Effects of proarrhythmic drugs on relaxation time and beating pattern in rat engineered heart tissue

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Received: 7 March 2014 / Revised: 27 August 2014 / Accepted: 29 August 2014 / Published online: 11 September 2014
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Abstract The assessment of proarrhythmic risks of drugs remains challenging. To evaluate the suitability of rat engineered heart tissue (EHT) for detecting proarrhythmic effects. We monitored drug effects on spontaneous contractile activity and, in selected cases, on action potentials (sharp microelectrode) and Ca$^{2+}$ transients (Fura-2) and contraction under electrical pacing. The $I_{\text{to}}$-blocker inhibitor 4-aminopyridine increased action potential duration and $T_2$ and caused aftercontractions, which were abolished by inhibitors of ryanodine receptors (RyR2; JTV-519) or sodium calcium exchanger (NCX; SEA0400). 77 Drugs were then tested at 1-10-100 $\times$ free therapeutic plasma concentrations (FTPC): Inhibitors of $I_{\text{Kr}}$, $I_{\text{ks}}$, $I_{\text{to}}$, antiarrhythmics (8), drugs withdrawn from market for torsades des points arrhythmias (TdP, 5), drugs with measurable (7) or isolated TdP incidence (13), drugs considered safe (14), 28 new chemical entities (NCE). Inhibitors of $I_{\text{Kr}}$ or $I_{\text{ks}}$ had no effect alone, but substantially prolonged relaxation time ($T_2$) when combined at high concentration. 15/33 drugs associated with TdP and 6/14 drugs considered non-torsadogenic (cibenzoline, diltiazem, ebastine, ketoconazole, moxifloxacin, and phenytoin) induced concentration-dependent $T_2$ prolongations (10-100 $\times$ FTPC). Bepridil, desipramine, imipramine, thioridazine, and erythromycin induced irregular beating. Three NCE prolonged $T_2$, one reduced force. Drugs inhibiting repolarization prolong relaxation in rat EHTs and cause aftercontractions involving RyR2 and NCX. Insensitivity to $I_{\text{Kr}}$ inhibitors makes rat EHTs unsuitable as general proarrhythmia screen, but favors detection of effects on $I_{\text{to}}$, $I_{\text{ks}} + I_{\text{to}}$ or $I_{\text{ks}} + I_{\text{Kr}}$. Screening a large panel of drugs suggests that effects on these currents, in addition to $I_{\text{Kr}}$, are more common than anticipated.

Keywords Arrhythmia · Torsades des points · Drugs · In vitro screening · Engineered heart tissue

Introduction

Proarrhythmic side effects of drugs can be life-threatening and have led to a number of drug withdrawals from the market [17]. Predicting the arrhythmogenic potential in preclinical drug development is difficult for several reasons [25, 37]. (1) Mechanisms of arrhythmias are complex. This is exemplified by the fact that not only loss-of-function, but also gain-of-function mutations of K$^+$- and Na$^+$-channels...
can cause arrhythmias by promoting ectopic activity, dispersion, and/or re-entry circuits [1]. This corresponds with clinical experiences that both class I (Na+-channel blockers) and class III (K+-channel blockers) antiarrhythmic drugs have significant proarrhythmic effects [10]. In addition, dysfunction of the sarcoplasmic reticulum Ca2+-release channel (RyR2) or the SR Ca2+-storage protein calsequestrin underlie catecholaminergic polymorphic ventricular tachycardia, characterized by increased ectopic automaticity in situations of stress. (2) The heart is equipped with several safety mechanisms, explaining why one hit is rarely sufficient to cause symptomatic arrhythmias. Even patients with inherited rhythm disorders experience clinically relevant arrhythmias relatively late in life and/or only under certain trigger situations such as increased sympathetic drive, hypokalemia, drugs, ischemia, or myocardial scars. (3) The existing preclinical test systems have shortcomings [14, 25, 37]. In current routine, new chemical entities (NCE) are tested on cells over-expressing the human eag-related gene (hERG), and those with significant inhibitory activity are excluded from further development. Whereas hERG-tests have documented high sensitivity and specificity for this single ion current and the principal relevance for hERG-inhibition in causing torsades des pointes (TdP) is undisputed, the predictive value of this test is limited [21, 31]. Several drugs have hERG-inhibitory activity without being associated with TdP-arrhythmias (e.g., verapamil). Others have no relevant hERG-activity at clinically used concentrations, but increase the risk of arrhythmias (e.g., mefloquine and phenytoin; [31]).

The Food and Drug Administration and the European Medicines Agency currently recommend an integrated risk assessment, which includes the results of several experimental tests (e.g., hERG, rabbit Purkinje fibers, and dog telemetry) as well as in silico and clinical data [11, 12, 38]. The content of current tests may be improved by characterizing NCEs in a whole panel of cell lines, each expressing a different cloned ion channel. This approach provides a more comprehensive picture of a drugs channel-affecting activity [3, 42], but needs modeling to predict the integrated effect. An alternative strategy is to test drugs directly on cardiac myocytes or more complex cardiac tissues as the “real substrate” for arrhythmias, assuming that any reproducible effect is relevant, independent of its exact mechanism. Unfortunately, isolated adult cardiac myocytes do not beat and, similar to Purkinje fibers or Langendorff-perfused hearts, cannot be examined in large series. Instrumented dogs or rabbits are the most valid models, but cannot be used for screening large number of NCEs, both for ethical and financial reasons.

