Proactive effective of exogenous adenosine triphosphate on hypothermically preserved rat liver

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INTRODUCTION

Improving the quality of cold stored organs and prolonging the effective preservation time are the pivotal contents in the investigation of hypothermic preservation of transplant grafts. Several investigators have reported that the intracellular level of adenosine triphosphate (ATP) in cold stored organs was closely correlated with the viability of transplant grafts[1-3]. Bowers reported that ATP level in cold stored pretransplant organs was a sensitive parameter for examining the activities of cold stored organs[4]. Therefore, providing direct energy substrate ATP to cold stored organs[5] should be a simple and effective method to sustain the high level of intracellular ATP.

However, thus far, whether exogenous ATP could enter cells or not is controversial[6-9]. Furthermore, there were few reports which elucidated the protective effect of ATP on cold stored transplant grafts. In this study, a continuously hypothermic machine perfusion model of rat liver was applied to reveal the protective effect of ATP on cold stored rat livers and its mechanism.

MATERIALS AND METHODS

Experimental Animals

Wistar rats weighing 180-220 g, both male and female, were randomly used.

Experiment groups and protocol

Cold storage study on rat livers The rats mentioned above were divided into 3 groups at random, group A (containing neither ATP nor MgCl₂ in the perfusate), group B (containing 5 mmol/L ATP but no MgCl₂ in the perfusate), group C (containing either ATP or MgCl₂ in the perfusate), respectively. There were 6 rats in each group. The rat liver was weighed immediately after resection by the method described previously[10-12], then these grafts were put into the modified Hoffmann perfusate[13] (0-4 °C) for 30 min (Table 1). Finally, the livers were preserved in a hypothermic preservation incubator by continuously hypothermic preservation perfusion model (Figure 1). The perfusate temperature was 6-8 °C[14], perfusion speed was 0.1 mL/min/g[15], the total volume of perfusate was 120 mL.

 Autoradiography study Six rats were chosen randomly, the livers were resected with the same method. One mCi [α-³²P] ATP, 5 mmol/L MgCl₂, 200 µL and 40 U phenol kinase were added into 1 L perfusate, and the same liver preservation mechanism was applied.

Table 1 Components of perfusate

| Component                  | Concentration |
|----------------------------|---------------|
| Hydroxyethyl starch        | 50 g/L        |
| Calcium gluconate          | 80 mmol/L     |
| Raffinose                  | 10 mmol/L     |
| K₂HPO₄                     | 25 mmol/L     |
| Hydroxyethyl piperazine    | 10 mmol/L     |
| Dexamethasone              | 12 mg/L       |
| Penicillin                 | 2×10⁵ units/ L|
| Insulin                    | 100 units/ L  |
| MgCl₂                      | 5 mmol/L      |
| ATP                        | 5 mmol/L      |

The pH value was modulated to 7.35 with NaOH, and the osmotic pressure was 300-320 mOsm/L; addition of MgCl₂ and ATP was dependent on the different groups.

Biochemical detection

Detection of energy status in cold stored rat livers The rat liver samples were used to detect the intracellular ATP, ADP, AMP, TAN and EC at 0, 1, 2, 6, 12, 24 and 36 h after resection.
preservation by HPLC method\(^{[15]}\) (TAN=ATP+ADP+AMP, EC=[ATP+0.5ADP]/TAN\(^{[16]}\)). One milliliter of perfusate was taken to detect the LDH and AST activities\(^{[17,18]}\) at 6, 12, 24 and 36 h respectively after preservation.

**Histological and morphological findings** Paraffin sections of HE staining were made after 24 h preservation of the rat livers\(^{[19]}\), and observed by a light microscope.

**Autoradiography of [\(\alpha\)-\(32\)P] ATP** The rat liver samples were made into paraffin sections of HE staining after 4-h preservation. Moreover, whether ATP entered the cells of cold stored rat livers or not was examined by autoradiography of [\(\alpha\)-\(32\)P] ATP\(^{[20]}\).

**Statistical analysis**
The average values were presented as mean±SD, \(t\)-test was applied and \(P<0.05\) was considered to be statistically significant.

**RESULTS**

**Energy status in cold stored rat livers (\(\mu\)mol/g wet liver)**
In group A, following the prolongation of preservation time, the ATP and EC levels in rat liver cells were significantly decreased. The ATP and EC levels were also rapidly decreased in group B, there was no statistical difference between these two groups \((P>0.05)\). However, the ATP and EC levels were slowly decreased in group C \((P<0.01, \text{Table 2})\).

**LDH and AST activities in hypothermic preservation perfusate**
LDH and AST activities in the perfusate were increased in groups A and B, there was no significant difference between two groups \((P>0.05)\). On the other hand, compared with those in groups A and B, the relevant activities were slowly increased in group C \((P<0.05, \text{Table 3})\).

