Serelaxin in addition to standard therapy in acute heart failure

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Serelaxin in addition to standard therapy in acute heart failure: rationale and design of the RELAX-AHF-2 study

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Patients admitted for acute heart failure (AHF) experience high rates of in-hospital and post-discharge morbidity and mortality despite current therapies. Serelaxin is recombinant human relaxin-2, a hormone with vasodilatory and end-organ protective effects believed to play a central role in the cardiovascular and renal adaptations of human pregnancy. In the phase 3 RELAX-AHF trial, serelaxin met its primary endpoint of improving dyspnoea through day 5 in patients admitted for AHF Compared to placebo, serelaxin also reduced worsening heart failure (WHF) by 47% through day 5 and both all-cause and cardiovascular mortality by 37% through day 180. RELAX-AHF-2 (ClinicalTrials.gov NCT01870778) is designed to confirm serelaxin’s effect on these clinical outcomes. RELAX-AHF-2 is a multicentre, randomized, double-blind, placebo-controlled, event-driven, phase 3 trial enrolling ~6800 patients hospitalized for AHF with dyspnoea, congestion on chest radiograph, increased natriuretic peptide levels, milder-to-moderate renal insufficiency, and systolic blood pressure ≥125 mmHg. Patients are randomized within 16 h of presentation to 48 h intravenous infusions of serelaxin (30 μg/kg/day) or placebo, both in addition to standard of care treatments. The primary objectives are to demonstrate that serelaxin is superior to placebo in reducing: (i) 180 day cardiovascular death, and (ii) occurrence of WHF through day 5. Key secondary endpoints include 180 day all-cause mortality, composite of 180 day combined cardiovascular mortality or heart failure/renal failure rehospitalization, and in-hospital length of stay during index AHF. The results from RELAX-AHF-2 will provide data on the potential beneficial effect of serelaxin on cardiovascular mortality and WHF in selected patients with AHF.

Keywords
Acute heart failure • Serelaxin • Worsening heart failure • Mortality • Phase 3 trial

Introduction

Acute heart failure (AHF) is the most common cause of hospitalization in patients 65 years and older.1,2 In part due to the ageing of the population and more effective treatment of chronic heart failure (HF), its prevalence is expected to increase by 25% over the next 20 years3 and the problem has expanded worldwide.4,5 Patients hospitalized for HF have a 40–50% rate of HF exacerbation, of which 10–15% is in-hospital worsening heart failure (WHF)6–10 and 30–40% is rehospitalization, within the first 6 months after discharge as well as a 10–15% mortality rate.11 Compared with ambulatory patients with stable chronic HF, patients hospitalized for AHF have a dramatic increase in their risk of death, similar or worse than that after a hospitalization for
Rationale and design of the RELAX-AHF-2 trial

FLUID OVERLOAD
FLUID REDISTRIBUTION TO THE LUNGS

ACUTE HEART FAILURE
SYMPTOMS AND SIGNS
IN-HOSPITAL WHF

MICROVASCULAR DYSFUNCTION
HYPOPERFUSION/ISCHEMIA
REACTIVE OXYGEN SPECIES

NEUROHORMONAL/
INFLAMMATORY ACTIVATION

MYOCARDIAL, RENAL
HEPATIC DAMAGE

INCREASED MORTALITY

Figure 1 Mechanisms of increased mortality and worsening heart failure (WHF) in acute heart failure.

