Highly Selective SERCA2a Activators: Preclinical Development of a Congeneric Group of First-in-Class Drug Leads against Heart Failure

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ABSTRACT: The stimulation of sarcoplasmic reticulum calcium ATPase SERCA2a emerged as a novel therapeutic strategy to efficiently improve overall cardiac function in heart failure (HF) with reduced arrhythmogenic risk. Istaroxime is a clinical-phase IIb compound with a double mechanism of action, Na+/K+ ATPase inhibition and SERCA2a stimulation. Starting from the observation that istaroxime metabolite PST3093 does not inhibit Na+/K+ ATPase while stimulates SERCA2a, we synthesized a series of bioisosteric PST3093 analogues devoid of Na+/K+ ATPase inhibitory activity. Most of them retained SERCA2a stimulatory action with nanomolar potency in cardiac preparations from healthy guinea pigs and streptozotocin (STZ)-treated rats. One compound was further characterized in isolated cardiomyocytes, confirming SERCA2a stimulation and in vivo showing a safety profile and improvement of cardiac performance following acute infusion in STZ rats. We identified a new class of selective SERCA2a activators as first-in-class drug candidates for HF treatment.

INTRODUCTION

Heart failure (HF) is a life-threatening syndrome characterized by an inability of the heart to meet the metabolic demands of the body. It is age-dependent, ranging from less than 2% of people younger than 60 years to more than 10% of individuals older than 75 years. Most patients with HF have a history of hypertension, coronary artery disease, cardiomyopathies, valve disease, or a combination of these disorders.1 The calculated lifetime risk of developing HF is expected to increase, and those with hypertension are at higher risk.2 Clinical symptoms in HF are caused by a cardiac double pathological feature that consists of diminished systolic emptying (systolic dysfunction) and impaired ability of the ventricles to receive blood from the venous system (diastolic dysfunction). The impaired contractility and relaxation are also caused by an abnormal distribution of intracellular Ca^{2+}, resulting from reduced Ca^{2+} uptake by the sarcoplasmic reticulum (SR).3 SR Ca^{2+} uptake is operated by the SR Ca^{2+} ATPase, SERCA2a, a 110 kDa membrane protein. During ion transport across the membrane, SERCA2a undergoes large conformational changes switching between Ca^{2+}-bound E1 and Ca^{2+}-free E2 states.4 SERCA2a activity is physiologically inhibited by the interaction with phospholamban (PLN), a small protein5 that stabilizes the E2 state, incompatible with Ca^{2+} binding. SERCA2a inhibition by PLN is normally relieved by PLN phosphorylation with protein kinase A, a signaling pathway severely depressed as a consequence of myocardial remodeling.8 SERCA2a activators are therefore promising drugs that might improve overall cardiac function in HF with reduced arrhythmogenic risk. Various therapeutic approaches that increase SERCA2a function have been recently investigated.9–13 Small-molecule SERCA2a activators have been recently discovered. Among them, a pyridone derivative directly binds to PLN displacing it from SERCA2a14 and a small molecule activates SERCA2a by promoting its SUMOylation [small ubiquitin-related modifier (SUMO)].15 Overall, new SERCA2a activators might be very useful in HF treatment together with the first line of therapeutic agents, β-blockers, and angiotensin-converting enzyme (ACE) inhibitors.

The work of our laboratory led to the successful completion of phase IIb clinical trials of the steroid derivative istaroxime,16 which is endowed with a double mechanism of action, that is, Na’/K’ ATPase inhibition17 and SERCA2a activation.18 Istaroxime is an inotropic/lusitropic agent, which is capable of improving both systolic and diastolic functions (HORIZON...
study)\textsuperscript{19} with a much lower proarrhythmic effect than digoxin, a pure \(\text{Na}^+ / \text{K}^+\) ATPase inhibitor.\textsuperscript{18} This suggests that, by improving \(\text{Ca}^{2+}\) clearance from the cytosol,\textsuperscript{20} SERCA2a stimulation may minimize the proarrhythmic effect of \(\text{Na}^+ / \text{K}^+\) ATPase blockade\textsuperscript{18,19,21} while preserving its inotropic effect.