We have recently developed an automated miniaturized drug screening assay based on our EHT technology and neonatal rat cardiac myocytes, which appears to combine some of the advantages of a relatively intact 3D cardiac tissue, availability at large numbers, robustness and high-content readout, particularly analysis of contractile force and kinetics [7, 16, 20]. Experiments with a limited number of model compounds indicated assay sensitivity to detect proarrhythmic effects of drugs [16]. The aim of the present study was to systematically determine the predictive value of the assay by testing a larger number of clinically used compounds with characterized hERG-inhibitory activity and proarrhythmic potential in humans [31] and randomly picked NCEs as well as underlying mechanisms.

Materials and methods

The investigation conforms to the guide for the care and use of laboratory animals published by the NIH (Publication No. 85-23, revised 1985). A detailed description of methods can be found in the supplemental file.

Cell isolation and EHT-generation

Heart cells of postnatal d0–d3 Wistar rats and EHTs were prepared as previously described [16, 44]. Fresh neonatal heart cells were mixed with medium, fibrinogen and thrombin, and casted into strip-format (12 × 3 × 3 mm) molds in agarose, in which pairs of elastic silicone posts were placed from above. EHTs maintained for up to 4 weeks.

Measurement of contractile parameters

Contractile parameters were evaluated as previously described [16]. In principle, the 24-well-plates with EHTs (14–21 days) were put in a gas-, temperature-, and humidity-controlled incubator with glass roof and customized software-controlled video camera placed on top. Contractile parameters of spontaneously beating EHTs were evaluated using an automated figure recognition algorithm. Deflection of the silicone posts was recorded over time and, based on post geometry and elastic modulus of the silicone, used to calculate force, frequency, fractional shortening, contraction and relaxation time (bpm, T1 [from 20 % to peak] and T2 [from peak to 20 %], respectively).

Drug screening—video optical analysis

All measurements were performed with 14–21 day old EHTs in fresh serum-free DMEM (Biochrom F04115), supplemented with 10 mM HEPES for pH-steadiness, pre-incubated at 37 °C, 40 % O2, 7 % CO2, 90 % humidity for 2 h. Measurements were done routinely 1 day after feeding.
with standard EHT medium. The drugs were analyzed in three different concentrations (45 min each, cumulative, 1–100× free therapeutic plasma concentration [FTPC]). For drug details including solvents see Supplement Table 1. Prior to each measurement, 50 nM epinephrine (Sigma E4643) was added to each well to simulate “physiological” conditions and enhance the likelihood of contractile activity within the 60 s recording time. Drugs were added under sterile conditions. After 45 min incubation in a standard incubator, EHTs were transferred to the video optical system (Fig. 1). Due to a sequential mode of measurement (total time ~30 min), incubation time varied from the first to the last EHT (45–75 min).

Measurements under perfusion and electrical stimulation, Ca^{2+}-transients, action potentials

Intracellular Ca^{2+}-transients were analyzed in parallel with force under electrical stimulation and continuous perfusion using a novel setup as described previously [35]. The setup consisted of an inverted microscope, a temperature- and O₂/CO₂-controlled chamber for the 24-well EHT plate, a flow rate-controlled perfusion system, platinum–iridium wire electrodes for field stimulation, a fluorescence light source (IonOptix Hyperswitch), a photomultiplier, video cameras and software (both IonOptix) for the evaluation of contractile activity (edge detection mode). Experiments were done at 4 ml flow/min (per well) and 2–4 Hz stimulation in modified Tyrode’s solution.

For Fura loading, EHTs were incubated in Tyrode’s solution containing 10 μM Fura2-AM (Invitrogen F1221) and Cremophor EL (0.75 %; Sigma C5135) for 2 h at 37 °C. The ratio of light emission (510 nm) at excitation with 340 and 380 nm light (F340/380 ratio) was used as an index of cytosolic Ca^{2+}-concentration.

Action potentials (APs) were recorded with standard intracellular microelectrodes in intact EHTs. Bath solution contained (in mM): NaCl 127, KCl 4.5, MgCl₂ 1.5, CaCl₂ 1.8, glucose 10, NaHCO₃ 22, NaHPO₄ 0.42, equilibrated with O₂-CO₂ [95:5] at 36.5 ± 0.5 °C, pH 7.4. Preparations were field-stimulated for at least 1 h (2 Hz) before data acquisition. APs were analyzed off-line using the LabChart® software (ADInstruments, Spechbach, Germany).

