Modulation of cardiac ionic homeostasis by 3-iodothyronamine

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Abstract

3-iodothyronamine (T 1AM) is a novel endogenous relative of thyroid hormone, able to interact with trace amine-associated receptors, a class of plasma membrane G protein-coupled receptors, and to produce a negative inotropic and chronotropic effect. In the isolated rat heart 20–25 μM T 1AM decreased cardiac contractility, but oxygen consumption and glucose uptake were either unchanged or disproportionately high when compared to mechanical work. In adult rat cardiomyocytes acute exposure to 20 μM T 1AM decreased the amplitude and duration of the calcium transient. In patch clamped cardiomyocytes sarcolemmal calcium current density was unchanged while current facilitation by membrane depolarization was abolished consistent with reduced sarcoplasmic reticulum (SR) calcium release. In addition, T 1AM decreased transient outward current (Ito) and IK1 background current. SR studies involving 20 μM T 1AM revealed a significant decrease in ryanodine binding due to reduced B max, no significant change in the rate constant of calcium-induced calcium release, a significant increase in calcium leak measured under conditions promoting channel closure, and no effect on oxalate-supported calcium uptake. Based on these observations we conclude T 1AM affects calcium and potassium homeostasis and suggest its negative inotropic action is due to a diminished pool of SR calcium as a result of increased diastolic leak through the ryanodine receptor, while increased action potential duration is accounted for by inhibition of h0 and k1 currents.

Keywords: 3-iodothyronamine ♦ ionic current ♦ calcium ♦ potassium ♦ sarcoplasmic reticulum ♦ ryanodine receptor ♦ heart ♦ thyroid

Introduction

3-iodothyronamine (T 1AM) is a novel endogenously synthesized relative of thyroid hormone with significant physiological and behavioural effects in mammals [1]. In isolated rat hearts micromolar concentrations of T 1AM produced a reversible, dose-dependent negative inotropic and chronotropic effect [1, 2]. In cultured rat cardiomyocytes decreased cellular shortening and increased action potential duration were reported upon exposure to T 1AM [2]. These actions occur within minutes and consequently they appear to represent a non-genomic response. Interestingly, T 1AM does not interact with nuclear thyroid hormone receptors. Rather in vitro it can activate heterologously expressed cloned rodent and human trace amine associated receptor 1 (TAAR1), and possibly other members of the same family of G-protein coupled receptors [3, 4]. Messenger RNAs coding for at least five different TAAR subtypes are expressed in rat heart and it has been suggested that one or more TAAR subtypes mediate T 1AM’s diverse biological effects [2]. There is convincing evidence that transduction of T 1AM-stimulated signalling involves changes in the phosphorylation state of tyrosine residues in yet-to-be-determined proteins, but the final effectors responsible for the functional effects have not been determined. Because myocardial function is largely dependent on the modulation of ionic currents and requires an adequate supply of metabolic energy, the present study attempted to understand T 1AM’s effects on myocardial metabolism and ionic homeostasis.

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Materials and methods

Chemicals and radionuclides

T1AM and thyronamine were synthesized as described elsewhere [5]. [3H]-ryanodine and [45Ca]-CaCl2 were obtained from New England Nuclear (Milan, Italy). Unless otherwise specified all other reagents were from Sigma-Aldrich (St. Louis, MO, USA).

Isolated heart perfusion

This investigation conforms to the Declaration of Helsinki and the Guiding Principles in the Care and Use of Animals. The project was approved by the Animal Care and Use committee of the University of Pisa. Male Wistar rats (275–300 g body weight), fed a standard diet, were anesthetized with a mixture of ether and air. The heart was then quickly excised and perfused according to the working heart technique, as described previously [2]. The height of the atrial chamber was set at 20 cm, corresponding to a filling pressure of 15 mmHg. To measure oxygen consumption the pulmonary artery was cannulated and samples of perfusate were collected from the aortic and pulmonary cannulas without exposure to air, as described previously [6]. Glucose was assayed in the recirculating perfusion buffer by the glucose oxidase technique using a commercial Glucose Assay Kit (GAHK-20, Sigma-Aldrich), and the rate of glucose uptake was calculated by linear regression.