Although having an excellent pharmacodynamic profile, istaroxime is not optimal for chronic administration because of its poor gastrointestinal absorption, high clearance rate, and extensive metabolic transformation\textsuperscript{19} which leads to the formation of a final metabolite, named PST3093. A recent study from our laboratory\textsuperscript{22} indicates that PST3093 behaves as a selective SERCA2a activator showing a longer half-life than istaroxime and it is likely to account for the lusitropic effect of istaroxime in patients; nonetheless, the presence of the oxime function may still limit the chronic usage of PST3093. The main aim of this work has been the rational design and synthesis of bioisosteric PST3093 analogues with metabolically stable groups replacing the oxime function, with the purpose to maintain the selective activity on SERCA2a. We present here the ligand-based rational design, the synthesis, and the biochemical and pharmacological in vitro and in vivo characterization of a new class of alkene-based PST3093 derivatives that turned out to have selective activity on SERCA2a.

\section*{RESULTS}

Ligand-Based Rational Design and Synthesis of PST3093-Derived Compounds. The metabolite PST3093 (Figure 1A) possesses a 17-androstanone core of istaroxime.
with two main structural differences: a carboxylic acid group instead of amino group on the C3-oxime linker and a hydroxyl group with R configuration (or β-configuration) at C6 instead of istaroxime’s carbonyl. A series of PST3093 variants (compounds 1–12, Figure 1B) has been designed by a ligand-based approach to carry a carboxylic group (acid or ethyl ester) attached through a linker to the C3 position of the 6-hydroxy-17-oxo androstane core. Although structural information and quantitative structure-activity relationship data allowed the identification of the pharmacophore of istaroxime derivatives that binds to Na+/K+ ATPase,16 no structural information is available on the interaction between compound PST3093 and SERCA2a and/or PLN. The ligand-based design of compounds 1–12 involved the selective variation of substituents at positions C3 and C6, allowing the pharmacophore identification. The 12 compounds presented here belong to a larger library of synthetic androstane derivatives that were screened for their SERCA2a activity. The need to replace the oxime function with a bioisosteric group guided the design of compounds 1–12.

The single or double carbon–carbon bonds in C3 would ensure a greater metabolic stability in all compounds compared to PST3093. The length of the C3 linker was also varied: a four-atom linker is present in compounds 1–7 exactly reproducing the PST3093 structure, while a shorter two-atom linker is present in compounds 8, 9, and 10, and a longer six-atom linker is present in compounds 11 and 12. All synthetic compounds have a hydroxyl group at C6 as in PST3093; compounds 3, 4, 6, and 9 are in the (R) configuration as PST3093, all the others are in the (S) configuration.

Compounds 1–12 were synthesized through the synthetic strategy depicted in Scheme 1. Compounds 13 and 14, 6R- and 65-hydroxyandrostan-3,17-dione, respectively, were prepared as described from the commercially available prasterone (3β-hydroxyandrost-5-en-17-one).16 The lower reactivity of C17 carbonyl compared to that of C3 allowed for the regioselective reaction of the C3 ketones in Wittig or Horner–Emmons reactions, obtaining the insertion of the carboxylic acid or ester functionalities at position C3.

Compounds 1 and 3 were synthesized by reacting compounds 13 and 14, respectively, with (3-carboxypropyl)-tri phenylphosphonium bromide and sodium hydride in dry dimethylsulfoxide (DMSO), giving the corresponding carboxylic acids (mixture of E + Z isomers 1:2 at C3 alkene) as products of the Wittig reaction that were converted into methyl esters by dissolving in methanol and treating with N-(3-dimethylaminopropyl)-N-ethylcarbodiimide (EDC) and 4-(dimethylamino)pyridine. The E and Z isomers of esters were separated by chromatography, giving compounds 15 and 17, respectively. Ethyl ester hydrolysis with aqueous lithium hydroxide in tetrahydrofuran (THF) afforded compounds 1 and 3. The same Wittig reaction using (3-carboxypentyl)-triphenylphosphonium bromide followed by ethanol esterification gave compound 19 as a mixture of E + Z isomers. Although the use of sodium hydride as a base for the Wittig reaction gave a mixture of E + Z isomers, by reacting 13 and 14 with the same phosphonium salt in the presence of lithium bis(trimethylsilyl)amide (LiHMDS) in THF, it was possible to obtain stereoselectively only the Z olefins, compounds 2 and 4.