Statistical analysis

Data were expressed as mean ± SEM. Statistical differences were analyzed using the one-way analysis of variance (ANOVA) followed by the Dunett’s (all compared to baseline) or Tukey’s (all compared to all) adjustment for post hoc multiple comparison, or by paired or unpaired Student’s t test, as indicated in the legend of each figure. Results were considered statistically significant if a paired Student’s t test revealed a p value of less than 0.05 and the deviation from baseline was at least 15 %. This limit was defined after initial series of experiments had shown that formally significant (t test), but not concentration-dependent effects of drugs often amounted to ±11 %. Further support for the 15 % threshold came from quantifying the mean ± SD of all baseline measurements (n = 221 independent EHTs), which amounted to 99.6 ± 11.4 % (SEM ± 0.77 %).

Results

Mechanisms of twitch prolongation and irregular beating pattern

In a previous study with EHTs, we found that the experimental I_{Kr}-inhibitor chromanol 293b and two drugs known
to inhibit I\textsubscript{Kr} and cause TdP in humans, quinidine and erythromycin, caused concentration-dependent prolongations of relaxation time T2 [16]. These observations suggested that T2 is a useful surrogate for drug-induced prolongations of repolarization and proarrhythmic effects. To test this hypothesis, we evaluated the effects of the I\textsubscript{Kr}-inhibitor 4-aminopyridine (4AP; IC\textsubscript{50} on I\textsubscript{Kr} in adult rat ventricular myocytes: 980 \mu M [41]) on EHTs. 4AP prolonged T2 concentration dependently reaching significance at 3 mM (Fig. 2). At 30 mM, EHTs showed extremely prolonged relaxation (+502 %), comparable to what have been seen previously with chromanol (+710 %; [16]). Prolonged contractions can either be caused by altered intracellular Ca\textsuperscript{2+} transients or myofilament response to Ca\textsuperscript{2+}. To differentiate between these mechanisms, EHTs were subjected to sharp electrode measurements of action potentials and calcium transients (Fura-2; Fig. 3). Action potential characteristics under basal conditions and electrical stimulation were similar as described previously in this model [16]. 4AP increased action potential duration (APD\textsubscript{90}) from a mean of 113 to 175 ms (Fig. 3a, b). 4AP also prolonged intracellular Ca\textsuperscript{2+} transients, whereas the \beta-adrenergic agonist epinephrine shortened it, both in the absence and presence of 4AP (Fig. 3c–f). Quinidine, a still prescribed drug for the treatment of atrial fibrillation, also increased APD\textsubscript{90} from 179 to 232 ms at 100 \mu M (Supplemental Fig. 5d). These data supported the interpretation that alterations in T2 reflect similar changes in APD and the kinetics of intracellular Ca\textsuperscript{2+} transients. Accordingly, a high concentration of caffeine, known to open ryanodine receptors (RyR2) in the sarcoplasmic reticulum (SR), reduced beating rate and induced a large and wide twitch (Supplemental Fig. 1).

4AP not only prolonged T2, but also induced beat-to-beat irregularities, variations of twitch amplitude and aftercontractions falling into the relaxation phase of prolonged twitches (Fig. 4). To investigate the role of different cellular effector systems in the T2-prolonging effect of repolarization-prolonging compounds, we determined the effect of 4AP in the absence and presence of tetrodotoxin (INa), tetracaine (INa\textsubscript{L}), verapamil (ICa), thapsigargin (SERCA), SEA0400 (sodium calcium exchanger, NCX) or JTV519.

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**Fig. 2** Concentration-dependent effects of the I\textsubscript{Kr}-blocker 4-aminopyridine (4AP).

a–e Representative original recordings of spontaneously beating EHTs in the absence (baseline, only epinephrine 50 nM) and presence of increasing concentrations of 4AP. f Statistical evaluation of relaxation time (T2; each dot represents one analyzed EHT; **p < 0.003, ***p < 0.0001 vs. baseline one-way ANOVA + Dunnett’s post test). The concentration of 4AP was cumulatively increased with 45 min incubation steps per concentration. Note the concentration-dependent prolongation of contraction twitches and the aftercontractions at high concentrations.
Verapamil had no discernible effect, thapsigargin, tetrodotoxin and tetracaine all increased the T2-prolonging effect of 4AP (Supplemental Fig. 2). In contrast, SEA0400 and JTV519 reduced the effect of 4AP (Fig. 4). Both compounds completely abolished beating irregularity and aftercontractions, but only partially T2-prolongation. T2 values were 127 (SEA + 4AP) vs. 78 ms (SEA alone) and 195 (JTV + 4AP) vs. 80 ms (JTV alone). This suggests that part of the 4AP-induced T2-prolongation was a direct consequence of action potential prolongation, whereas the SEA- and JTV-sensitive after contractions and pronounced T2-prolongations were mediated by RyR2 and NCX. Similar data were obtained with thioridazine (30 µM, data not shown).