The efficacy and safety of serelaxin as a continuous intravenous (i.v.) infusion for up to 48 h in AHF patients have been evaluated in two multicentre, randomized, double-blind, placebo-controlled trials: (i) the dose-finding phase 2 study Pre-RELAX-AHF and (ii) the phase 2 registration study RELAX-AHF. In both of these trials, patients were admitted for acute heart failure with persistent dyspnoea despite i.v. diuretics with normal-to-elevated systolic blood pressure (SBP >125 mmHg), congestion on chest radiograph, elevated natriuretic peptides, and mild-to-moderate renal insufficiency, and enrolled within 16 h of presentation. The 234 patients enrolled in Pre-RELAX-AHF were randomized to four doses of serelaxin ranging from 10 to 250 μg/kg/day or matching placebo. While each dose suggested some clinical benefit, patients assigned to serelaxin 30 μg/kg/day had the greatest overall improvement in signs and symptoms of HF and trends toward improved long-term outcomes with minimal adverse effects. In the RELAX-AHF study, 1161 patients admitted for AHF were randomized to receive either serelaxin (n = 581) or matching placebo (n = 580), both in addition to standard-of-care AHF treatment. The 48 h i.v. infusion of serelaxin at the dose of 30 μg/kg/day produced dyspnoea relief as demonstrated by a 19.4% treatment improvement compared to placebo measured over 5 days by visual analogue scale, representing one of the two primary efficacy endpoints in the study. However, there was no significant effect on the other primary endpoint of dyspnoea relief through 24 h measured by a Likert scale. Serelaxin...
treatment significantly reduced the incidence of WHF through day 5 which was the main component of the improvement in dyspnoea and contributed to the reduced length of index hospital stay. These clinical effects were associated with significant improvements in biomarkers suggestive of less end-organ damage and dysfunction in serelaxin-treated patients.\textsuperscript{20} Analysis of all-cause mortality through day 180 revealed fewer deaths with serelaxin compared to placebo with a total of 42 [Kaplan–Meier (K–M) estimate, 7.3%] deaths in the serelaxin group compared to 65 (K–M estimate, 11.3%) in the placebo group [hazard ratio (HR) 0.63; 95% confidence intervals (CI) 0.43–0.93; \( P = 0.020 \)]. The mortality difference was largely driven by cardiovascular (CV) death through day 180 (K–M estimates 9.6% and 6.1% in placebo and serelaxin groups, respectively, HR 0.63; 95% CI 0.41–0.96; \( P = 0.028 \)). Including all patients treated with serelaxin 30 \( \mu \)g/kg/day from the two trials, CV mortality was reduced by 44% (Figure 3; HR 0.56; 95% CI 0.37–0.86; \( P = 0.007 \)). These findings support a sustained benefit of serelaxin beyond the initial 48 h of administration. Better relief of congestion and protection from damage to the myocardium, kidneys, and liver seem the most likely mechanisms for these long-term beneficial effects.\textsuperscript{19,20} These results were also consistent with a reduction in WHF episodes in serelaxin-treated patients. Worsening HF is associated with poorer outcomes independently of AHF severity.\textsuperscript{20}

Given that mortality was not a primary efficacy endpoint of the RELAX-AHF trial, a global, phase 3 trial designed to evaluate the efficacy, safety and tolerability of i.v. infusion of 30 \( \mu \)g/kg/day serelaxin for 48 h, when added to standard therapy. The trial is event-driven: \( \sim 6800 \) AHF patients in 545 centres in 34 countries will be enrolled to obtain at least 547 confirmed CV deaths. Primary efficacy will be determined based on the relative risk reductions in CV death through day 180 and in WHF through day 5. Secondary efficacy endpoints include 180 day all-cause mortality, length of hospital stay, and CV death or HF/renal failure rehospitalization through day 180.

The RELAX-AHF-2 Executive Committee (Appendix) designed the trial and wrote the study protocol in collaboration with the clinical team from Novartis. The protocol must be approved by the Ethics Review Committee/Institutional Review Board affiliated with each centre and is being conducted in accordance with Good Clinical Practice and the 2002 Declaration of Helsinki. All participants provide written informed consent. The trial is registered on ClinicalTrials.gov, NCT01870778). The protocol was amended five times (see Supplementary material online, Appendix S1).

**Study design and methods**

The RELAX-AHF-2 trial (Figure 4) is a multicentre, randomized, double-blind, placebo-controlled trial to evaluate the efficacy, safety and tolerability of i.v. infusion of 30 \( \mu \)g/kg/day serelaxin for 48 h, when added to standard therapy. The trial is event-driven: \( \sim 6800 \) AHF patients in 545 centres in 34 countries will be enrolled to obtain at least 547 confirmed CV deaths. Primary efficacy will be determined based on the relative risk reductions in CV death through day 180 and in WHF through day 5. Secondary efficacy endpoints include 180 day all-cause mortality, length of hospital stay, and CV death or HF/renal failure rehospitalization through day 180.

