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Kinetics of adenylate metabolism in human and rat myocardium

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Abstract

Pathways producing and converting adenosine have hardly been investigated in human heart, contrasting work in other species. We compared the kinetics of enzymes associated with purine degradation and salvage in human and rat heart cytoplasm assaying for adenosine deaminase, nucleoside phosphorylase, xanthine oxidoreductase, AMP deaminase, AMP- and IMP-specific 5'-nucleotidases, adenosine kinase and hypoxanthine guanine phosphoribosyltransferase (HGPRT). Xanthine oxidoreductase was not detectable in human heart. The \( K_m \)-values of the AMP-catabolizing enzymes were 2-5 times higher in human heart; the substrate affinity of the other enzymes was in the same order of magnitude in both species. The maximal activity (\( V_{max} \)) of adenosine kinase was the same in both species, but HGPRT in man was only 12% of that in the rat. For human heart the \( V_{max} \)-values of adenosine deaminase, nucleoside phosphorylase, AMP- and IMP-specific 5'-nucleotidases, and AMP deaminase were 25-50% of those for rat heart. We conclude that human heart is less geared to purine catabolism than rat heart as is evident from the lower activities of the catabolic enzymes. Maintenance of the nucleotide pool may thus play a more important role in human heart.

Keywords: Adenosine; Adenylate metabolism; Enzymology; Heart; Human; Rat

1. Introduction

Adenosine plays an important role in several cardiovascular processes. It is a potent vasodilator with depressant effects on contractility, heart rate and conductance of the pacemaker nodes [1]. The nucleoside may protect the heart by 'preconditioning' through A\(_1\)-receptor activation [2]. The pathways that produce and catabolize adenosine in hearts of several nonprimates have extensively been studied. In contrast, the activity and regulation of enzymes involved in purine-nucleotide metabolism, like AK, ADA, NP, AMP-D and HGPRT, have been investigated to a lesser extent in human tissues (see footnote for abbreviations). Also the presence of two isozymes of 5'-NT differing in substrate preferences (AMP- and IMP-specific) found in various animal hearts [3] has not been confirmed in human heart. Fig. 1 shows the pathways involved. Deactivation of adenosine is possible through rapid breakdown by ADA. This happens both inside and outside the cell [4]; the cellular process has been studied more thoroughly and is likely to be more important. ADA activity varies considerably in hearts of different species [5]. Metabolic inhibitors for purine degradation, such as allopurinol and EHNA, are in use in cardioplegic solutions [6-8], as are adenosine catabolites like inosine to treat heart failure [9,10]. Those clinical trials assume the presence and activity of certain enzymes in the human heart. However, the enzymological background for such studies is absent. The purpose of this study was to investigate the kinetics of cytosolic purine-metabolizing enzymes in human heart, with emphasis on the regulation of the adenosine level. This could give therapy and diagnostics with adenosine a
4°C. To get rid of small molecules, PD-10 Sephadex was washed with buffer. Extracts were immediately used in the assays or stored at —80° C.

For transplantation for clinical reasons. Because the kinetic data obtained were comparable to the ones from explanted diseased hearts, the data were pooled. Hearts were also excised and cooled on ice; they arrived in the laboratory within 10 min. Nonfibrotic samples from the left-ventricular wall were cut into pieces. Some control experiments (n = 4) were done on healthy donor hearts, unused and 4°C, whereas the 20% (w/v) homogenates were made in 20 mM Na+ -dimethylglutarate (pH 7.4).

2.2. Inhibitors

The AK inhibitor NSC 113939 was a gift of the National Cancer Institute (Bethesda, MD, USA). 5-Iodotuberculocidin (4-amino-5-ido-7-(β-d-ribofuranosyl)pyrrolo[2,3-d]pyrimidine), another AK inhibitor, was obtained from Research Biochemicals (Natick, MA, USA). The ADA inhibitor pentostatin (2'-deoxycoformycin, i.e., R-3-(2-deoxy-β-D-erythro-pentofuranosyl)-3,6,7,8-tetrahydro[4,5-d][1,3]diazapin-8-ol) was from Parke-Davis (Detroit, MI, USA). EHNA, an alternative ADA inhibitor, was a gift from Wellcome Research Laboratories (Beckenham, Kent, UK). AOPCP, a potent membrane 5'-NT inhibitor, was bought from Sigma (St. Louis, MO, USA).