Cardiomyocyte experiments

Left ventricular cardiomyocytes were prepared from male Wistar rat hearts by enzymatic digestion in a Langendorff apparatus. The isolated cells were resuspended in Tyrode’s solution, as described previously [7]. The following solutions were used (in mM): Tyrode’s solution, i.e. NaCl 140, KCl 5.4, MgCl2 1.2, glucose 10, HEPES-NaOH 5 (pH 7.3), CaCl2 1.8; modified Tyrode’s solution used for L-type calcium current recording, i.e. KMeSO4 125, KCl 25, EGTA 1, amphotericin B 0.13, HEPES-KOH 5 (pH 7.0). The experimental set-up for patch-clamp recording and data acquisition was similar to that described previously [8]. Dissociated cardiomyocytes grown in culture were placed in an experimental bath set on the stage of an inverted microscope (Nikon Diaphot TMD, Kawasaki, Japan). The patch-clamped cell was superfused by means of a temperature-controlled (36 ± 0.5°C) micro-superfuser that allows rapid changes in the cell’s surrounding solution to be made. Patch-clamp pipettes, prepared from glass capillary tubes by means of a two-stage horizontal puller (P-87 Flaming/Brown micropipette puller, Sutter Instrument, Novato, CA, USA), had a resistance of ~2.5 MΩ when filled with pipette solution.

Recordings were performed with a patch amplifier (Axopatch 200B, Molecular Devices, Sunnyvale, CA, USA) in whole-cell configuration for current recording or by means of the perforated-patch technique for calcium transient recordings. Signals were digitized via a DAC/ADC interface (Digidata 1200B, Molecular Devices) and acquired by means of pClamp software. Currents were recorded in the voltage-clamp mode. Cell membrane capacitance was measured by applying a ±10 mV pulse starting from a holding potential of –70 mV, as previously reported [8]. Offline data analysis was performed with pClamp (vers. 9, Molecular Devices) and Origin 7.5 (GraphPad Software, San Diego, CA, USA).

The amplitudes of L-type calcium currents (I Ca-L) were measured by applying a voltage protocol for steady-state activation (from –35 to +55 mV after a pre-step of –40 mV, to inactivate sodium currents) and inactivation (pre-steps in the range –75 to –5 mV followed by a test step at 0 mV to record maximal current amplitude). Bi- or mono-exponential fitting was used to determine the time constants of deactivation. Background currents were elicited by a ramp protocol from –120 to –50 mV and currents were measured at +40 mV and at –100 mV, chosen because they are representative of the ultra-rapid delayed rectifier current (IKur) and the inward rectifier current (IK1), respectively. Transient outward current (IO) was measured by applying depolarizing steps from –40 to +60 mV and maximal current density was plotted versus step potential to obtain an activation curve.

To evaluate the effect of T1AM on calcium transient, left ventricular cardiomyocytes were incubated with Tyrode’s solution containing 1–5 μM fluo-3-acetoxymethyl ester (Fluo-3) and Pluronic F127 (Molecular Probes, Eugene, OR, USA) for 20 min. at 37°C to load Fluo-3. The preparation was set in the patch-clamp setup previously described, properly modified for fluorescence signals production and detection. Fluo-3 was excited with the 488 nm line of a xenon lamp (LPS 220, Photo Technology International, Birmingham, NJ, USA). Fluorescence was detected at 540 ± 15 nm with a photomultiplier tube (model 01–614, Photo Technology International) and signal was digitized by a DAC/ADC interfaces (Digidata 1200B, Molecular Devices). Oscilloscopes and computer screens allowed online data view. Experimental control, data acquisition and preliminary analysis were performed by means of the integrated software package pClamp. Calcium transient was recorded during a pulse at 0 mV from a holding potential at –40 mV.