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**Study population**

The study population includes male and female patients (\( \geq 18 \) years old) admitted to the hospital for AHF with dyspnoea, congestion on chest radiograph, elevated BNP or NT-proBNP, normal-to-elevated SBP \( \geq 125 \) mmHg, and mild-to-moderate renal impairment [estimated glomerular filtration rate \( \geq 25 \) and \( \leq 75 \) mL/min/1.73 m\(^2\), calculated using the standardized Modification of Diet in Renal Disease (sMDRD) equation] who are anticipated to require i.v. therapy for at least 48 h
Rationale and design of the RELAX-AHF-2 trial

Figure 3 Cardiovascular mortality of patients with acute heart failure in the serelaxin programme treated with serelaxin 30 μg/kg/day compared to placebo.

Figure 4 Schematic diagram of study design for RELAX-AHF-2 trial. HF, heart failure; IV, intravenous.

(Table 1). Patients must remain symptomatic after initial treatment with at least 40 mg i.v. loop diuretic. Patients receiving i.v. nitrates at a dose <0.1 mg/kg/h with a blood pressure >150 mmHg are eligible. There is no ejection fraction criterion, such that both HF patients with preserved (HFrEF) or reduced ejection fraction (HFrEF) are enrolled. Patients are randomized within 16 h of the earlier of first administration of i.v. loop diuretic or hospital presentation for the AHF episode in a 1:1 ratio into the two study arms.

Study treatment

Serelaxin or matching placebo is administered as an i.v. infusion beginning no more than 4 h after randomization. Infusion continues for up to 48 h according to a weight range-adjusted dosing regimen at the nominal dose of 30 μg/kg/day. Similar to the protocol adopted in the previous Pre-RELAX-AHF and RELAX-AHF trials, blood pressure is monitored frequently during study drug administration.

If at any time during the study drug administration the patient’s SBP decreases by >40 mmHg from baseline but the absolute SBP is ≥100 mmHg in two consecutive measurements 15 min apart, the study drug infusion rate is decreased by 50% for the remainder of the infusion period. If the patient’s SBP falls to <100 mmHg in two consecutive measurements 15 min apart, the study drug is permanently discontinued. Randomized patients are required to receive standard-of-care background HF management during both the index hospitalization and the follow-up period of 180 days. After randomization, the investigator may prescribe any additional medications dictated by the patients’ condition, including i.v. loop diuretics and vasoactive medications.

Study assessments

Patients are assessed daily while hospitalized through day 5 or discharge, whichever comes first. They are also assessed at days 14, 60, 120 (phone contact), and 180 (Table 2). Heart failure signs and symptoms...
### Table 1 Key inclusion and exclusion criteria in RELAX-AHF-2