Fig. 3 Effect of 4-aminopyridine (4AP; 10 mM), epinephrine (50 nM) or their combination on action potential duration (APD₉₀), Fura-2 F340/380 ratio and relaxation time (T2) of electrically stimulated EHTs (2 Hz). For APD₉₀ measurements two independent groups of EHTs (n = 3, each) were directly perfused with vehicle control (VC) or 4-aminopyridine (4AP), electrically stimulated and contractions recorded. Preparations were stimulated for at least 1 h before drug exposure and data acquisition. For Fura-2 F340/380 ratio and T2 the two independent groups (n = 8, each) were preincubated with VC or 4AP. Thereafter, EHTs were perfused and electrically stimulated (2 Hz; baseline) for 10 min, before epinephrine (50 nM) was added to the perfusion system. a Average peaks for action potentials in the absence (VC, black line) and presence of 4AP (red line). b Statistical evaluation of APD₉₀. c Original recordings of F340/380 ratio transients in the absence (VC, left) and presence of epinephrine (right). d Original recordings in the presence of 4AP alone (left) and 4AP plus epinephrine (right). The red line displays the unfiltered signal, whereas the blue line displays the filtered signal. e Statistical evaluation of time to 90 % return of the F340/380 ratio and F T2. Each point represents one EHT. *p < 0.05, **p < 0.003, ***p < 0.0001, Student’s t test, paired for comparison within one group (baseline vs. epinephrine) and unpaired for comparison between the two groups (VC vs. 4AP); y-axes for C, D: F340/380 ratio in arbitrary units (AU). Note that 4AP increased T2 and time to 90 % return of the F340/380 ratio, both in the absence and presence of epinephrine. Epinephrine alone shortened T2 and F340/380 transients. Note also the failure of capture under 4AP (RyR2), respectively. Verapamil had no discernible effect, thapsigargin, tetrodotoxin and tetracaine all increased the T2-prolonging effect of 4AP (Supplemental Fig. 2). In contrast, SEA0400 and JTV519 reduced the effect of 4AP (Fig. 4). Both compounds completely abolished beating irregularity and aftercontractions, but only partially T2-prolongation. T2 values were 127 (SEA + 4AP) vs. 78 ms (SEA alone) and 195 (JTV + 4AP) vs. 80 ms (JTV alone). This suggests that part of the 4AP-induced T2-prolongation was a direct consequence of action potential prolongation, whereas the SEA- and JTV-sensitive after contractions and pronounced T2-prolongations were mediated by RyR2 and NCX. Similar data were obtained with thioridazine (30 µM, data not shown).
Role of IKr and IKs in T2 prolongations

The role of IKr and IKs for action potential repolarization in rat heart is still poorly understood [32]. We studied the involvement of these two currents in rat EHTs by applying the reference IKr blocker E-4031 (IC50 7.7 nM [43]) and the selective IKs blocker HMR-1556 (IC50 10.5 nM [36]) alone or in combination (1–1,000 nM, Fig. 5). Neither E-4031 nor HMR-1556 affected T2 even at high concentrations (1,000 nM). The combined application also did not affect T2 at up to 100 nM, but caused a substantial increase at 1,000 nM (8-fold). Given the high selectivity of E-4031 and HMR-1556 for IKr and IKs (HMR-1556 IC50 on Ito: 33.9 μM; on Ica: 27.5 μM; on IKr: 12.6 μM [36]), respectively, the effect of the combination suggests a role of these two currents for determining the repolarization reserve in rat EHTs.

Screening of proarrhythmic compounds under spontaneous beating

To test the usefulness of our screening system for the detection of proarrhythmic compounds, we analyzed a large panel of drugs associated with arrhythmias. The selection was made according to a list of drugs published by Redfern and colleagues [31] which related various levels of proarrhythmic risk with inhibition of IKr (hERG). We tested the effect of the 46 compounds of this list which were commercially available (details Supplemental Table 1) at 1-, 10-, and 100-fold FTPC, n = 4–8 each. In addition, we tested moxifloxacin, an important inhibitor of bacterial gyrases, associated with prolongation of the QT-interval, but not with arrhythmias [28]. The selection of compounds encompassed 8 clinically used antiarrhythmic drugs (Group I), 5 drugs withdrawn from the market for TdP (Group II), 7 drugs with measurable incidence of TdP in humans (Group III), 13 drugs with isolated reports of TdP (Group IV) and 14 drugs devoid of TdP reports and therefore considered safe (Group V; Figs. 6, 7). Under the experimental conditions (which included a *EC50 epinephrine concentration, 50 nM) and in the absence of interventional drugs, EHTs showed a typical and reproducible beating pattern, consisting of periods with high frequency (4.4–4.6 Hz, called “bursts”, 8–10 s length) and periods of contractile inactivity (30–40 s; Supplemental Fig. 3). Single twitch kinetics showed a mean ± SD
T1 and T2 of 67 ± 8 (±1) and 92 ± 18 (±2) ms, respectively. Given that the bursts occurred by chance either fully inside the 60 s recording window or only partially, the total number of beats per 60 s was relatively meaningless. In contrast, the frequency of beating in the burst was stable and systematically affected by epinephrine (715–20 %) and carbachol (reversed epinephrine effect and induced partial stop; Supplemental Fig. 3).