Sarcoplasmic reticulum (SR) experiments

Ventricles were obtained from perfused hearts after 30 min. of exposure to T1AM or an equivalent period of control perfusion. The tissue was finely minced and homogenized in 5 volumes of 300 mM sucrose and 10 mM imidazole (pH 7.0) at 4°C by 15 ± 15 passes in a Potter-Elvejheim homogenizer set at 800 rpm. A microsomal fraction enriched in SR was prepared by high-speed differential centrifugation as described elsewhere [9]. The techniques to assay SR function, previously described and validated [9–11], are summarized below. Measurements were typically performed on crude homogenates because SR purification has a low yield and may not be representative of the intact SR [12].

Active Ca2+ transport was evaluated on the basis of oxalate-supported Ca2+ uptake. Aliquots of homogenate (protein concentration 0.5–0.8 mg/ml) were pre-incubated at 37°C for 5 min. in the presence of 900 μM ryanodine to block the SR Ca2+ release channel. The uptake medium and the experimental technique were the same as described previously [9]. In a few experiments, different amounts of CaCl2 were added to obtain free Ca2+ concentrations ranging from 0.1 to 40 μM. SR Ca2+-induced Ca2+ release was determined after passive loading with 10 mM [45Ca]-CaCl2 [10]. Ca2+ release was induced by washing the loaded vesicles with release buffer containing (in mM): HEPES potassium 20 (pH 6.8), KCl 100, CaCl2 1.01 and EGTA 1 (free calcium concentration was 18 μM) 1. A rapid filtration system with time resolution on the order of 10 msec. was used (RFS-94, Bio-Logics, Grenoble, France) and the rate constant of quick Ca2+ release was calculated over the first 100 msec. by exponential fitting.
SR Ca$^{2+}$ leak was measured under conditions promoting SR channel closure. For this purpose SR vesicles were actively loaded by 2 min. incubation in the presence of 100 mM $[^{45}Ca]$-CaCl$_2$ and 2 mM acetylphosphate [11, 13]. The preparation was filtered using the RFS-04 equipment charged with the following buffer (in mM): Tris-MOPS (pH 7.0) 20, KCl 80, MgCl$_2$ 10, Tris-EGTA 1 and tetracaine 0.3. EGTA is used as a calcium chelator, while magnesium and tetracaine are used to produce channel blockade. Release rates were calculated by exponential fitting over 2 sec.

To determine high affinity ryanodine binding aliquots of homogenate were incubated at 37°C for 60 min. with 0.2 to 50 nM $[^{3}H]$-ryanodine, in a buffer containing (mM): imidazole 25 (pH 7.4 at 37°C), KCl 1000, EGTA 0.950 and CaCl$_2$ 1.013 (free Ca$^{2+}$ concentration: 20 μM) [9]. Saturation experiments were analysed by nonlinear fitting of a single binding site model.

Statistical analysis

Results are expressed as the mean ± S.E.M. Differences between groups were evaluated by ANOVA and individual groups were compared versus the control group using the Dunnett post hoc test. When only two groups were compared, the paired or unpaired t-test was used, as deemed appropriate.

Data represent mean ± S.E.M. of 10–16 hearts per group. Measurements were obtained after 20 min. of perfusion. HF, heart rate; AF, aortic flow; CF, coronary flow; CO, cardiac output and AoP, systolic aortic pressure.

Electrophysiological effects

The electrophysiological effects of T1AM were studied in adult ventricular cardiomyocytes using the whole-cell patch-clamp technique. On the basis of our isolated heart results we decided to use 20 μM T1AM. T1AM had no effect on peak density of calcium currents but affected $I_{Ca-L}$ inactivation kinetics (Fig. 1). Under control conditions $I_{Ca-L}$ inactivated with a typical bi-exponential decay, characterized by two deactivation constants ($\tau_t$: 9.0 ± 2.6 msec.; $\tau_d$: 50.0 ± 10.0 msec.; n = 4). However, in the presence of T1AM $I_{Ca-L}$ inactivated with a mono-exponential decay profile and the deactivation constant averaged 26.0 ± 2.0 msec. (n = 4, P < 0.01 versus control).