The reaction of 13 and 14 with triethylphosphonoacetate in the presence of sodium hydride (Horner–Wadsworth–Emmons reaction) gave the esters 20 and 21 (E + Z mixture).

The catalytic hydrogenation (H2, Pd/C) of compounds 15, 17, 19, 20, and 21 afforded compounds 7, 22, 12, 10, and 23, respectively (Scheme 1). Interestingly, the hydrogenation at the C3 carbon–carbon double bond was found to be completely stereoselective for all derivatives, affording only the β (S) isomers. This is very likely due to the steric hindrance of the C19 methyl group on the upper face of the androstane A ring. Hydrolysis of the esters 7, 22, 12, 10, and 23 by treatment with aqueous lithium hydroxide in THF gave final carboxylic acids 5, 6, 11, 8, and 9, respectively.

**New Synthetic Compounds Do Not Affect Na+/K+ ATPase and Stimulate SERCA2a in a PLN-dependent Way.** It has been recently shown that although istaroxime inhibits the Na+/K+ ATPase activity (IC50 0.14 μM in dog renal preparations), its metabolite PST3093 is inactive against Na+/K+ ATPase. In contrast, both molecules improve disease-induced SERCA2a depression with a similar potency.22 To assess whether the new PST3093 derivatives retain the pharmacological activity of the parent compound, Na+/K+ ATPase and SERCA2a activities were evaluated in vitro assays.

Compounds were tested in a concentration range from 10−9 to 10−4 M on a purified renal Na+/K+ ATPase preparation with a specific activity of 14 μmol/min/mg protein. None of the tested molecules inhibited Na+/K+ ATPase activity up to 10−4 M (Figure 2), similar to PST3093.22

![Figure 2](image-url)

**Figure 2.** Na+/K+ ATPase activity inhibition in a dog purified enzyme preparation. All compounds were tested in a concentration range from 10−9 to 10−4 M in comparison to ouabain. (●) Cpd 9 (~62% at 10−4 M), (○) Cpd 3 (~55% at 10−4 M), and all the other compounds showed <40% inhibition at 10−4 M (N = 2).

SERCA2a ATPase activity was assessed in SR preparations from healthy guinea pig hearts. Ca2+ dependency of ATPase activity was measured, and kinetic parameters (KCa and Vmax) were estimated at compound concentrations of 10 and 100 nM. Four compounds (2, 3, 4, and 9) were inactive, while the other eight compounds significantly increased SERCA2a−Ca2+ affinity (decreased KCa) at nanomolar concentrations (Figure 3). The maximal effect of compounds on KCa reached ~26% at 100 nM, close to that of PST3093 (~25% at 100 nM). No compound affected SERCA2a Vmax activity in healthy guinea pig SR preparations.

Compounds 5–8 and 10 had similar potencies in increasing SERCA2a−Ca2+ affinity (i.e., in stimulating SERCA2a). The activity of compounds 5 and 8 on SERCA2a was further investigated at 1 nM in comparison to PST3093. Also, at this
concentration, compounds 5 and 8 increased SERCA2a−Ca$^{2+}$ affinity ($K_d$Ca reduced by 16%, $N = 7$, $p < 0.05$ and 14%, $N = 12$, $p < 0.05$, respectively) similar to PST3093 (18%, $N = 8$, $p < 0.05$), without affecting $V_{max}$; this makes them good candidates for SERCA2a activation.

Istaroxime$^{24}$ and PST3093$^{22}$ stimulated SERCA2a by relieving SERCA2a−PLN interaction. In a range of concen-
trations from 30 to 1000 nM, compounds 5 and 8 failed to affect skeletal SERCA1 activity in the absence of PLN (Figure 4A). As expected, reconstitution with the PLN_{1−32} fragment markedly reduced SERCA1 affinity for Ca^{2+} (K_{Ca} increased by 23−25%), without affecting V_{max}. Under this condition, compounds 5 and 8 dose-dependently reversed the PLN-
induced shift in $K_dCa$, leaving SERCA1 $V_{\text{max}}$ unchanged, as previously reported for istaroxime 24 and PST3093.22 The present results indicate that in the absence of PLN, SERCA1 is insensitive to compounds 5 and 8; however, sensitivity is restored after reconstitution of SERCA1 with PLN, suggesting that the compounds act by weakening SERCA−PLN interaction, similar to istaroxime24 and PST3093.22