Given our previous results with erythromycin and quinidine [16], we focused on T1, T2, and the beating pattern as possible surrogates for arrhythmias. A threshold of 15% T2 prolongation was considered significant (see Statistics for details). Supplemental Fig. 4 depicts typical examples of drug responses, Figs. 6 and 7 show the T2 effect of 47 drugs according to the Redfern classification. Significant T2 prolongations were observed with 4/8 Redfern Class I drugs (antiarrhythmics; quinidine, disopyramide, sotalol, procanamide; Fig. 6a), 1/5 drugs withdrawn from the market due to TdP (Redfern Class II; cisapride; Fig. 6b), 4/7 drugs with measurable incidence of TdP (Redfern Class III; aprindine, bepridil, flecaïnide, erythromycin; Fig. 6c), 6/13 drugs with isolated TdP reports (Redfern Class IV; domperidone, sparflaxacin, propafenone, desipramine, fexofenadine, clarithromycin; Fig. 7a), and 6/14 drugs considered safe (ebastine, ketoconazole, diltiazem, cibenzoline, phenytoin, moxifloxacin; Fig. 7b). Whereas most drugs prolonged T2 only at 100× FTPC, phenytoin exerted this effect already at 1× FTPC. In addition, five compounds (bepridil, desipramine, imipramine, thioridazine, and erythromycin) induced irregular beating as the primary effect, characterized by a “sinusoidal” fluctuation of twitch amplitudes (Supplemental Fig. 4a). The effect was time dependent and was followed by complete stop of beating. It occurred at concentrations of 30–100 fold FTPC, i.e., 3–10 (−600 in case of thioridazine) fold hERG IC50. Bepridil, desipramine, and erythromycin increased T2 in addition to inducing beating irregularity.

In aggregate, the rat EHT responses categorized the 47 drugs in 4 groups (Fig. 8). Group 1 caused irregular beating as the main effect (n = 5), group 2 induced a concentration-dependent increase in T2 (n = 18). T2-effective concentrations were 5 fold (domperidone, disopyramide) to 100 fold FTPC (e.g., ebastine, diltiazem, sotalol, or moxifloxacin). Group 3 (n = 7) had variable effects on EHT contraction including prolongation of contraction time T1 (haloperidol, sertindole, diphenhydramine, and mefloquine), shortening of T2 (terfenadine) and a decrease in force (verapamil and astemizole). The negative inotropic effect of verapamil
was already seen at 300 nM, approximately 4-fold FTPC. Group 4 encompassed 17 compounds without any effect on the parameters studied. It contained specific IKr-blockers such as ibutilide and dofetilide, which was in contrast to the results obtained with human EHTs [33].

Effects of drugs under electrical pacing

Drugs which caused prolongations of T2 also reduced beating rate in the bursts in many cases (Supplemental Fig. 4), raising the question which effect is the cause and which the consequence. We therefore measured EHT contractility under continuous electrical stimulation and perfusion and carefully evaluated the time course of the effects of quinidine (100 μM) and erythromycin (1 mM; Supplemental Fig. 5). Quinidine first induced a marked, time-dependent prolongation of T2 and a reduction in force, which was then followed by the development of a slower rhythm independent of pacing. Erythromycin also first prolonged T2, reduced force and then induced an autonomous chaotic contraction pattern. Effects of both drugs were fully reversible after washout. The data suggest that the drugs exert primary effects on processes underlying relaxation and that slowing of rate is a consequence.

Screening of new chemical entities (NCEs) under spontaneous beating

To get a rough estimate of the frequency of rat EHT effects in a non-selected group of drugs a chemical library was analyzed. NCEs were purchased from Maybridge and randomly chosen by respecting the rule of five [26]. 28
NCEs were tested at 0.1, 1, and 10 µM. Most NCEs had no effect on EHT contractility. Three NCEs prolonged T2 and one reduced force. In two additional cases EHTs stopped beating and didn’t react to electrical pacing (Supplemental Table 2). Overall, an effect on EHT contractility was observed in 21 % of the investigated NCEs.

Discussion

Assessing the proarrhythmic risks of drugs remains a challenge in preclinical drug development. Current experimental models in preclinical toxicology determine the effect of NCEs on ion channels commonly involved in arrhythmogenic drug effects, particularly hERG channels [39], on electrophysiological surrogates of arrhythmias such as triangulation in Langendorff-perfused rabbit hearts [23, 24], on action potential duration in rabbit Purkinje fibers [15] or the QTc interval in telemetrically surveyed dogs [13]. Newer models with higher throughput and/or a human cardiomyocyte context include measurements of heart rate in zebrafish [27], of Ca²⁺ transients in isolated guinea-pig [30] or human pluripotent stem cell (hPSC)-derived cardiomyocytes [5] or of electric field potentials in hPSC-cardiomyocytes [2]. The present study in rat EHTs is, to the best of our knowledge, the largest head-to-head comparison of proarrhythmic drugs performed so far. It showed that time of relaxation (T2) in rat EHTs is essentially insensitive to selective blockers of I Kr or I Ks, but prolonged by inhibition of I to, combined full inhibition of IKr and IKs or combined inhibition of IKs and Ito. This suggests that rat EHTs monitor mainly Ito and IKs effects of drugs. The Ito-blocker 4AP also prolonged action potential duration and Ca²⁺ transients at T2-effective concentrations, indicating that, in this experimental model, T2 prolongation follow similar prolongations of repolarization and Ca²⁺ transients. The data suggest that, for drug screening purposes, T2 is a useful surrogate for time of repolarization in rat EHTs. The high percentage of known proarrhythmic drugs that induced concentration-dependent T2 prolongations, aftercontractions and/or irregular beating (group 1 and 2, 23/38 = 61 %) indicates that many clinically used drugs possess effects on Ito and/or IKs in addition to their well-characterized inhibitory action of IKr (hERG).