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<tr>
<th>Key inclusion criteria</th>
<th>Key exclusion criteria</th>
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<tr>
<td>1. Male or female ≥18 years of age who sign the informed consent, with body weight ≤160 kg</td>
<td>1. Dyspnoea due to non-cardiac causes such as acute or chronic respiratory disorders or infections (i.e., severe COPD, bronchitis, pneumonia), which may interfere with the ability to interpret the primary cause of dyspnoea</td>
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<td>2. Hospitalized for AHF with the anticipated requirement of i.v. therapy (including i.v. diuretics) for at least 48 h; AHF is defined as including all of the following measured at any time between presentation (including the emergency department) and the end of screening:</td>
<td>2. Known history of respiratory disorders requiring the daily use of i.v. or oral steroids; need for intubation or the current use of i.v. or oral steroids for COPD</td>
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<td>- Persistent dyspnoea at rest or with minimal exertion at screening and at the time of randomization, despite standard background therapy for AHF including the protocol required i.v. furosemide of at least 40 mg total (or equivalent)</td>
<td>3. Patients with blood pressure &gt;180 mmHg at the time of randomization or persistent heart rate &gt;130 b.p.m.</td>
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<td>- Pulmonary congestion on chest radiograph</td>
<td>4. Temperature &gt;38.5°C (oral or equivalent) or sepsis or active infection requiring i.v. anti-microbial treatment</td>
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<td>- BNP ≥500 pg/mL or NT-proBNP ≥2000 pg/mL for patients ≥75 years of age or with current atrial fibrillation (at the time of randomization), BNP ≥750 pg/mL or NT-proBNP ≥3000 pg/mL</td>
<td>5. Clinical evidence of acute coronary syndrome currently or within 30 days prior to enrolment. (Note that the diagnosis of acute coronary syndrome is a clinical diagnosis and that the sole presence of elevated troponin concentrations is not sufficient for a diagnosis of acute coronary syndrome, given that troponin concentrations may be significantly increased in the setting of AHF)</td>
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<td>3. Systolic blood pressure ≥125 mmHg at the start and at the end of screening and impaired renal function defined as an eGFR&lt;sup&gt;b&lt;/sup&gt; between presentation and randomization of ≥25 and ≤75 mL/min/1.73 m&lt;sup&gt;2&lt;/sup&gt;, calculated using the sMDRD equation</td>
<td>6. AHF due to significant arrhythmias, which include any of the following: sustained ventricular tachycardia, bradycardia with sustained ventricular rate ≤45 b.p.m., or atrial fibrillation/flutter with sustained ventricular response of &gt;130 b.p.m.</td>
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<td>4. Able to be randomized within 16 h of presentation to the hospital, including the emergency department&lt;sup&gt;a&lt;/sup&gt;</td>
<td>7. Patients with severe renal impairment defined as pre-randomization eGFR &lt;25 mL/min/1.73 m&lt;sup&gt;2&lt;/sup&gt; calculated using the sMDRD equation, and/or those receiving current or planned dialysis or ultrafiltration.</td>
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<tr>
<td>5. Received i.v. furosemide of at least 40 mg total (or equivalent) at any time between presentation (this includes outpatient clinic, ambulance, or hospital including emergency department) and the start of screening for the study for the treatment of the current AHF episode. Time from presentation to start of furosemide administration should be less than 6 h</td>
<td>8. Patients with haematocrit &lt;25% or a history of blood transfusion within the 14 days prior to screening, or active life-threatening GI bleeding.</td>
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<td>9. Known hepatic impairment (as evidenced by total bilirubin &gt;3 mg/dL, or increased ammonia levels, if performed) or history of cirrhosis with evidence of portal hypertension such as varices</td>
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<td>10. Significant, uncorrected, left ventricular outflow obstruction, such as obstructive hypertrophic cardiomyopathy or severe aortic stenosis (i.e. aortic valve area &lt;1.0 cm&lt;sup&gt;2&lt;/sup&gt; or mean gradient &gt;50 mmHg, on prior or current echocardiogram), and severe mitral stenosis</td>
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<td>11. Severe aortic insufficiency or severe mitral regurgitation for which surgical or percutaneous intervention is indicated</td>
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<td>12. Documented, prior to or at the time of randomization, restrictive amyloid myocardiopathy, or acute myocarditis or hypertrophic obstructive, restrictive, or constrictive cardiomyopathy (does not include restrictive mitral filling patterns seen on Doppler echocardiographic assessments of diastolic function)</td>
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<td>13. Current (within 2 h prior to randomization) or planned (through the completion of study drug infusion) treatment with any i.v. vasoactive therapies, including vasodilators (including nesiritide), positive inotropic agents and vasopressors, or mechanical support (endotracheal intubation, mechanical ventilation; intra-aortic balloon pump or any ventricular assist device; haemofiltration, ultrafiltration or dialysis), with the exception of i.v. furosemide (or equivalent), or i.v. nitrates at a dose of ≥0.1 mg/kg/h if the patient has a systolic blood pressure &gt;150 mmHg at the start of screening</td>
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<td>14. Any major solid organ transplant recipient or planned/anticipated organ transplant within 1 year or major surgery, including implantable devices (e.g., ICD, CRT), or major neurological event including cerebrovascular events, within 30 days prior to screening</td>
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<td>15. History of malignancy of any organ system other than localized basal cell carcinoma of the skin, treated or untreated, within the past year with a life expectancy less than 1 year</td>
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<td>16. Women of child-bearing potential, defined as all women physiologically capable of becoming pregnant, unless they are using highly effective methods of contraception during dosing of study treatment plus 5 days after cessation of study drug</td>
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</table>

<sup>a</sup>Assessed based on local laboratory

<sup>b</sup>Presentation starts as the earliest of (i) time of presentation at either the emergency room/department, intensive/cardiac care unit or ward (excludes emergency medical service or other pre-hospital care); or (ii) time of first i.v. loop diuretic prior to arrival at the hospital (this includes outpatient clinic, ambulance, or hospital including emergency department) for the current AHF episode. Time from presentation to start of furosemide administration should be less than 6 h