Compounds 5 and 8 were further characterized by testing their effects in cardiac preparations from a diabetic rat model [streptozotocin (STZ)-induced] with impaired SERCA2a function.22,25 Consistent with our previous reports,22,25 baseline SERCA2a $V_{\text{max}}$ activity was 30% lower in STZ (0.239 ± 0.012 μmol/min/mg protein, $N = 9$) than in healthy rats (0.343 ± 0.02 μmol/min/mg protein, $N = 8$, $p < 0.05$); SERCA2a $K_dCa$ was instead unchanged in STZ rats (healthy 259 ± 22 nM, STZ 293 ± 23 nM, NS). Thus, as previously reported,22,25 $V_{\text{max}}$ may represent a better readout of SERCA2a activity in this species. Over the whole range of concentrations tested, compounds 5 and 8 increased SERCA2a $V_{\text{max}}$ in STZ rats (+26% and +25%, +17% and +28%, at 300 and 500 nM, respectively) (Figure 5), thus reversing STZ-induced SERCA2a depression. SERCA2a $K_dCa$ was unchanged by all compounds. Both compounds failed to affect ATPase Ca$^{2+}$ dependency in healthy rats in terms of both parameters $K_dCa$ and $V_{\text{max}}$ (Table S1).

In summary, compounds 5 and 8 displayed similar potency in recovering disease-induced depression of SERCA2a ATPase activity in STZ rats, as previously shown for istaroxime25 and PST3093.22

The absence of effects on Na$^+$/K$^+$ ATPase implies that compounds 5 and 8 may represent new selective SERCA2a activators, similar to their precursor PST3093. Compound 5 was selected for further in vitro and in vivo effects.

**Compound 5 Stimulates SR Ca$^{2+}$ Uptake in Isolated STZ Cardiomyocytes.** A proof-of-principle evidence that compound 5 stimulates SERCA2a was provided by testing its effects on SR Ca$^{2+}$ uptake function in isolated STZ cardiomyocytes through the “SR loading” protocol. This protocol (see Methods, Figure S2 and refs 22 and 25) is suitable to assess SR Ca$^{2+}$ uptake kinetics following caffeine-induced SR depletion under conditions emphasizing the SERCA2a role, that is, in the absence of the Na$^+$/Ca$^{2+}$ exchanger (NCX) function. In particular, through this protocol, we recently showed22 that voltage-induced SR Ca$^{2+}$ reloading is significantly depressed in STZ myocytes, a functional readout of the depressed SERCA2a function in this model.

Cells were incubated for at least 30 min with compound 5 at 1 μM, a concentration not affecting Na$^+$/K$^+$ ATPase. Indeed, as shown in Figure 6A, in isolated rat cardiomyocytes, a detectable inhibition of the Na$^+$/K$^+$ ATPase current ($I_{\text{NaK}}$) was observed at concentrations higher than 20 μM (−19.8 ± 3.1% at 100 μM, $N = 22$), likely for PST3093.22

$$\text{Figure 7. Effect of compound 5 on in vivo echocardiographic parameters in STZ diabetic rats. Compound 5 was i.v. infused (0.2 mg/kg/min) in rats 8 weeks after STZ treatment. Echocardiographic parameters were measured before (basal) and at 15 and 30 min during drug infusion and 10 min after drug interruption under urethane anesthesia. Data are the mean ± SEM; all the results of echocardiographic indexes are summarized in Table S2; N = 13; \ast p < 0.05 (one-way RM ANOVA).}$$
In STZ myocytes, compound 5 (1 μM) sharply accelerated Ca\(^{2+}\) transient (Ca\(_T\)) decay (reducing the Ca\(_T\) decay time constant) and increased excitation release (ER) gain at each pulse of the reloading protocol; Ca\(_T\) amplitude was not significantly affected by the drug (Figure 6B). Comorable results have been obtained with 1 μM PST3093. 22

Overall, compound 5 stimulates SR function in diseased myocytes, most likely through SERCA2a enhancement, within the context of an intact cellular environment.