Characterization of rat EHTs as a test system for testing proarrhythmic drugs

The underlying hypothesis of this study was that repolarization-inhibiting drugs prolong action potential duration
Fig. 8  Grouping of drugs according to the type and the concentration-dependence of the effects on EHTs. All drugs were tested in 1×, 10×, and 100× free therapeutic plasma concentration (FTPC) in the presence of 50 nM epinephrine. The absolute concentration range is indicated in the first row. Group 1 encompasses drugs (red) that induced polymorphic arrhythmias, group 2 (orange) prolongations of relaxation (T2), group 3 prolongations of contraction (T1, yellow), shortening of T2 (blue), or negative inotropic effects (violet). Group 4 (white) drugs exerted no significant effect. The colored bars in group 1–3 range from the lowest tested concentration in which the indicated effect occurred (generally 10–100× FTPC) to 3 mM, regardless of whether this concentration has been tested. The white bars in group 4 indicate the tested concentration range. Also, indicated are hERG/I Kr lowest published IC50 upward; lowest published IC50 values on Ino upward; lowest published IC50 values on IKr upward; concentrations associated with 10–20% increases in APD90; with decreases in APD90; QTc-prolongation in vivo; increased QTc in humans; estimated plasma concentration (ep) for TdP case reports in humans; ep for TdP case reports in combination with other drugs/diseases; ep for TdP case reports in combination with a cytochrom P450 inhibitor; ep for TdP case reports in children; ep QTc case reports; asterisk TdP in humans known; double asterisk other types of arrhythmias in humans known. The type of depiction and most information independent of EHTs were adopted from Redfern et al. (2003). Case reports (indicated by asterisk), estimated plasma concentration and IC50 values on Ino/IKr were obtained by a literature search.
and Ca$^{2+}$ influx and thereby time of relaxation, a parameter easily assessable in EHT. EHTs are engineered three-dimensional cardiac tissue constructs in which cardiac cells are embedded in a fibrin matrix and, over 10–14 days, form a highly organized longitudinally oriented network [16]. The 24-well mini-EHTs system is designed for a robust, automated measurement of spontaneous or electrically stimulated contractile activity at a medium throughput scale. It is stable for weeks and has the advantage that measurements can be done at near-physiological conditions: 37 °C, steady state beating (not dying preparations such as isolated myocytes, papillary muscles, Purkinje fibers, and Langendorff hearts), auxotonically work-performing heart muscle constructs, and absence of invasive experimental interventions (e.g., patch clamp or micro-electrodes). On the other hand, it is not well suited for measuring effects of drugs on action potentials, field potentials, or ion channels directly. We therefore performed several experiments to evaluate whether the relaxation time T2 is indeed a suitable surrogate of time of repolarization.

The following observations favor this assumption. (1) T2 is easily determined and highly reproducible (11 % SD, 0.77 % SEM, n = 221). (2) The duration of contraction and relaxation corresponds well with that of action potentials in rat EHTs and both were prolonged in the presence of 4AP as well as quinidine. Action potential duration (APD$_{90}$) measured with sharp microelectrodes at 36 °C amounted to a mean of 113 ms, time to 90 % Ca$^{2+}$ return to a mean of 124 ms at 37 °C, and T1 + T2 (at 80 % relaxation) amounted to a mean of 159 ms at 37 °C (this study and [16]). Thus, it is reasonable to assume that prolongations of repolarization affect T2. (3) Drugs with well-characterized actions on I$_{to}$ [41], I$_{Kr}$ [43], and I$_{Ks}$ [36] had reproducible, concentration-dependent and reversible effects on T2. 4AP started to prolong T2 at 3 mM, which corresponds to 3-fold IC$_{50}$ in adult rat cardiac myocytes [41] and confirms the prominent role of I$_{to}$ for rodent heart repolarization [19]. The lack of effect of E-4031, doftelide, ibutilide, and HMR-1556 even at 100-fold IC$_{50}$ concentrations confirms the minor role of I$_{Kr}$ and I$_{Ks}$ for repolarization in rat cardiac myocytes [29]. Interestingly, however, the combined application of high concentrations of E-4031 and HMR-1556 markedly prolonged T2. Given the high selectivity of both drugs for I$_{Kr}$ and I$_{Ks}$ [36, 41], respectively, the data suggest that I$_{Kr}$ and I$_{Ks}$ can substitute for each other, but together play a role for repolarization in rat EHTs. The strong effect of HMR-1556 in the presence of 3 mM 4AP (and the lack of an E-4031 effect under this condition) suggests a greater role for I$_{Ks}$ than I$_{Kr}$. The necessary channel subunits are expressed in rat EHTs (KCNH2a, KCNQ1a, and KCNE1/E2 mRNAs are present in rat EHTs [16]).

