig. 2

Dissolved protein (%)

Additives (M)
Fig. 3

Ln Remaining ATPase (relative) vs. Incubation time (min)
Fig. 4

Ln Remaining ATPase (relative) vs. Incubation time (min)

-6.0 -5.5 -5.0 -4.5
0 30 60 90 120
Fig. 5

Ln Remaining ATPase (relative) vs. Incubation time (min)

-6.0 -5.5 -5.0 -4.5 -4.0 -3.5 -3.0 -2.5 -2.0 -1.5 -1.0 -0.5 -0.0 0.5 1.0
0 30 60 90 120