In addition, T1AM influenced the so-called L-type current facilitation, a typical feature of cardiac cells [14, 15]. This is illustrated in Fig. 1A: under control conditions, the current evoked from a holding potential of ~45 mV was ~30% greater than that evoked from a holding potential of ~75 mV. T1AM completely abolished this phenomenon because the currents measured at ~45 and ~75 mV had the same amplitude (Fig. 1B). The steady state inactivation

Table 1  Haemodynamic and metabolic variables

| Group     | HR (bpm) | AF (ml/min.) | CF (ml/min.) | CO (ml/min.) | AoP (mmHg) | Glucose uptake (μmol/min. per g) | Oxygen consumption (μmol/min. per g) |
|-----------|----------|--------------|--------------|--------------|------------|---------------------------------|-------------------------------------|
| Control   | 271 ± 14 | 43.4 ± 0.8   | 19.6 ± 0.6   | 62.9 ± 1.0   | 177 ± 9    | 7.6 ± 1.1                       | 40.0 ± 3.9                          |
| T1AM (2.5 μM) | 231 ± 18 | 37.3 ± 2.9   | 19.3 ± 2.4   | 56.7 ± 0.7   | 173 ± 7    | 7.3 ± 0.8                       | 37.4 ± 3.5                          |
| T1AM (20 μM)  | 210 ± 12* | 22.7 ± 3.3†  | 17.7 ± 1.0   | 40.4 ± 3.3†  | 158 ± 8    | 7.2 ± 0.8                       | 37.1 ± 5.8                          |
| T1AM (25 μM)  | 154 ± 16† | 13.2 ± 2.2†  | 14.4 ± 0.8†  | 27.6 ± 2.4†  | 127 ± 18* | 4.5 ± 1.0                       | 21.2 ± 3.2*                         |

Data represent mean ± S.E.M. Differences between groups were analyzed by ANOVA while individual groups were compared versus the control group using the Dunnett post hoc test, as deemed appropriate.

*P < 0.05, †P < 0.01 versus control, by Dunnett’s test.

In the isolated perfused working rat heart model baseline average values of the haemodynamic variables were as follows: cardiac output 62.9 ± 1.0 ml/min., aortic flow 43.4 ± 0.8 ml/min., coronary flow 19.6 ± 0.6 ml/min., systolic aortic pressure 177 ± 9 mmHg, heart rate 271 ± 14 beats per minute. In accordance with our previous findings, 2.5 μM T1AM did not modify contractile performance while 20 μM and 25 μM caused a significant decrease in cardiac output, aortic pressure, coronary flow and heart rate. The time course of these changes was the same as reported previously [2], and the maximum response occurred after 10–20 min. The values of the haemodynamic variables recorded after 20 min. are summarized in Table 1. However, these haemodynamic effects were not associated with comparable reductions of oxygen consumption or glucose uptake. Actually with 20 μM T1AM glucose uptake and oxygen consumption were not significantly modified (7.2 ± 0.8 versus 7.6 ± 1.1 μmol/min. per g and 37.1 ± 5.8 versus 40.0 ± 3.9 μmol/min. per g, respectively) in spite of 36% and 22% decrease in cardiac output and heart rate, respectively. A significant reduction in oxygen consumption was observed only with 25 μM T1AM, which determined 56% and 43% reduction of cardiac output and heart rate, respectively.

Results

Functional and metabolic effects

In the isolated perfused working rat heart model baseline average values of the haemodynamic variables were as follows: cardiac output 62.9 ± 1.0 ml/min., aortic flow 43.4 ± 0.8 ml/min., coronary flow 19.6 ± 0.6 ml/min., systolic aortic pressure 177 ± 9 mmHg, heart rate 271 ± 14 beats per minute. In accordance with our previous findings, 2.5 μM T1AM did not modify contractile performance while 20 μM and 25 μM caused a significant decrease in cardiac output, aortic pressure, coronary flow and heart rate. The time course of these changes was the same as reported previously [2], and the maximum response occurred after 10–20 min. The values of the haemodynamic variables recorded after 20 min. are summarized in Table 1. However, these haemodynamic effects were not associated with comparable reductions of oxygen consumption or glucose uptake. Actually with 20 μM T1AM glucose uptake and oxygen consumption were not significantly modified (7.2 ± 0.8 versus 7.6 ± 1.1 μmol/min. per g and 37.1 ± 5.8 versus 40.0 ± 3.9 μmol/min. per g, respectively) in spite of 36% and 22% decrease in cardiac output and heart rate, respectively. A significant reduction in oxygen consumption was observed only with 25 μM T1AM, which determined 56% and 43% reduction of cardiac output and heart rate, respectively.
curve obtained by normalization of calcium current versus conditioning step potential (Fig. 1C) confirmed that T1AM abolished the overshoot observed in the control curve at holding potential between –50 and –30 mV. However, T1AM had no effect on the increase of the calcium current produced by exposure to 1 μM isoproterenol (data not shown), thus suggesting that the activity of protein kinase A is not hampered by exposure to T1AM.