AHF, acute heart failure; COPD, chronic obstructive pulmonary disease; CRT, cardiac resynchronization therapy; eGFR, estimated glomerular filtration rate; GI, gastrointestinal; HF, heart failure; ICD, implantable cardioverter defibrillator; i.v., intravenous; sMDRD, standardized Modification of Diet in Renal Disease.
symptoms are assessed through day 60, and local haematology and clinical chemistry tests performed through day 5. Adverse events are reported from signing of the informed consent form through day 5 for non-serious adverse events and through day 14 for serious adverse events. All deaths and hospitalizations reported through day 180 are adjudicated by a Clinical Events Committee (Appendix). Rehospitalization is defined as an unplanned hospitalization (including admission to a hospital or any attendance in an acute care setting, e.g. emergency department, or in another health care facility) of 24 h or greater, regardless of whether the patient was admitted to the hospital.

The occurrence of WHF is reported by the investigator through day 5 post-randomization, and is defined as worsening signs and/or symptoms of HF that require an intensification of i.v. therapy or mechanical ventilatory, renal or circulatory support for HF. Such treatment can include the institution or up-titration of i.v. diuretic, i.v. nitrates, or any other i.v. medication for HF, or institution of mechanical support such as ventilation, ultrafiltration, haemodialysis, intra-aortic balloon pump, or ventricular assist device. Worsening HF can occur within the 5 day post-randomization period either during the index admission, or after discharge as an unplanned rehospitalization or unscheduled physician office/emergency department visit due to a primary diagnosis of HF. This endpoint also includes patients who die in this 5 day period from any cause.

ECGs are performed and interpreted locally for all patients at screening and at day 5 or discharge, whichever occurs first. In addition, ECGs are performed locally and interpreted by a central core lab at baseline and end of study infusion in a substudy of at least 500 patients in selected centres to evaluate the impact of seleralxin on ECG variables. Samples through day 14 are collected for central haematology, blood chemistry, and biomarker assays in ~1600 patients at selected centres.

Statistical considerations

Primary and secondary efficacy outcomes, except all-cause mortality, will be compared between treatment groups on an intent-to-treat basis using a sequentially rejective multiple testing procedure controlling the overall two-sided α at 5%. Treatment groups will be compared regarding the time to CV death through day 180 with a log-rank test at an initial significance level of (4/5)α, and regarding time to WHF through day 5 at an initial significance level of (1/5)α using a Gehan’s generalized Wilcoxon test. All-cause mortality will be tested independently at the two-sided 5% significance level, if the test of either or both of the primary endpoints is significant. The significance level for the final test will be adjusted to account for the interim efficacy analysis, planned to occur after ~60% (i.e. 329) of confirmed CV deaths have accrued. A Lan–DeMets spending function approximating an O’Brien–Fleming stopping boundary will be employed to control the overall one-sided statistical testing of the CV mortality endpoint at the 2% level and the interim analysis will only be performed on the CV mortality endpoint. An independent Data Monitoring Committee (Appendix), supported by an independent, unblinded statistical centre, regularly reviews safety data as well as the interim efficacy analysis.

Accounting for the one interim efficacy analysis, 547 confirmed CV deaths are needed for 80% power to detect a 22% relative risk reduction. Assuming the 180 day CV death rate is 9.0% in the placebo group, which is ~80% of 11.3% of all-cause death observed in the placebo group in RELAX-AHF, ~6800 patients will need to be enrolled. The observed overall rate of the primary CV mortality endpoint will be assessed on a blinded basis, and adjustments to patient enrolment made in order to achieve the required number of events. Using the proposed multiple testing procedure, the power for WHF is at least 80% with the sample size of 6800 assuming at least a 20% relative risk reduction with 12.2% placebo event rate based on RELAX-AHF data.

