β-adrenergic receptor responsiveness in aging heart and clinical implications

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INTRODUCTION

Elderly healthy individuals have a reduced exercise tolerance and a decreased left ventricle inotropic reserve related to increased vascular afterload, arterial-ventricular load mismatching, physical deconditioning and impaired autonomic regulation (so called “β-adrenergic desensitization”). Adrenergic responsiveness is altered with aging and the age-related changes are limited to the β-adrenergic receptor density reduction and to the β-adrenoceptor-G-protein(s)-adenyl cyclase system abnormalities, while the type and level of abnormalities change with species and tissues. Epidemiological studies have shown an high incidence and prevalence of heart failure in the elderly and a great body of evidence correlate the changes of β-adrenergic system with heart failure pathogenesis. In particular it is well known that: (a) levels of catecholamines are directly correlated with mortality and functional status in heart failure, (b) β1-adrenergic receptor subtype is down-regulated in heart failure, (c) heart failure-dependent cardiac adrenergic responsiveness reduction is related to changes in G proteins activity. In this review we focus on the cardiovascular β-adrenergic changes involvement in the aging process and on similarities and differences between aging heart and heart failure.

Keywords: β-adrenergic receptors, β-adrenoceptor desensitization, β-adrenoceptor down-regulation, G-protein coupled receptor kinase, aging heart, failing heart, exercise
reduced plasma clearance of catecholamine (Ng et al., 1993; Esler et al., 1995). The “β-adrenergic desensitization,” at least in part, is due to the reduction of β-adrenergic receptor plasma membrane density described in hearts of both senescent animals and elderly humans (White et al., 1994; Xiao et al., 1998). The β-adrenergic receptors are members of the G-protein-coupled receptor family, which acts by coupling with guanine nucleotide binding proteins, and the age-induced decrease in β-adrenoceptor responsiveness is characterized at the molecular level by decreased activation of adenylyl cyclase and reduced production of cAMP. Beside β-adrenergic receptor down-regulation, another crucial age-related alteration of this signaling pathway seems to be the coupling of the β-adrenergic receptor to adenylyl cyclase via the Gs protein, which leads to a reduction in the ability to increase cAMP and to activate protein kinases. Some studies have also reported an increase in Gsat activity as a possible additional mechanism in “β-adrenergic desensitization.” Moreover, the reduction in the efficacy of cardiac β-adrenoceptor stimulation with aging could be also related to other mechanisms, such as the upregulation of G protein-coupled receptor kinases (Rengo et al., 2012a), whereas the role of these kinases in aging heart is controversial. From an overall data analysis on the role of aging in β-adrenoceptor regulation in human and animal hearts it is possible to conclude that the reduced response to β-agonists is common to all species and all cardiac tissues investigated. Moreover, the age-related changes are limited to β-adrenoceptor-G-protein (s)-adenylyl cyclase system abnormalities, while the type and level of abnormalities change with species and tissues. These differences could explain the inconsistency in results obtained in different experimental models of aged heart. Interestingly, several evidence suggest that the β-adrenergic receptor system plays an important role also in heart failure pathogenesis. In fact, it is well known that (a) the levels of catecholamines are directly correlated with mortality and functional status in heart failure, (b) cardiac β-receptors, in particular β1 subtype, are downregulated in heart failure and (c) heart failure-dependent cardiac adrenergic responsiveness reduction is related to adrenoceptor kinases and Gsat increased activities.

AIMS
This review focuses on (a) the development of knowledge on aging heart over the years, (b) the changes involving the sympathetic system in relationship to the cardiovascular aging in different species, (c) the clinical implications of changes in β-adrenergic mechanisms in the aging heart and (d) the similarity between aging and failing heart.

β-ADRENERGIC SIGNALING IN THE HEART AT MOLECULAR LEVEL
For the first time the existence of β-adrenergic receptors (β-AR) was described in 1948 by Alquist (1948). At the present three subtypes of β-AR: β1-AR, β2-AR, β3-AR have been recognized. A fourth subtype has been proposed and investigations have been recently clarified its functioning and localization (Lewis et al., 2004). At the beginning it was thought that only β1-AR subtype was expressed in the cardiac cells. However, many studies provided evidence that both heart β1-AR subtype and β2-AR subtype (Lemoine and Kaumann, 1991; Altschuld et al., 1995; Lonardo et al., 2005), coexist in humans as well as in animals. In the human heart approximately 80% of the β-AR subtype expressed belong to the β1-AR, followed by 20% of the β2-AR subtype (Lakatta and Levy, 2003). It is important to underline that this β-AR expression proportion has been observed in the non-failing young, but not in elderly human heart.