2.3. Determination of enzyme kinetics

Assays were carried out at 30° C. The half-saturation constant (Km) and the maximal activity (Vmax) were determined, using Lineweaver-Burk linearization of the Michaelis-Menten equation [12]. One unit is 1 µmol substrate converted in one min or 1 µmol product formed in one min at 30° C. Values are presented as the mean ± S.E.

The activity of AK was measured with the radiometric method described by De Jong et al. [13]. There was no difference in results obtained with fresh and frozen (for ≤2 days) supernatant fluids. The incubation mixture contained 40 mM potassium phosphate (pH 7.0), 1.0 mM GTP-lithium salt, 1.0 mM MgCl2, bovine serum albumin 0.05 mg/ml, and [U-14C]adenosine (0.2–10 µM). After 10 min incubations were stopped by distribution of aliquots from the reaction mixture on squares of DEAE-cellulose (Whatman DE81) to bind the reaction product, [14-C]AMP. The squares were washed with 1 mM ammonium formate, water and ethanol.

ADA activity was determined spectrophotometrically (Hitachi U-2000 double-beam) at 265 nm according to Coddington [14]. During 5 min incubations the conversion of adenosine to inosine was measured in 50 mM potassium phosphate buffer (pH 7.4) with substrate concentrations between 2 and 100 µM.

The NP activity was analyzed spectrophotometrically (Hitachi U-2000 double-beam) at 293 nm during 5 min based on the method of Coddington [15] with modifications. Present were 0.1 M Tris (pH 7.4), xanthine oxidase 0.1 U/ml, and either various concentrations of KH2PO4 (0.1–50 mM) with 0.3 mM inosine, or various concentrations of inosine (0.02–1.0 mM) with 0.1 M K+ -phosphate buffer (pH 7.4).

The incubation mixture for the HGPRT determination contained 0.4 mM hypoxanthine, 0.5 M Tris/0.05 M MgCl2 (pH7.4), 8 mM PRPP and 10% Triton X-100 in the ratio 10:10:10:1. The reaction was started by adding 10 µl supernatant fluid to 50 µl incubation mixture. After 120 min at 30° C the reaction was stopped with HClO4 (0.4 M final concentration). Samples were kept on ice for 10
min and centrifuged for 4 min, 15,000 × g at 4°C. K₂HPO₄ was added to 50 µl supernatant fluid in a final concentration of 0.14 M. The samples were analyzed on HPLC as described previously [16].

For AMP-D determination Moss' method [17] modified by Meghji et al. [5] was used. The reaction mixture contained 20 mM sodium dimethylglutarate (pH 7.0), 1 mM EDTA, 0.1 mM dithiothreitol, 0.15 M KCl, 1 µM EHNA, 1 mM ATP, and AMP in the range 0.5–15 mM. The reaction was stopped with 50% HClO₄ after 20 min. After centrifugation for 5 min at 4°C, 15,000 × g, supernatant fluids were taken and neutralized with 1.2 M K₂CO₃/30 mM EDTA. After centrifugation the supernatant fluid was taken to estimate ammonia spectrophotometrically at 625 nm according to Chaney and Marbach [18].

5'-NT was determined at 37°C as described by Skladanowski et al. [19]. The assay is based on the method of Newby [20] with some modifications for 5'-nucleotidases of different specificity [21]. For both assays the conditions adopted secured lack of cross reactivity of the two forms. Briefly, the final concentrations in the incubation mixture for the AMP-specific form of 5'-NT were: 100 mM imidazole/HCl, pH 6.5, 50 mM MgCl₂, 30 mM NaCl, 1 mM ADP and AMP between 0.5–20 mM. To prevent conversion of adenosine to AMP and/or inosine, 2 µM 5-iodotubercidin and 10 µM EHNA were included. The reaction was stopped with 50 µl 1.3 M HClO₄ after 1.5 to 3 min. Samples were kept on ice for 30 min, deproteinized, centrifuged and analyzed with HPLC [22]. The incubation conditions for the IMP-specific form of 5'-NT were: 100 mM TES (or Hapes)/NaOH, pH 7.0, 10 mM MgCl₂, 30 mM NaCl, 1 mM ADP and AMP between 0.5–20 mM. The time course was linear for both assays. We recalculated the results from the earlier study [19] for 30°C, using factors of 0.75 for human and 0.71 for rat AMP- and IMP-preferred 5'-NT, which we determined experimentally.