Discussion

Over the last two decades, morbidity and mortality from chronic HFrEF has decreased dramatically with the adoption of ACE inhibitors/angiotensin receptor blockers, beta-blockers, and mineralocorticoid receptor antagonists. These therapies comprise the cornerstone of pharmacological chronic HF treatment. More recently, further progress has been made with the addition of an angiotensin receptor/neprilysin inhibitor (ARNI) as an alternative to ACE inhibitors, and ivabradine as an adjunct to maximally tolerated beta-blocker therapy in patients in sinus rhythm.1,2 However, these patients remain at high risk for acute decompensation, an event associated with a marked increase in mortality and HF recurrence either in the form of in-hospital or post-discharge WHF. Despite the urgent need to improve the outcomes of patients with AHF, only three therapies have gained regulatory approval in the last two decades: i.v. milrinone and nesiritide in the US and levosimendan in Europe.1,2 Despite their approval, none of these drugs has demonstrated favourable effects on outcomes. Current treatment of AHF is based on drugs that have limited evidence of efficacy based on formal randomized controlled data. While diuretics have a Class I recommendation with a C level of evidence for the treatment of patients with congestion for symptom relief, other therapies such as nitrates have Class IIa recommendations with a B level of evidence.1 None of these therapies has an indication for improvement in clinical outcomes.

On the other hand, trials of novel therapies for AHF have failed to significantly improve symptoms or outcomes.24 Given the number of failures, variables other than the drug itself may have influenced these results. The heterogeneity of AHF patients, the inclusion of patients with non-cardiac causes of symptoms, and the failure to align the drug’s mechanism of action with the optimal patient population most likely to benefit from the study treatment are also potentially contributing causes.

To address these limitations, the RELAX-AHF trials have enrolled patients admitted for AHF with persistent symptoms well defined by objective clinical diagnostics, including congestion on chest radiograph and elevated natriuretic peptide plasma levels. The patients are required to remain symptomatic despite initial treatment with i.v. diuretics, and in RELAX-AHF-2 only patients with severe enough AHF anticipated to require 48 h of i.v. therapy are enrolled. The inclusion of at least mild renal dysfunction among the entry criteria allows selection of a higher-risk patient population most likely to benefit from the study treatment are also potentially contributing causes.

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### Table 2 Assessment schedule

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BP, blood pressure; ECG, electrocardiogram; HF, heart failure; HR, heart rate; PSW, premature patient withdrawal; X, assessment.

a ECGs will be performed and interpreted locally at screening and at day 5 or discharge, whichever occurs first.
b ECGs will be collected in a subset of randomized patients participating in the ECG substudy and sent to a central ECG vendor for evaluation. ECGs will be collected at baseline and at the end of study drug infusion.
c The echocardiogram should be performed as soon as possible post-randomization, but prior to discharge. If an echocardiogram is performed during the screening period (i.e., within the 16 h window) and the patient is subsequently randomized, the screening echocardiogram will qualify as the index hospitalization echocardiogram and a repeat echocardiogram post-randomization will not be necessary.
d A complete physical examination will be performed at screening; an abbreviated physical examination will be performed at all other specified time points.
e BP and HR measurements are to be performed at 30 and 60 min and then every hour for the first 6 h of study drug infusion, and then every 3 h during study drug infusion, including night-time hours. Post-infusion, BP and HR are to be measured every 3 h until 12 h following end of infusion, then every 6 h for 48 h and then every 24 h until the earlier of day 5 or discharge. BP and HR are to be measured with the patient in the same position and with the same equipment using the same arm, throughout study drug infusion. These measurements may be made and recorded by trained health care personnel as part of their routine clinical duties, as well as study personnel.