Facilitation has been attributed to the effect of SR calcium release on the L-type calcium channel and it is abolished when the SR calcium pool is depleted [15, 16]. Therefore we investigated the influence of 20 μM T1AM on intracellular calcium transients. Cardiomyocytes were loaded with Fluo3 to obtain a fluorescence signal proportional to the intracellular concentration of free calcium. Figure 2 shows the results of a typical experiment in which the calcium transient was recorded during a pulse at 0 mV from a holding potential of –40 mV, and the substantial reduction of its amplitude that occurred in the presence of T1AM. In eight independent preparations T1AM produced significant decreases both in the peak of the signal (0.06 ± 0.01 versus 0.10 ± 0.02 arbitrary units, P < 0.05) and its decay time (86 ± 26 versus 176 ± 40 msec., P < 0.01).

We previously reported T1AM can produce a negative chronotropic effect which is associated with action potential prolongation [2]. This was confirmed in the present investigation. Cardiomyocytes exposed to 20 μM T1AM displayed an APD90 that increased from 84 ± 16 to 157 ± 29 msec. (n = 10, P < 0.05). To gain insight into the mechanism responsible for action potential prolongation we investigated the effects of 20 μM T1AM on Ito current, responsible for the first phase of the action potential repolarization, and on background currents (IK1 and IKur), responsible for the last phase of repolarization.

Background currents were elicited by a ramp protocol and currents were measured at –100 and –40 mV, representative of IK1 and IKur, respectively (Fig. 3). IK1 was significantly reduced after exposure to 20 μM T1AM (pA/pF averaged –12.8 ± 1.5 versus –17.6 ± 2.2; n = 7, P < 0.05), whereas the changes in IKur did
not reach statistical significance. The activation curve of \( h_0 \) current, obtained by plotting maximal \( h_0 \) density versus step potential (Fig. 4), revealed peak amplitude to be significantly reduced after exposure to T1AM at 20–50 mV (* = \( P < 0.05 \)). See text for analytical results.

**Effects on SR function**

To investigate whether the observed modifications of the calcium transient are associated with changes in SR function we determined the effects of perfusing with 20 \( \mu \)M T1AM on ryanodine binding, SR calcium release and oxalate-supported calcium uptake.

Ryanodine is a selective ligand of the SR calcium release channels and changes in ryanodine binding provide indirect evidence of changes in channel structure or gating [17]. Saturation binding curves obtained in control hearts and in hearts perfused with T1AM are shown in Fig. 5 (left panel). After perfusion with T1AM \(^{[3]H}\)-ryanodine binding was significantly reduced, due to decreased \( B_{\text{max}} \) (265 ± 19 \( \text{fmol/mg of protein} \), \( P < 0.05 \)), while the \( K_D \) was not significantly altered (1.7 ± 0.5 \( \text{versus} \ 2.1 \pm 0.2 \text{ nM} \)). Incubation of control SR vesicles with 20 \( \mu \)M T1AM did not affect ryanodine binding (Fig. 5, right panel), demonstrating the observed effect was not due to direct interaction of T1AM with the SR calcium channel.

In other experiments SR vesicles obtained from hearts which had been perfused with T1AM were loaded with radiolabelled calcium and the kinetics of calcium induced calcium release were determined by a quick filtration technique, using a buffer containing 18 \( \mu \)M unlabelled calcium to determine full channel activation. As shown in the left panel of Fig. 6 no significant effect could be demonstrated, although there was a trend for the average value of the first-order release constant to be higher in the T1AM group (89 ± 18 \( \text{versus} \ 60 \pm 15 \text{ sec.}^{-1}, P = 0.28 \)).