**In Vivo Administration of Compound 5 is Safe and Improves STZ-Induced Diastolic Dysfunction.** *Toxicity.* Compound 5 was selected to evaluate in vivo effects of the new class. Acute toxicity (LD\(_{50}\)) was preliminarily evaluated in mice following i.v. and oral drug administration. For i.v. administration, compound 5 had an LD\(_{50}\) of 300 mg/kg; for comparison, the LD\(_{50}\) of PST3093 and istaroxime was >250 22 and 23 mg/kg, respectively. Oral toxicity of compound 5 was >800 mg/kg, as compared to >200 and 200 mg/kg for PST3093 and istaroxime, respectively. The main signs of toxicity were prostration, gasping, and convulsions. No overt signs of acute toxicity were observed in the surviving animals.

Collectively, the present data indicate low toxicity of compound 5, particularly as compared to i.v. istaroxime. This is likely due to lack of inhibitory activity on the Na\(^+/K^+\) pump; therefore, the reason for this discrepancy remains to be established.

**In Vivo Hemodynamics.** Features of the STZ-induced diabetic cardiomyopathy have been previously assessed by comparing morphometric, echocardiographic, and cellular parameters between healthy and STZ-treated rats under urethane anesthesia. 22 Echo measurements in STZ rats indicated primarily an impairment of diastolic function, evidenced by decreased early filling velocity (E\(_1\)), increased E wave deceleration time over the E ratio (DT\(_{E}\)/E), and decreased protodiastolic TDI relaxation velocity (\(\epsilon_1\)). The systolic function was moderately altered, as shown by larger LV end-systolic diameter (LVESD), reduced ejection fraction, depressed fractional shortening (FS) and systolic tissue velocity (\(s'\)).

Compound 5 was i.v. infused in STZ rats at a rate of 0.2 mg/kg/min, and the effects on echo parameters were investigated at 15 and 30 min of infusion and 10 min after discontinuation (Figure 7 and Table S2 for all echo parameters). Compound 5 positively affected transmitral Doppler flow indexes by increasing E and A waves; it shortened the DT\(_{E}\), reduced the DT\(_{E}/E\) ratio, and increased both protodiastolic (\(\epsilon_1\)) and telediastolic (\(\epsilon_d\)) TDI relaxation velocities. Compound 5 increased CO, without significantly affecting HR or systolic indexes, such as FS, systolic TDI velocity (\(s'\)), or LVESD. Drug effects reached a plateau at 15 min of infusion; 10 min after discontinuation of the infusion, most echo indexes affected by the compound returned to the basal level.

To summarize, in vivo hemodynamic data indicate a specific lusitropic effect of compound 5 in STZ rats, compatible with rescue of SERCA2a function and largely comparable to the in vivo effect of the parent drug PST3093. 22

**DISCUSSION AND CONCLUSIONS**

In spite of the intense research targeting the discovery of small molecules or gene therapy aimed at selectively activating SERCA2a, recognized to be depressed in HF, no promising clinical outcomes have been reached so far. Therefore, there is still a compelling medical need for a compound with positive lusitropic activity. In this context, we successfully developed a new class of derivatives of PST3093 (the long-lasting metabolite of istaroxime) 22 with a selective stimulatory action on SERCA2a but devoid of inhibitory activity on Na\(^+/K^+\) ATPase. This was achieved by replacing the metabolically unstable oxime of PST3093 with more stable saturated and unsaturated carbon–carbon bonds, without altering the molecular geometry. Biological effects of the new compounds have been investigated by in vitro and in vivo assays.