f At hours 24, 48, 72, 96, and 120, physician assessment of HF signs and symptoms will include an assessment of the occurrence of worsening HF in the interval preceding the visit.
g Blood will be locally collected and analyzed daily during hospitalization. If discharge occurs prior to day 5, local blood collection will not be required at the day 5 hospital/domicile visit.
h Blood will be collected in a subset of randomized patients participating in the biomarker substudy for measurement of biochemistry, haematology and plasma glycated haemoglobin by the central laboratory. Urine dipstick will be measured locally at screening to rule out any conditions requiring further diagnostic evaluation or treatment.
i Major cardiovascular and non-cardiovascular classes of medication taken by a subject from -30 days prior to study drug initiation and on a daily basis while hospitalized through day 5, at discharge, and at days 14, 60, 120, and 180 will be recorded. Only those medications currently being taken or that were taken within 24 h prior to the visit will be collected.
j Non-serious and serious adverse events will be reported from the signing of the informed consent form through days 5 and 14, respectively.
and clinical practice. Enrolling patients early enough to maximize the potential end-organ protective and haemodynamic benefits of serelaxin, yet late enough to be confident in the diagnosis of AHF, is central to the design of the RELAX-AHF trials. Other studies that focused on dyspnoea relief suggested that earlier initiation of therapy provides greater symptom relief. However, in one study, only two-thirds of the enrolled patients with a suspected diagnosis of AHF within 1 h of presentation were confirmed to have AHF at 6 h after presentation, suggesting that too rapid enrolment might result in patients without AHF being included in the trial. In both Pre-RELAX-AHF and RELAX-AHF, patients were randomized within 16 h of presentation at a median of 8–9 h; this time frame is the goal for RELAX-AHF-2 as well. In contrast with studies in patients with chronic HF where a reduced ejection fraction has been a useful criterion for patient selection, the serelaxin trials enrolled patients with both HFrEF and HfPEF. This is consistent with previous trials testing drugs acting on peripheral vessels and/or renal function in patients with AHF. The prognosis of the patients with AHF was not found to be related to left ventricular ejection fraction and a similar benefit of serelaxin on clinical outcomes was observed in both patients with preserved and reduced ejection fraction. However, to provide additional information on the potential beneficial effects of serelaxin in these groups of patients, echocardiograms are obtained during the index hospitalization in RELAX-AHF. The timing of the echocardiograms was not specified, since a prior study suggested that left ventricular function does not change significantly in these types of AHF patients and it would be undesirable to have the echocardiogram interfere with randomization and initiation of study drug.

There is no limitation on concomitant oral therapies during study drug administration, but i.v. vasoactive therapies are either excluded or limited. Given the entry criterion of SBP ≥125 mmHg, there is no indication for vasopressors or inotropic agents, and while relatively safe, nesiritide has demonstrated no clinical benefits and has limited regulatory approval globally. Intravenous nitrates are the only i.v. vasoactive drugs allowed in addition to diuretics at the time of enrolment. Despite lack of evidence about their efficacy in the treatment of AHF and a recent guideline specifically stating, ‘Do not routinely offer nitrates to people with AHF,’ many authorities continue to support their use. This may be especially true in patients with increased blood pressure at the time of randomization and hence, administration of i.v. nitrates was allowed only in this subgroup of patients (i.e. SBP >150 mmHg at the time of screening). The dose of nitrates allowed in RELAX-AHF-2 is clinically relevant and not particularly restrictive; an 80 kg person could be receiving up to 133 μg/min nitroglycerin or three times the dose achieved in VMAC, and over 6–10 times the recommended starting dose of the ESC guidelines.

While few would challenge the clinical importance of reducing CV mortality, some have questioned the biological plausibility of a 48 h infusion of any drug having a significant effect on 180 day mortality in patients with AHF. While the ability of a brief infusion of a thrombolytic can clearly improve survival in patients with acute myocardial infarction, our understanding of AHF has not yet revealed a similarly specific ‘clot’ to target. However, an emerging concept of AHF as a combination of a haemodynamic, neurohormonal, inflammatory, and cytokine storm that results in small, but clinically significant end-organ damage suggests that early and effective interventions could have long-term, beneficial effects. Results from RELAX-AHF support this hypothesis, where evidence of myocardial, renal, liver, and other organ protection by serelaxin was associated with a 37% reduction in both CV and all-cause mortality. In addition, there is evidence to support the converse, where a 48 h infusion of milrinone decreased long-term survival compared to placebo with the survival curves continuing to diverge beyond the infusion. While the Pre-RELAX-AHF and RELAX-AHF studies suggest a survival advantage of serelaxin in patients with AHF, mortality was not a primary endpoint in either study and the HF literature is replete with programmes that have failed to confirm early signals of improved survival. Consequently, RELAX-AHF-2 is appropriately powered to detect a clinically meaningful 22% reduction in risk of CV mortality.