Calcium release experiments were also performed under conditions which induce SR calcium channel closure in an effort to measure the so-called SR calcium leak. The results are shown in the right panel of Fig. 6. Under our experimental conditions the rate constant of calcium release was significantly increased following perfusion with 20 \( \mu \)M T1AM (0.32 ± 0.03 \( \text{versus} \ 0.13 \pm 0.04 \text{ sec.}^{-1}, P < 0.01 \)).

To evaluate active calcium transport by the SR calcium ATPase we measured oxalate-supported calcium uptake in the crude homogenate obtained from control hearts, and from hearts perfused with T1AM. With this technique oxalate precipitates calcium in the SR and allows the uptake rate to be linear for several minutes, so that it can be easily determined even in unfractionated preparations [9, 12]. As shown in Fig. 7, at a free calcium concentration of 10 \( \mu \)M the uptake rate was not modified by perfusion with 20 \( \mu \)M T1AM. In vitro incubation with 20 \( \mu \)M T1AM was also ineffective (data not shown). In additional experiments calcium uptake was measured at five different calcium concentrations in an effort to estimate its calcium-dependence. In two different preparations the EC50 for calcium uptake averaged 2.6 \( \mu \)M both in control hearts and in hearts which had been perfused with 20 \( \mu \)M T1AM. The maximum uptake rate was not significantly modified, as well (11.8 ± 2.5 \( \text{versus} \ 12.0 ± 0.4 \text{ nmol/min per mg}, P = \text{NS} \)).
T1AM is a recently identified, naturally occurring compound that is most likely derived in vivo from thyroid hormone through decarboxylation and deiodination. In vitro evidence suggests T1AM activates a novel aminergic system coupled to the so-called TAARs [1, 2, 18]. Several functional effects of T1AM have already been described, and the focus of the present work was to unravel the molecular mechanism underlying its effects on contractility and action potential duration.

In the isolated rat heart a negative inotropic response to T1AM was observed before any decrease in oxygen consumption or glucose uptake is produced. Therefore, it seems unlikely that perfusion with T1AM impaired contractile performance because of energy shortage. On the other hand, cardiomyocyte experiments showed that T1AM substantially reduced the amplitude and duration of depolarization-induced calcium transients. A 20 μM concentration reduced the area under the calcium-time curve by more than 50%. On the basis of literature data and mathematical models of calcium homeostasis [19] this response is consistent with the observed reduction in cellular shortening, which averaged about 40% in our cardiomyocyte model [2]. Consequently, we hypothesize that the negative inotropic effect of T1AM is linked to reduced calcium mobilization.

The calcium transient is dependent on calcium influx through L-type sarcolemmal calcium channels and calcium release from the SR via the SR calcium channel–ryanodine receptor. Whole-cell patch-clamp experiments did not show any evidence of reduced L-type sarcolemmal calcium influx, because peak current density was unchanged. However, the phenomenon known as facilitation, namely increased current amplitude at more positive holding
that T1AM reduces SR calcium release. However, the so-called calcium leak, i.e. calcium release deter-
Determined under conditions able to lock the SR channel closed, was significantly increased after perfusion with T1AM.

Additional evidence that T1AM affects SR calcium release is provided by the observation that ryanodine binding was reduced after perfusion with T1AM. This does not appear to represent the consequence of a direct interaction between T1AM and the SR channel, because incubation of SR preparations with T1AM did not modify ryanodine binding. In general, a large number of physiological and pharmacological modulators of SR calcium release affect ryanodine binding, and altered ryanodine binding is usually, although not invariably, associated with altered calcium release [17]. When we determined the kinetics of SR calcium-induced calcium release by quick filtration experiments the first-order rate constant was not significantly modified after perfusion with T1AM. However, the so-called calcium leak, i.e. calcium release determined under conditions able to lock the SR channel closed, was significantly increased after perfusion with T1AM.