### 3. Results and discussion

#### 3.1. Adenosine kinase

Use of the AK inhibitors NSC 113939 (50 µM) and 5-iodotubercidin (5 µM), proved that our AK measurements were specific; they inhibited the reaction almost completely (94% and 96%, respectively). Inclusion of the ADA inhibitor pentostatin (1 µM) in the reaction mixture did not change the Vₘₐₓ, which shows that ADA did not interfere with the assay. Addition of AOPCP (50 µM), a potent membrane 5'-NT inhibitor [23], had no influence on the AK activity. We avoided thus possible product conversion by ecto-5'-NT in the assay. The Vₘₐₓ of AK in human myocardial homogenates was the same as in rat heart (Table 1). In both species, the Kₘₐₓ-values were in the low micromolar range (Table 1). The observed species-differences are in line with those suggested earlier [25].

#### 3.2. Adenosine deaminase

The inhibitor pentostatin (3 µM) completely abolished ADA activity, showing that other reactions did not interfere with our assay. In human heart the Vₘₐₓ was about 30% of that in rat heart (Table 1); however, the Kₘₐₓ-values were almost identical (Table 1). Our results on ADA activity in human and rat heart agree with those published by Meghji et al. [5], although they made only one-point estimations. The kinetic data on rat heart are in line with Arch and Newsholme [26].

### Table 1

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Substrate</th>
<th>Vₘₐₓ (mU/g wet weight)</th>
<th>Kₘ (µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Rat</td>
<td>Human</td>
</tr>
<tr>
<td>AK</td>
<td>adenosine</td>
<td>38.9 ± 3.2 (3)</td>
<td>37.7 ± 0.8 (9)</td>
</tr>
<tr>
<td>ADA</td>
<td>adenosine</td>
<td>715 ± 36 (3)</td>
<td>212 ± 17 (11)</td>
</tr>
<tr>
<td>NP</td>
<td>inosine</td>
<td>738 ± 52 (3)</td>
<td>245 ± 25 (8)</td>
</tr>
<tr>
<td>XOD</td>
<td>xanthine</td>
<td>38.6 ± 1.8 (5) a</td>
<td>&lt; 1</td>
</tr>
<tr>
<td>HGPRT</td>
<td>hypoxanthine</td>
<td>38.1 ± 4.3 (5) a</td>
<td>&lt; 1</td>
</tr>
<tr>
<td>AMP-D</td>
<td>AMP</td>
<td>1837 ± 152 (3)</td>
<td>976 ± 20 (5)</td>
</tr>
<tr>
<td>5'-NT</td>
<td>AMP</td>
<td>362 ± 17 (3)</td>
<td>128 ± 37 (6)</td>
</tr>
<tr>
<td>IMP</td>
<td>IMP</td>
<td>533 ± 11 (5)</td>
<td>35 ± 9 (6)</td>
</tr>
</tbody>
</table>

Enzyme assays were done on myocardial supernatant fluids. The activity of the catabolizing enzymes proved to be in human heart 12-50% of that in rat heart; the activity of adenosine kinase was the same in both species. In human heart the Kₘₐₓ-values were in the same order of magnitude as in rat heart, although they were at least twice higher for mononucleotide-catabolizing enzymes. Mean ± S.E. with number of experiments in parentheses. For abbreviations, see legend to Fig. 1. n.d., not detectable. N.D., not determined.

a Data from [24].
3.3. Nucleoside phosphorylase

In human myocardial tissue, the $V_{\text{max}}$ of NP with inosine or KH$_2$PO$_4$ as variable substrate was about 30% of that in rat (Table 1). The $V_{\text{max}}$ estimated with varying inosine concentrations was about twice higher than with KH$_2$PO$_4$ as variable substrate. This could be attributed to inhibition by the Tris-buffer used when phosphate was varied. The $K_m$, however, was similar in both species (Table 1). We are unaware of kinetic data on human NP activity except for a qualitative estimation of activity in human and rat heart [27].