While reducing CV mortality is an undisputedly important goal, WHF has only more recently emerged as a clinically meaningful endpoint in itself with increasing recognition also by regulators. In-hospital WHF is generally defined as WHF symptoms and signs requiring an intensification of therapy and occurs in a variable proportion of patients admitted for AHF, ranging from 5 to 42%. Worsening HF is associated with a prolonged length of hospitalization, increased release of biomarkers related to myocardial damage and renal dysfunction and, more importantly, with a poorer long-term outcome both with respect to rehospitalizations and mortality. The clinical importance of WHF has been demonstrated in retrospective analyses of patient databases and intervention trials, as well as in a recent pooled analysis of 3691 patients from AHF trials. The occurrence of WHF is also sensitive to treatment as it may be reduced by drugs active on symptoms in the patients with AHF, and serelaxin treatment in RELAX-AHF was associated with a 30% decrease in WHF within 14 days. Due to these encouraging results and the importance of this event to patients, RELAX-AHF-2 was designed from its initiation to include robust report forms and detailed documentation to appropriately collect and characterize WHF events, an endpoint elevated from exploratory in RELAX-AHF to key secondary in RELAX-AHF-2 since the trial initiation. Moreover, after a strategic reconsideration following the reviews of marketing authorization applications submitted based on RELAX-AHF results to health authorities worldwide and after further consultations with regulators, the Sponsor and the Executive Committee decided to further elevate WHF through day 5 to a second primary endpoint in RELAX-AHF-2, in addition to CV mortality through day 180. Thus, RELAX-AHF-2 is designed to definitively assess the effect of serelaxin on WHF as an additional primary endpoint of the trial.

In conclusion, RELAX-AHF-2 is expected to answer a key question in AHF therapeutics: can a 48 h infusion of serelaxin reduce 180 day CV mortality of patients admitted for AHF or reduce the occurrence of WHF? If this question is answered in agreement with previous evidence from Pre-RELAX-AHF and RELAX-AHF, it will revolutionize therapy for patients with AHF and for the first time provide an addition to the AHF therapeutic armamentarium with definitive outcome data.

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**Supplementary Information**

Additional Supporting Information may be found in the online version of this article:

Appendix S1. RELAX-AHF-2 Amendments.

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**Conflict of interest:** J.R.T. reports grants, personal fees, and non-financial support from Novartis, during the conduct of the study; grants, personal fees, and non-financial support from Amgen, Bayer, Bristol-Myers Squibb, and Novartis; personal fees and non-financial support from Cytokinetiks, and Trevena; personal fees from Relypsa, and ZS Pharma, outside the submitted work. M.M. reports personal fees from Novartis, during the conduct of the study; personal fees from Bayer, Servier, Relypsa, and Stealth Therapeutics, outside the submitted work. A.A.V. reports grants and personal fees from Novartis, during the conduct of the study; personal fees from Boehringer, personal fees from GSK, grants and personal fees from Vifor, grants and personal fees from Stealth, grants and personal fees from Bayer, personal fees from Trevena, outside the submitted work. P.P. reports personal fees from Novartis, during the conduct of the study; grants and personal fees from Vifor Pharma, personal fees from Amgen, personal fees from Servier, personal fees from Bayer, grants and personal fees from Coridea, personal fees from Celladon, personal fees from Cardiorentis, grants from Singulex, outside the submitted work. P.S.P. reports personal fees and non-financial support from Novartis, during the conduct of the study; personal fees and non-financial support from Bristol-Myers Squibb, Relypsa, scPharmaceuticals, personal fees from Medtronic and Trevena, personal fees and other from Roche Diagnostics, and grants from PCORI, outside the submitted work. B.H.G. reports personal fees and non-financial support from Novartis, during the conduct of the study; he is Committee member of trials sponsored by Novartis, Bayer, Cardiorentis, Vifor, Servier, outside the submitted work. G.M.F. reports grants and personal fees from Novartis, during the conduct of the study; grants and personal fees from Amgen, grants from Trevena, personal fees from GSK, personal fees from Bristol-Myers Squibb, personal fees from Myokardia, grants from Otsuka, outside the submitted work. B.A.D. and G.C. report grants and personal fees from Novartis, during the conduct of the study; grants from Celyad, Trevena, Inc., grants from Sorbent Therapeutics, ChanRx, Laguna Pharmaceuticals, Singulex, outside the submitted work. B.A.D. and G.C. are employees of Momentum Research, Inc. L.B.M., C.G., M.W., T.A.H., and T.S. report personal fees from Novartis, during the conduct of the study; personal fees and other from Novartis, outside the submitted work, and are employees of Novartis.

**Appendix**


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