**Histological and morphological findings after 24-h hypothermic perfusion preservation**
In group A (Figure 2), the hepatocytes were obviously swollen, cytosol and part of nucleus were faintly stained. Part of the endothelial cells entered the hepatosinus.

In group B (Figure 3), the hepatocytes were also expanded, cytosol was faintly stained. Some nuclei were strongly stained. Some endothelial cells entering the hypatosinus were also found.

In group C (Figure 4), the hepatocytes were lightly expanded. There was no apparent bubble in cytosol, and the morphology of nucleus was normal. The endothelial cells of hepatosinus were continuous.

**Figure 1** Preservation of continuously hypothermic machine perfusion. A: Organ hypothermic preservation box, B: Temperature displayer, C: Perfusion pump, D: Perfusion pump, (Displaying the perfusion pressure), E: Perfusate container, F: Organ preservation container, G: pH displayer, H: Entry of wire, (Being sealed while preservation).

**Figure 2** Histological and morphological findings in hypothermically preserved hepatocytes. Group A: 24-h preservation, HE staining 100×.

**Figure 3** Histological and morphological findings in hypothermically preserved hepatocytes. Group B: 24-h preservation, HE staining 100×.

**Figure 4** Histological and morphological findings in hypothermically preserved hepatocytes. Group C: 24-h preservation, HE staining 100×.

**Figure 5** Autoradiography of [\(\alpha\)-\(32\)P] ATP in hypothermically preserved hepatocytes. Black spots in hepatocytes are the autoradiographies of [\(\alpha\)-\(32\)P] ATP, 4-h preservation, HE staining 100×.
Autoradiography of [α-32P] ATP
Numerous silver spots of [α-32P] ATP were found to be limited within the rat hepatocytes, while no silver spots were found in the hepatosinus and central vein (Figure 5). This observation demonstrated that [α-32P] ATP entered the cold stored rat hepatocytes.

DISCUSSION

Protective effects of exogenous ATP on hypothermically preserved rat livers

Up to now, some reports have revealed that ATP-MgCl2 had a protective effect on the therapy of hemorrhagic shock27,28, but reports revealing the protective effect of ATP-MgCl2 on hypothermically preserved transplant organs were few29,30. The results of this study demonstrated that the intracellular level of ATP in the group containing no ATP-MgCl2 or the group containing ATP alone decreased rapidly after hypothermic preservation. Simultaneously, the release of intracellular enzymes was increased, indicating severe damages of the membrane functions. Moreover, significant swelling of the hepatocytes and obvious infiltration of neutrophils were found histologically. On the contrary, the intracellular ATP level in the group containing ATP-MgCl2 was almost maintained at the normal level for quite a long time, and decreased much slower after hypothermic preservation. Furthermore, because of the protective effect of ATP on cell membranes23,24, the metabolic function of hepatocytes was restored, and the release of intracellular enzymes (LDH and AST) was significantly inhibited. The histological observations also showed that the swelling of hepatocytes was milder than that in groups B and C. These results suggested that ATP-MgCl2 could directly provide the energy or energy substrates for intracellular Na+-K+ ATPase as well as Ca2+-ATPase to remain the extracellular and intracellular ion balance25-29, and lighten the intracellular acidosis and cell swelling21,22. In addition, ATP-MgCl2 also had effects on the amelioration of microcirculation, restoration of membrane voltage, restoration of normal membrane permeability and improvement of cellular functions30,31.

Together, ATP showed protective effects on cold stored rat livers, and it might be a synthetical effect of multiple actions.

Mechanism of protective effect of ATP on hypothermically preserved rat livers

ATP had a very strong effect on vascular expansion32, but our current study demonstrated that exogenous ATP protected cold stored rat livers not through vascular expansion.

If ATP-MgCl2 protected the cold stored rat livers through vascular expansion, then addition of ATP alone to the perfusate should also exhibit a protective effect. But no protective effect was observed by the addition of ATP alone in our study (Data not shown). Moreover, addition of MgCl2 alone to the perfusate also showed no protective effect33,34. Addition of ADP-MgCl2 complex, which has a more effective action of vascular expansion, showed no protective effect as ATP-MgCl2 (Data not shown). An even more important finding was that, ATP-MgCl2 could enter cold stored rat liver cells in our study. This also directly confirmed that exogenous ATP-MgCl2 could protect cold stored rat livers through the intracellular mechanism. By our knowledge, no report has revealed that exogenous ATP could enter hepatocytes through the membrane, and the mechanism is still unclear. We suspect that the possible pathway might be considered as followings. First, as ATP is a large biomolecule, the membrane is impermeable to it under normal status. But the permeability is increased to ATP due to the activation of some membrane carrier proteins by hypothermia and anoxia. Second, ATP enters hepatocytes through the disrupted hepatocyte membrane. In addition, how does ATP play the protective effect after entering the cells is still poorly understood.