Whereas all this argues for repolarization as an important parameter of relaxation time T2 in our system, it is obvious that other mechanisms have to be considered and that T2 prolongations alone do not prove effects on repolarization. Any effect on beating rate will affect T2 because of the well-known frequency-dependent acceleration of relaxation. Such effect could be excluded in case of quinidine and erythromycin (Supplemental Fig. 5), but may play a role in other cases. Drugs that directly affect intracellular Ca$^{2+}$ handling could alter T2 independently of repolarization. Indeed, caffeine at a high concentration (5 mM) which enhances the open probability of RyR2 reduced beating rate and caused widening of contraction twitches (Supplemental Fig. 1). Thapsigargin, a selective inhibitor of SERCA, did not affect time of relaxation as such (3–30 nM), but aggravated the relaxation-slowing effect of 4AP (Supplemental Fig. 2). Another possibility to affect T2 independently of repolarization is an effect on myofilament Ca$^{2+}$ sensitivity. Ca$^{2+}$ sensitizers such as EMD 57033 [34] shift the force-pCa curve to the left and thereby prolong relaxation as recently shown in mouse EHTs [35]. The unexpected T2-prolongation under diltiazem may represent this mechanism. A study in isolated adult rat ventricular myocytes reported a mild myofilament Ca$^{2+}$ sensitizing effect of diltiazem [9]. Such an effect may also explain the unexpected lack of significant negative inotropic effects of this compound at the highest tested concentration (10 μM) which was in contrast to verapamil. Conversely, epinephrine (Supplemental Fig. 3) and all other cAMP-dependent drugs increase the PKA-dependent phosphorylation of troponin I, myosin binding protein C and others and shift the curve to the right. This favors relaxation and abbreviates T2. Taken together, T2-prolongation of rat EHTs can be caused not only by blockers of repolarization (mainly I$_{to}$, I$_{Kr}$, very little contribution of I$_{Ks}$), but also rate-slowing drugs, Ca$^{2+}$ sensitizers or blockers of cAMP-dependent pathways such as carbachol (Supplemental Fig. 3), which needs to be considered in drug screening efforts.

The pronounced prolongation of relaxation at high concentrations of 4AP and aftercontractions were sensitive to both SEA0400 and JTV519 (Fig. 4), but relaxation remained significantly prolonged in their presence (~200 %). This finding is interesting as it suggests that the primary effect of 4AP, the inhibition of repolarization, has time-dependent (see time course in Fig. 4) secondary effects that affect time of relaxation. Very likely, this secondary effect corresponds to increased filling of the SR with Ca$^{2+}$ with the final consequence of RyR2-mediated spontaneous Ca$^{2+}$ release and NCX-mediated Ca$^{2+}$ extrusion. The latter transport is electrogenic and contributes to prolonged depolarization and after depolarizations, visible in our system as marked prolongations of T2 and
aftercontractions. An interesting speculation is that group 1 (Fig. 8) drugs directly interfere with RyR2 and/or NCX or SR function and therefore induce irregular beating, whereas group 2 drugs primarily affect repolarization and therefore prolong T2 before causing extra beats. On the other side, group 1 and 2 effects cannot be firmly separated because three group 1 drugs also caused T2 prolongations. Interestingly, tetracaine (also flecainide (0.5 μM); data not shown) did not mimic the effect of JTV519 (Supplemental Fig. 2), although both have inhibitory effects on RyR2-mediated Ca\(^{2+}\) release [18, 40]. Possibly, their main effect on \(I_{\text{Na}}\) which is not shared by JTV519 could explain the difference.

Drug-induced relaxation slowing and beating irregularities in rat EHTs

We selected drugs according to a list published by Redfern and colleagues [31] that grouped drugs according to their proarrhythmic potential. 15/33 drugs with various degree of risk for TdP (Class I–IV) showed T2 prolongation or beating irregularities in our system (group 1 and 2). The percentage did not clearly differ between those that were withdrawn from the market for TdP (1/5) and those with only isolated TdP reports (6/13). In addition, T2-prolongation was observed in 6/14 drugs considered safe. This clearly indicates that the rat EHTs test does not sufficiently discriminate between high and low risk drugs, very likely due to the lack of \(I_{\text{Kr}}\) sensitivity.