When interpreting these findings it should be emphasized that modulation of SR calcium activation is less important than modulation of diastolic leak through the SR, insofar as inotropic status is concerned. Transmembrane calcium flux is the product of membrane permeability and concentration gradient; therefore, SR calcium release depends both on channel gating and on the extent of the SR calcium pool. Increased calcium release due to increased channel opening in systole would decrease SR calcium concentra-

Potentials (~50 and ~30 mV), was abolished. Facilitation has generally been attributed to L-channel modulation by SR calcium release and is impaired after depletion of SR-sequestered calcium [15]. While facilitation may be also favoured by L-channel phosphorylation by protein kinase A [15], this mechanism can be ruled out in the present context, because isoproterenol-induced facilitation was not affected by T1AM. Therefore, our findings suggest that T1AM reduces SR calcium release.

Additional evidence that T1AM affects SR calcium release is provided by the observation that ryanodine binding was reduced after perfusion with T1AM. This does not appear to represent the consequence of a direct interaction between T1AM and the SR channel, because incubation of SR preparations with T1AM did not modify ryanodine binding. In general, a large number of physiological and pharmacological modulators of SR calcium release affect ryanodine binding, and altered ryanodine binding is usually, although not invariably, associated with altered calcium release [17]. When we determined the kinetics of SR calcium-induced calcium release by quick filtration experiments the first-order rate constant was not significantly modified after perfusion with T1AM. However, the so-called calcium leak, i.e. calcium release determined under conditions able to lock the SR channel closed, was significantly increased after perfusion with T1AM.

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**Fig. 7** Effect of perfusion with 20 μM T1AM on oxalate-supported calcium uptake in the crude rat heart homogenate at 10 μM free calcium concentration. Data represent mean ± S.E.M. obtained in four different preparations. The difference between groups was not statistically significant by unpaired t-test.
TAAR1 but rather may well occur via a different TAAR subtype [18]. The identities of the G proteins which are coupled to these receptors and their downstream effectors have also not been established although there is growing evidence that changes in tyrosine phosphorylation may play a critical role [2]. Interestingly, tyrosine kinases and tyrosine phosphatases are known to modulate ionic channels in several cell types, including the heart, possibly by modulating the subcellular distribution of channel proteins and/or their association with ancillary proteins [24, 25]. In the current context it would be particularly interesting to assess whether T1AM can influence the interaction of the ryanodine receptor with FKBP12, a protein thought to regulate the diastolic calcium leak [19].

Several limitations of our investigation must be acknowledged. Electrophysiological recordings were performed in a single cell type, and were not integrated with single channel recordings. In addition, we did not investigate sodium/calcium exchange nor mitochondrial calcium transport. Sodium/calcium exchange might indirectly affect the SR calcium pool because the exchanger competes with the SR calcium-ATPase for cytosolic calcium. Mitochondria are present in unfractionated preparations. They are known to both accumulate and release calcium although they are believed to play a minor role in the short-term regulation of calcium homeostasis in vivo [19], and their contribution to the calcium fluxes measured in our in vitro preparation should be negligible, because calcium uptake was measured in the presence of oxalate and calcium release was measured in the absence of sodium. Finally, we did not evaluate the calcium buffering capacity of the cytosol which could play a major role in determining free calcium concentration given that less than 1% of cytosolic calcium is unbound [19].

In conclusion, we have observed that T1AM signalling modulates calcium homeostasis and repolarizing potassium currents. These actions may account for its negative inotropic and possibly for its negative chronotropic action, leading to functional effects which are opposite those produced on a longer time scale by thyroid hormone. At present it is difficult to assess the physiological role of this aminergic system because the physiological concentration of myocardial T1AM is uncertain; pilot assays suggest average concentration to be in the mid-nanomolar range – i.e. 20-fold and 2-fold higher than 3,5,3'-triiodothyronine and thyroxine content – but the concentration at receptor level might be substantially higher [22]. Changes in local T1AM concentration might contribute to the cardiac manifestations of thyroid disease, and might be involved in pathological conditions such as heart failure, in which the role of thyroid hormone is certainly relevant but remains to be comprehensively clarified [26, 27]. In any case pharmacological manipulation of this interesting new signalling system could provide an exciting new approach to modulating cardiac function.

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