In the in vitro tests on SERCA2a clearly point out that replacing the C=N group of PST3093 with an alkene C=C, the activity on SERCA2a is partially retained in the case of E isomers (Figure 3). Indeed, although compound 1 (100 nM), carrying the alkene bond in the E configuration, reduced SERCA2a K\(_r\)Ca by 15%, compound 2 with the same bond in the Z configuration was inactive. However, albeit in the E configuration, compound 3 was less active than compound 1, that is, it showed only a trend to SERCA2a K\(_r\)Ca reduction. For comparison, PST3093 (100 nM) reduced SERCA2a K\(_r\)Ca by 25% in the same guinea pig preparation. The fact that compounds with C=C=C in the E configuration are active while Z isomers are inactive parallels the observation that the oxime isomer E is the most active in istaroxime. 16 The replacement of oxime with a saturated C=C bond in C3 in the beta configuration led to compounds 5–12. Compounds 5–8 and 10, in which the C3 linker is two- and four-carbon atom long, have similar activities. Compounds 11 and 12 with a six-carbon linker are slightly less active.

Among the compounds with significant SERCA2a stimulatory activity, compounds 5 and 8, had the best balance between potency and efficiency of chemical synthesis; thus, they were selected as leads for further evaluations. At nanomolar concentrations, the new compounds enhanced in vitro SERCA2a activity in healthy guinea pig preparations and in diseased (STZ) rat preparations; furthermore, the stimulatory effect on SERCA2a depended on the presence of PLN. This pattern is superimposable on that previously reported for the parent compound PST3093 and suggests that the new compounds may also act by partially relieving SERCA2a from PLN-induced inhibition. 22

SR Ca\(^{2+}\) uptake function in diseased (STZ) myocytes was stimulated by compound 5 at a concentration not affecting Na\(^+/K^+\) ATPase. According to the protocol specificity, this effect is surely attributable to SERCA2a stimulation under conditions of depressed function (STZ-induced SERCA2a downregulation). The parent compound PST3093 also in this case shows similar effects. 22

In vivo studies investigated the effects of compound 5 on cardiac function in STZ diabetic rats, a disease model characterized by diastolic dysfunction, as assessed by echocardiography. 22 Compound 5 i.v. infusion at a single dose improved diastolic echo indexes and, because of a small increase in the heart rate, cardiac output; however, it did not affect systolic function significantly (Table S2). Although improvement of diastolic indexes mimics the effect of the parent compound, PST3093 also improved systolic indexes. 22 Similar to compound 5, PST3093 does not inhibit the Na\(^+/K^+\) pump; therefore, the reason for this difference remains to be verified.

Evaluation of in vivo acute toxicity after i.v. administration yielded an LD\(_{50}\) of compound 5 comparable to that of PST3093 whereas the oral one was higher, indicating a lower toxicity. The difference between i.v. and oral toxicity may result

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from the absence of enteric transit/absorption and first-pass hepatic metabolism, which characterize the former administration route. The remarkable reduction of istaroxime toxicity when orally administered can be accounted for by its fast conversion to PST3093, missing Na⁺/K⁺ pump inhibition.

Overall, the new PST3093 derivatives provide a tool for pharmacological enhancement of SERCA2a function, leading to improvement of in vivo diastolic function. With respect to PST3093, its derivatives are devoid of the oxime function and thus suitable for chronic usage and have a lower acute oral toxicity. They further differ from istaroxime, the progenitor of "PLN antagonists" already tested for clinical use because of lack of Na⁺/K⁺ pump inhibition. Considering the proarrhythmic potential of the latter, this may represent a substantial advantage in terms of safety; nonetheless, at variance with istaroxime, the new compounds should be seen as purely "lusitropic" agents, that is, devoid of "inotropic" effects beyond that expected from systo–diastolic coupling.

Although the mechanism of action of PST3093 and synthetic analogues has still to be investigated, the reversal of electrostatic properties from cationic ammonium for istaroxime to anionic carboxylate for PST3093 and analogues could account for the inactivity toward Na⁺/K⁺ ATPase.

Because PST3093 has a half-life (~10 h) much longer than that of istaroxime (~1 h) and its derivatives have been obtained by introducing groups with higher chemical stability, pharmacokinetics of the new compounds is likely compatible with chronic oral dosing. Nonetheless, suitability of the new compounds for chronic therapy, their natural destination, remains to be directly tested.

**EXPERIMENTAL SECTION**

Methods are succinctly described here; details are given in the Supporting Information.

**Animal Models.** All experiments involving animals confirmed to the guidelines for Animal Care endorsed by the Milano-Bicocca and Chang Gung Universities and to the Directive 2010/63/EU of the European Parliament on the protection of animals used for scientific purposes.