3.4. Xanthine oxidoreductase

Earlier our group could not detect XOD activity in human-heart homogenate. We found a very low activity in perfused diseased human hearts [28], agreeing with the results of others [see, e.g., [29,30]]. In rat heart we determined activity [31] and kinetic data for XOD with hypoxanthine and xanthine as substrate [24]. For both substrates the $V_{\text{max}}$-values were the same; the $K_m$-values were in the lower micromolar range (Table 1).

3.5. Hypoxanthine guanine phosphoribosyltransferase

In human heart the $V_{\text{max}}$ of HGPRT with hypoxanthine as substrate was only 12% of the value for rat heart. Purine degradation, but also purine salvage to IMP via HGPRT, appears slow in human heart compared with rat heart.

3.6. AMP deaminase

Side reactions only interfered in rat heart at substrate concentrations higher than 15 mM by an artificial sharp increase of ammonia production. The dependence of ammonia production on AMP concentration was biphasic in rat- but not in human-heart homogenate. HPLC analysis of the postreactional mixtures revealed that the accelerated ammonia production at $\geq$ 15 mM AMP correlated with formation of adenosine and inosine. Then the EHNA concentration (1 $\mu$M) seemed too low to inhibit ADA completely. At low concentrations of AMP, the production of adenosine, if detectable, was not accompanied by increased inosine, showing effective inhibition of ADA. We decided to estimate activity values at substrate concentrations $\leq$ 10 mM. Then we observed a $V_{\text{max}}$ for rat heart AMP-D twice higher than for human heart, where the delayed increase of ammonia did not take place (Table 1). The $K_m$-values for AMP-D were similar in human and rat heart, i.e., in the lower millimolar range (Table 1). We found lower activities of AMP-D in both species than Meghji et al. [5]. This is presumably due to their overestimation caused by EHNA's incomplete inhibition of ADA at 25 mM AMP. The presence of 1 mM ATP in the AMP-D assay ensured that the sigmoidal curve turned into a hyperbolic one. Our findings partly confirm other literature data on rat [32,33] and human heart [34].

3.7. 5'-Nucleotidase

In human heart the $V_{\text{max}}$ of AMP- and IMP-preferred 5'-NT was 25–30% of that in rat heart (Table 1). The $K_m$-values for both substrates were 2–4 times higher in human heart (Table 1), suggesting a lower affinity of 5'-NT for AMP and IMP in man. Our results in rat heart were comparable with those from Truong et al. [35]. We found a 13-fold lower initial concentration of inosine accumulated in freshly prepared human-heart supernatant fluids compared to rat-heart ones. Concentrations were $8.6 \pm 1.2 \ \mu$M ($n = 7$) and $116 \pm 3 \ \mu$M ($n = 7$), respectively. Additive effects of lower AMP dephosphorylation by 5'-NT and adenosine deamination by ADA in the human heart caused this difference with rat heart.

3.8. Physiological implications

One has to be cautious in the interpretation of the $V_{\text{max}}$ of a particular reaction. We realize that we calculated kinetic values on enzymes present in cytosol, not on purified proteins. On the other hand, we avoided the instability problems sometimes seen with purified enzymes. In human myocardium the maximum velocities of the enzymes catabolizing adenylate to hypoxanthine (AMP-D, AMP- and IMP-specific 5'-NT, ADA and NP) were in a highly regular order 25–50% of those in rat heart. The $V_{\text{max}}$ of the anabolic enzyme AK was the same in both species, whereas the maximal HGPRT activity for human heart was only 12% of that observed for rat. We found the $K_m$-values of AK, ADA and NP to be either the same or in the same order of magnitude for both species. Thus we expect that the affinity of these enzymes to their substrates is similar. The $K_m$-values of AMP-D and 5'-NT (AMP/IMP) were higher in human than in rat heart, suggesting a lower affinity of these two enzymes in human heart. Slower purine breakdown but not purine salvage via AK possibly protects the human heart at an earlier stage through purine accumulation instead of rapid breakdown to urate. Maintenance of the cardiac nucleotide pool may play a more important role in man than in rat.