Nevertheless, screening the large number of drugs revealed a number of interesting novel informations. (1) It is apparent that T2 prolongations and beating irregularities were seen mainly in those drugs which reach high FTPC in clinical use and were therefore tested at high absolute concentrations (compare concentration range of Group 1 and 2 drugs with group 3 and 4; Fig. 8). This confirms the general rule in pharmacology that drugs with low potency at their target have a higher chance of off-target toxicity than high affinity drugs. (2) Most effects were seen at 30–100-fold FTPC and higher. This is at the upper limit generally considered a critical safety margin [31] and point to a relatively low sensitivity of our assay. Rather than indicating simple non-specific effects (drugs such as sematilide and ciprofloxacin were tested at up to 300 μM) the data with E-4031, HMR-1556 and 4AP indicate that almost complete block of one or more currents is required to overcome safety mechanisms of repolarization in a relatively intact system such as the EHT. As such, the low sensitivity probably reflects the situation in vivo better than isolated cells [28]. Pharmacokinetic peculiarities and time-dependent effects likely add to high concentration requirements. For example, amiodarone is very lipophilic and accumulates in cells over time. With an apparent volume of distribution of 20–200 l/kg, the low FTPC of amiodarone (0.3 nM) is orders of magnitude lower than cellular concentrations in the steady state. (3) Concentrations in which group 1 and 2 drugs exerted their effects on EHTs were generally well above their IC\(_{50}\) for hERG (gray bars in Fig. 8). Notable exceptions are domperidone and phenytoin, where relaxation slowing occurred at concentrations below hERG IC\(_{50}\). Since E-4031 at high concentration markedly potentiated the effect of HMR-1556 without having an effect alone, the data suggest that inhibition of \(I_{\text{Kr}}\) may participate in the effect of group 1 and 2 drugs, but is not sufficient to fully explain them. Inhibition of other currents must come into play. We found reports of \(I_{\text{to}}\)-inhibiting activity for 14 drugs (blue bars in Fig. 8). In four of these (imipramine, propafenone, quinidine, and clarithromycin) published IC\(_{50}\) values for hERG and \(I_{\text{to}}\) were at or below the threshold for T2 prolongation or irregularity, providing a likely mechanism of action. Moreover, we found reports of \(I_{\text{Ks}}\)-inhibiting effects for 20 drugs (green bars in Fig. 8). In six of these cases (bepridil, thiordizine, propafenone, quinidine, cibenzoline, and sotalol), the IC\(_{50}\) was at or below the threshold for group 1 or 2 effects. It is interesting that quinidine at the T2-threshold concentration (100 μM) inhibits all three currents \(I_{\text{Kr}}, I_{\text{to}}\), and \(I_{\text{Ks}}\). Other currents involved in rat cardiac repolarization such as \(I_{\text{Kr}}, I_{\text{ss}},\) and \(I_{\text{Ks}}\) [19] or \(I_{\text{KATP}}\) [4] and \(I_{\text{KNa}}\) are also potential targets, but not much is known about effects of the drugs investigated. Taken together, the data indicate that many clinically used drugs, some of them associated with TdP, others not, inhibit cardiac repolarizing currents in addition to \(I_{\text{Kr}}\). This supports recent data suggesting that combined channel block underlies clinical proarrhythmia [28].

Miscellaneous effects on EHT function

Some drugs caused T1 prolongations (haloperidol, sertindole, diphenhydramine, and mefloquine), T2 acceleration (terfenadine) or negative inotropic effects (verapamil). Whereas the latter is the expected main effect of the drug, the mechanism of the other is unclear at present. T1 prolongations could indicate reduced conduction velocity in the EHT, a typical consequence of INa inhibition. In fact, terfenadine, which also shortened action potential duration in the SCREENIT test system (10 μM [23]), has recently been suggested to cause
arrhythmias not by a TdP mechanism, but by its strong I_{Na} blocking activity [46].

Taken together, rat EHTs are not suitable as a general screening assay for proarrhythmic drug effects due to the small contribution of I_{Ks} for rat EHT repolarization. On the other hand, the assay is a simple and robust system for the analysis of non-hERG-related drug effects on cardiac function. Relaxation time was shown to be a particularly suited screening parameter, sensitive to drugs affecting cardiac repolarization, but also Ca^{2+} handling or myofilament function. The high fraction of drugs with known arrhythmogenic effects that prolonged relaxation or induced irregular beating in this hERG-insensitive system suggest that I_{to} and I_{Ks} effects add to the proarrhythmic risk of drugs and require further consideration.

Acknowledgments We are grateful for the technical help by Bülent Aksehirlioglu, Hamburg, in constructing the EHT setup and to Prof. Ursula Ravens, University of Dresden, for the kind gift of HMR-1556.

Conflict of interest None.

Funding sources This study was supported by grants from the German Research Foundation (DFG Es/88-12), the European Commission (FP7 Angioscaff and FP7 Biodesign) and the DZHK (German Research Center for Cardiovascular Research), BMBF (German Ministry for Education and Research).

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Appendix

EHTs were adopted from Redfern et al. 2003. Case reports (indicated by asterisk), estimated plasma concentration and IC50 values on Ito/IKs were obtained by a literature search.

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