**Chemistry. General.** All reagents and solvents were purchased from commercial sources and used without further purification. Reactions were carried out under an argon atmosphere unless otherwise noted and were monitored by thin-layer chromatography. Reactions were carried out under an argon atmosphere unless otherwise noted and were monitored by thin-layer chromatography. Reactions were carried out under an argon atmosphere unless otherwise noted and were monitored by thin-layer chromatography.

**Biochemical Measurements.** Total ATPase activity was assessed by measuring the rate of 32P-ATP release (μmol/min/mg protein) at 37 °C.

**Na⁺/K⁺ ATPase Activity Assay.** The inhibitory effect of compounds was tested at multiple concentrations on Na⁺/K⁺ ATPase-alpha1 isofrom purified from dog kidneys. Na⁺/K⁺ ATPase activity was identified as the ouabain (1 mM)-sensitive component of total one; compound efficacy was expressed as the concentration exerting 50% inhibition (IC₅₀).

**SERCA ATPase Activity Assay.** Measurements were performed in cardiac SR microsomes (guinea pig) and in whole tissue homogenates (rat). Cardiac preparations included SERCA2a and PLN. To test for PLN involvement in the effect of compounds, SERCA1 activity was measured in PLN-free microsomes (from guinea pig skeletal muscle) before and after reconstitution with the synthetic PLN₃−32 inhibitory fragment at a ratio of 300:1 for PLN/SERCA. The SERCA component, identified as the cyclopiazonic acid (10 μM)-sensitive one, was measured at multiple Ca²⁺ concentrations (100–1500 nM), and Ca²⁺ dose-response curves were fitted to estimate SERCA maximal hydrolytic velocity (Vₘₐₓ, μmol/min/mg protein) and the Ca²⁺ dissociation constant (Kₐₕ, CaₙM). Either an increase in Vₘₐₓ (rat) or a decrease in Kₐₕ (guinea pig) (increased Ca²⁺ affinity) (guinea pig) stands for enhancement of SERCA function.

**Functional Measurements in Isolated Cardiomyocytes.** LV myocytes were isolated from healthy and STZ rats as previously described with minor modifications.

Na⁺/K⁺ ATPase Current (IₖNaK). IₖNaK was recorded in isolated myocytes from healthy rats as the holding current at −40 mV under conditions enhancing IₖNaK and minimizing contamination by other conductances. IₖNaK inhibition by compound 5 was expressed as percentage reduction of ouabain (1 mM)-sensitive current.

**Intracellular Ca²⁺ Dynamics.** Ca²⁺-dependent fluorescence (Fluo4-AM) was recorded in patch-clamped LV myocytes from STZ rats and quantified by normalized units (F/F₀). The SR Ca²⁺ uptake rate was evaluated through a "SR loading" voltage protocol, specifically devised to examine the system at multiple levels of SR Ca²⁺ loading and to rule out NCX contribution. Current through the L-type Ca²⁺ channel (IₕCaₗ) was simultaneously recorded, and the ER "gain" was calculated as the ratio between Ca₉ amplitude and Ca²⁺ influx through IₕCaₗ up to the Ca₉ peak (protocol in Figure S2).

**In Vivo Studies. Acute Drug Toxicity Studies in Mice.** Acute toxicity of compound 5 was preliminarily evaluated in male Albino Swiss CD-1 mice for determining the dose causing 50% mortality (LD₅₀, mg/kg body weight) at 24 h after i.v. injection or oral treatment. Compound 5 was dissolved in saline solution and i.v. injected at 100, 200, 250, 275, and 300 mg/kg (one–four animals for each group) or orally administered at 100, 200, 400, 600, and 800 mg/kg (four animals for each group). Control animals received the vehicle only.

Acute toxicity following oral treatment with istaroxime and PST3093 was also evaluated for comparison; istaroxime was administered at 30, 100, and 300 mg/kg (four animals for each group) and PST3093 at 200 mg/kg (four animals for each group).