A possible difference in tissue adenosine levels between human and rat heart may be assessed by relating the flux rate determined from AMP to adenosine with the rate of adenosine degradation to inosine. From the data (Table 1) it appears that the lower $V_{\text{max}}$-value for 5'-nucleotidase in human heart is compensated by slower adenosine catabolism.

Borst et al. [36] determined kinetic values for SAH hydrolase in patients with hypertrophic obstructive cardiomyopathy. Its activity is very low compared to the other enzyme activities involved in purine metabolism in human heart. We reported earlier that SAH hydrolase activity does...
not contribute significantly to adenosine during ischemia, but could play a role during normoxia [37].

Our results give the maximal velocity obtainable and affinity toward substrates for the enzymes investigated. Free cytosolic AMP-concentrations in cardiomyocytes are probably in the submicromolar range even under ischemic conditions [38]. This is far below the $K_m$ of 5'-NT and AMP-D. We suggest that the rise in AMP-concentration due to hypoxia will give a large response in adenosine formation, reinforced by allosteric activation of 5'-NT. The next step is the rapid conversion to inosine by ADA (see Table 1).

The breakdown of inosine to hypoxanthine by NP depends on the concentration of phosphate. Its physiological concentration in rat heart exceeds the $K_m$ (phosphate) for NP by a factor of 1.5 [39]. When the free phosphate concentration rises (about three times) due to ischemia, the activity of NP might increase, approaching the $V_{max}$-value. Thus, the intracellular phosphate concentration is possibly rate-limiting for adenosine degradation to hypoxanthine. We confirmed this in our AMP-specific 5'-NT assay: the amount of inosine present in the supernatant fluid – stoichiometric with phosphate liberated from AMP – was converted to hypoxanthine by NP both in rat and human heart. Hypoxanthine salvage to IMP by HGPRT in human heart is possible but slow.

3.9. Cardiac IMP production

In skeletal muscle and liver, IMP produces inosine. There is consensus that in ischemic heart muscle like in brain, AMP degradation via adenosine is preferred. Olsson and Pearson suggested that the IMP pathway is dominant in cells generating ATP predominantly through anaerobic glycolysis; the adenosine pathway seems prominent in organs relying on oxidative phosphorylation [40]. Our in vitro data would show that the capacity of the IMP pathway exceeds the adenine pathway in both species (see Table 1).

Association with membrane phospholipids activates AMP-D [41]. Recently, Tanfani et al. suggested that synchronized AMP-D inhibition by ATP decrease and this association loss in ischemic heart shifts adenylate to adenosine production by 5'-NT [41]. The cardioplegia-arrested human heart produces considerable amounts of adenosine besides inosine [42,43]. However, IMP accumulates in human heart as well [43].

3.10. Limitations of the study

We measured the kinetics of the enzymes in homogenates, which consist of endothelial cells (ca. 4%) and cardiomyocytes. The localization of the enzymes in myocytes and/or endothelial cells differs in species. For example, nucleoside phosphorylase in guinea-pig heart is completely localized in the endothelial cells [44]; in rat heart we found this enzyme to be also present in the myocytes [11]. It is not known if the distribution of the enzymes discussed in this paper is the same in rat and human heart. We also did not deal with kinetics of nucleoside transport in cardiomyocytes and endothelial cells in this study. We only measured the kinetics of the enzymes involved in purine metabolism under optimal conditions in human and rat heart.

3.11. Conclusion

Human heart with its slower rate of adenosine formation from AMP, its lower activity of purine-catabalizing enzymes and its virtual absence of XOD, is less geared to adenosine production and subsequent degradation than rat heart.

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