Taken together, these results indicate that exogenous ATP-MgCl2 could protect cold stored rat livers through an intracellular rather than an extracellular mechanism.

Participation of Mg2+ in protection of cold stored rat livers by exogenous ATP

As we know, ATP could form chelate with other extracellular bivalent cations (Ca2+, Sr2+, Mg2+, etc.). However, addition of ATP-MgCl2 complex could inhibit the dephosphorylation and deamino action of ATP, suppress the extracellular hydrolysis of ATP, and prevent the different dynamic effects by interaction of ATP and other extracellular cations35. The other possible reason may be that participation of Mg2+ may be required while

Table 2  Energy status in hypothermically preserved rat livers (n=6, mean±SD)

| Time  | Group A | Group B | Group C |
|-------|---------|---------|---------|
| 0 h   | 2.760±0.302 | 2.760±0.302 | 2.760±0.302 |
| 1 h   | 2.337±0.202 | 2.263±0.282 | 2.514±0.298 |
| 2 h   | 1.914±0.209 | 1.971±0.205 | 2.391±0.276a |
| 6 h   | 1.509±0.211 | 1.506±0.180 | 2.523±0.269a |
| 12 h  | 1.145±0.177 | 1.136±0.150 | 2.715±0.298a |
| 24 h  | 0.755±0.082 | 0.842±0.088 | 2.547±0.279a |
| 36 h  | 0.603±0.065 | 0.706±0.080 | 1.782±0.200a |

EC

| Time  | Group A | Group B | Group C |
|-------|---------|---------|---------|
| 0 h   | 0.871±0.093 | 0.871±0.093 | 0.871±0.093 |
| 1 h   | 0.803±0.090 | 0.789±0.083 | 0.791±0.083 |
| 2 h   | 0.741±0.082 | 0.736±0.082 | 0.743±0.079 |
| 6 h   | 0.653±0.071 | 0.645±0.068 | 0.695±0.071a |
| 12 h  | 0.543±0.063 | 0.537±0.060 | 0.660±0.070a |
| 24 h  | 0.380±0.045 | 0.406±0.045 | 0.601±0.068a |
| 36 h  | 0.316±0.040 | 0.348±0.042 | 0.471±0.051b |

Table 3  Activities of AST and LDH in perfusate (IU/L kg liver) (n=6, mean±SD)

| Time  | LDH       | AST       | LDH       | AST       | LDH       | AST       |
|-------|-----------|-----------|-----------|-----------|-----------|-----------|
| 6 h   | 80.2±3.8  | 4.6±0.6   | 112.7±4.1 | 8.9±0.9   | 188.4±5.1 | 15.6±1.4  |
| 12 h  | 76.4±2.2  | 4.8±0.5   | 123.8±3.6 | 8.2±0.8   | 170.1±6.2 | 14.1±1.7  |
| 24 h  | 9.3±1.8   | 1.2±0.4a  | 26.7±2.3  | 2.4±0.5a  | 42.6±3.5  | 4.4±0.6a  |
| 36 h  | 80.2±3.8  | 4.6±0.6   | 112.7±4.1 | 8.9±0.9   | 188.4±5.1 | 15.6±1.4  |

P <0.05, P <0.01 vs group A.

Participation of Mg2+ in protection of cold stored rat livers by exogenous ATP

As we know, ATP could form chelate with other extracellular bivalent cations (Ca2+, Sr2+, Mg2+, etc.). However, addition of ATP-MgCl2 complex could inhibit the dephosphorylation and deamino action of ATP, suppress the extracellular hydrolysis of ATP, and prevent the different dynamic effects by interaction of ATP and other extracellular cations35. The other possible reason may be that participation of Mg2+ may be required while
ATP goes through the cell membrane. The carrier protein has been found on the intima of mitochondria. The functional mechanism was found to be: ATP-Mg\(^{2+}\)+HPO\(_4^{2-}\)\(\rightarrow\)ATP-Mg\(^{2+}\)+HPO\(_4^{2-}\) [33]. Further investigation is needed to confirm whether there is such a carrier protein on the outside membrane of hepatocytes or not, and whether ATP enters hepatocytes by interaction with Mg\(^{2+}\) or not. In addition, there is also the possibility that, as a co-factor of many intracellular functions, Mg\(^{2+}\) could participate in a diverse of ATP dependent intracellular actions, such as Na\(^{+}-K\)^+ ATPase, Ca\(^{2+}\)-ATPase, and glycolysis [34].

In summary, the results of the current study suggest that exogenous ATP could protect cold stored rat livers by entering hepatocytes. ATP-MgCl\(_2\) should be a pivotal component in the hypothermic preservation solution. Further study is required to clarify the protective mechanism of ATP on cold stored organs, which may contribute to the development of hypothermic preservation solution.

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