**Humodynamic Studies in Rats with Diabetic Cardiomyopathy.** Diabetes was selected as the in vivo model because of its association with reduced SERCA2a function. Diabetes was induced in Sprague Dawley male rats (150–175 g) by a single i.v. STZ group (50 mg/kg in citrate buffer) injection in the tail vein. Control (healthy group) rats received vehicle (citrate buffer). Fasting glycaemia was measured after 1 week, and rats with values >290 mg/dL were considered diabetic.

Rats were studied by transthoracic echocardiographic under anesthesia (1.25 g/kg urethane, i.p.). Left-ventricular end-diastolic (LVEDD) and end-systolic (LVESD) diameter, posterior wall (PW), and interventricular septal thickness (IVST) were measured according to the American Society of Echocardiography guidelines. FS was calculated as FS = (LVEDD − LVESD)/LVEDD. Transmirtal flow velocity was measured (pulsed Doppler) to obtain early and late filling velocities (E and A waves) and E wave deceleration time (DT). DT was also normalized to E wave amplitude (DT/E ratio). Peak myocardial systolic (s') and diastolic velocities (e' and a') were measured at the mitral annulus by tissue Doppler imaging (TDI).

Compounds 5 was i.v. infused at 0.2 mg/kg/min (0.16 mL/min); echocardiographic parameters were measured before and at 15 and 30 min during the infusion and 10 min after drug discontinuation.
Statistical Analysis. Individual means were compared by unpaired or paired t-test; multiple means were compared by one-way ANOVA for repeated measurements (RM) followed by post hoc Tukey's multiple comparisons. Data are reported as mean ± SEM; p ≤ 0.05 defined statistical significance of differences in all comparisons. The number of animals or cells are specified in each figure legend.

- **ASSOCIATED CONTENT**
  
  Supporting Information
  The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jmedchem.2c00347.
  Synthetic procedures, characterization of all compounds and intermediates (assignments of 1H and 13C NMR spectra, HRMS, and HPLC spectra), supplementary methods (animal models, biochemical measurements, functional measurements in isolated myocytes, and in vivo studies), supplementary tables (PDF).
  Molecular formula strings (CSV).

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  **Author Contributions**
  F.P., M.R., and A.Z. contributed equally as senior authors to the article. A.L. and A.R. conducted chemical assays; M.F. and P.B. conducted biochemical assays; M.A. and E.T. conducted functional measurements in isolated myocytes; S.-C.H. carried out in vivo measurements; C.R. carried out formal analysis; G.-J.C. supervised in vivo measurements; P.F. and G.B. supervised biochemical measurements; A.Z., M.R., and F.P. revised and supervised the study and wrote the article.

  **Notes**
  The authors declare the following competing financial interest(s): MF and PB are Windtree employees, PF and GB are Windtree consultants, S-CH is an employee of CVie Therapeutics Limited. All the other Authors declare no conflict of interest.

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  **ABBREVIATIONS**
  APT, attached proton test; 2D-COSY, two-dimension homonuclear correlation spectroscopy; CO, cardiac output; CPA, cyclopiazonic acid; Cpd, compound; 2D-HSQC, two-dimension heteronuclear single quantum correlation; 2D-NOESY, two-dimension nuclear Overhauser effect spectroscopy; DMF, dimethylformamide; DMSO, dimethylsulfoxide; DT, E wave deceleration time; E, early filling velocity; e’, TDI relaxation velocity; EDC, N-ethyl-N’-(3-dimethylaminopropyl) carbodiimide; EF, ejection fraction; EtOAc, ethyl acetate; EtOH, ethanol; FS, fractional shortening; HPLC, high-performance liquid chromatography; HF, heart failure; Iv, intraventricular; i.e., intravenous; HR, heart rate; IVST, interventricular septal thickness; LD50, lethal dose 50%; LVEDD, left-ventricular end-diastolic diameter; LVESD, left-ventricular end-systolic diameter; LiHMDS, lithium bis(trimethylsilyl)amide; PLN, phospholamban; PWT, posterior wall thickness; QSAR, quantitative structure–activity relationship; s’, systolic tissue velocity; SERCA, sarcoplasmic reticulum calcium ATPase; SR, sarcoplasmic reticulum; STZ, streptozotocin; SV, stroke volume; TDI, tissue Doppler imaging; THF, tetrahydrofuran; TLC, thin-layer chromatography

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