Modern molecular biology techniques and radio-ligand binding studies have shown that major expression and main contribution in the contractile functioning of cardiac cells belong to the β1-AR subtype (Benovic et al., 1991; Borea et al., 1992). In humans, as well as in other animals with relative big body weight like sheep, dogs or cats, the β1-AR and β2-AR are both significantly present, while in other small animals like rats or guinea-pigs the presence of β2-AR is undetectable. Some other studies found the presence of β2-AR in rat hearts but not localized in ventricular myocytes (Buxton and Bruton, 1985). Even the different specie-dependent contribution of β-AR and the difficulties to find a perfect experimental model, it is well known that in all different species the general mechanical pathway is always related to adenylyl cyclase (AC) activation, cyclic AMP (c-AMP) formation, Protein Kinase A (PKA) activation and G-Protein Coupled Receptor Kinase (GRK) activation. β-AR are members of the G-Protein Coupled Receptors (GPCRs) family which acts by coupling with Guanine nucleotide binding proteins (Rengo et al., 2012a). β1-AR subtype is coupled to the stimulatory G protein (Gs). Gs protein is a heterotrimetric protein made up of α, β, and γ subunits. The presence of β-AR agonists induces the dissociation of Gs protein in two subunits: α subunit and β-γ subunit. The primary effect of this dissociation is the activation of AC that catalyzes the conversion of ATP to c-AMP, a second intracellular messenger, and induces the activation of c-AMP dependent PKA. Serine and threonine residues of many regulatory proteins are phosphorylated by PKA. These regulatory proteins include: β-AR themselves, myofilament proteins (troponine I and C protein), membrane proteins (phospholamban—PLB, L-type Ca2+ channels, Sarco/Plasmatic Reticulum—SR, Ca2+/ATPase inhibitory protein). The stimulation of β-AR modifies not only the cardiac excitation and contraction but also other cellular functions such as gene transcription and growth, and can induce death. An important role for the above mentioned functions has played by the activation of Mitogenic-Activated Protein Kinase (MAPK). Moreover, these kinases are thought to be implicated in the regulation of several vital cellular processes, including differentiation, proliferation, growth, and death (Van Biesen et al., 1995). Ultra Violet light, osmotic stress and heat shock can activate MAPK signaling cascades and GPCRs play a pivotal role in the regulation of MAPKs, particularly of the extracellular signal-regulated kinase (ERK1/2) MAPK. One major pathway of GPCRs-mediated activation of MAPKs is dependent on “transactivation” of a group of receptor tyrosine kinases, such as epidermal growth factor and insulin-like growth factor. Additionally the activation of p38 MAPK, also called “stress-activated protein kinase,” is associated with the initial signs of cardiac hypertrophy in response to “in vivo” pressure overload or ischemic/reperfusion injury (Bogoyevitch et al., 1996; Wang et al., 2013).

The β-AR stimulation induced by catecholamines is also responsible for the Ca2+ influx, that by itself triggers a potential release of Ca2+ from SR, acting on the ryanodine receptors.
The intracellular Ca\textsuperscript{++} release activates contractile proteins, finalizing the muscular contraction (positive inotropic effect). Then, intracellular Ca\textsuperscript{++} is removed from the cytoplasm by the SR-Ca\textsuperscript{++}/ATPase pump and the Na\textsuperscript{+}/Ca\textsuperscript{++} exchange. The further acceleration of Ca\textsuperscript{++} removing leads to the muscle relaxation. The maximum velocity of relaxation is defined as positive lusitropic effect. As a result of PKA activation, the \(\beta\)-AR stimulation triggers the G-Protein Coupled Receptor Kinase family, like GRK2. GRKs are a family of serine/threonine protein kinases that phosphorylates GPCRs only when the receptors are in the activated (agonist-bound) state. When \(\beta\)-ARs are stimulated by agonists, the \(\beta\)-\gamma subunits interact with GRK2 bringing the kinase from the intracellular to the transmembranic localization, phosphorylating the \(\beta\)-ARs, becoming target for binding of \(\beta\)-Arrestin proteins. The \(\beta\)-Arrestin bounds to these receptors and prevents their further coupling to the G-protein, reducing the level of functional receptors, inducing the internalization of receptors and, as final result, their decreased density and desensitization (Freedman et al., 1995).

“In vitro” studies showed further mechanisms induced by stimulation of \(\beta_1\)-AR. For example, persistent stimulation of \(\beta_1\)-AR is able to activate Calmoduline Dependent Kinase II, without the implication of PKA pathway. This mechanism induces cardiomyocyte hypertrophy and could explain the well-known relationship between adrenergic stimulation and cardiac hypertrophy\textsuperscript{23-24} (Ramirez et al., 1997; Morisco et al., 2000). In addition to the cardiac effects, \(\beta_1\)-ARs regulate the release of renin, the activation of Renin-Angiotensin-Aldosteron (RAA)-system and the lipolysis.

\(\beta_2\)-AR, one of the first receptors identified, belongs to the GPCRs family and plays an important role in the cardiovascular and respiratory physiology. Its main effects are related to vasodilatation and bronchodilatation (Corbi et al., 2013b). In addition \(\beta_2\)-AR is responsible for glycogenolysis (Corbi et al., 2002) and relaxation of uterine muscle. Despite similarities \(\beta_1\)-AR and \(\beta_2\)-AR present different signaling pathways. The \(\beta_2\)-AR is coupled to the G\(_i\) protein and to the G\(_s\) protein too (dual coupling of \(\beta_2\)AR to G\(_i\) and G\(_s\) protein). There is also evidence that \(\beta_2\)-AR signaling is coupled to an independent pathway like the Na\textsuperscript{+}/H\textsuperscript{+} exchanged regulatory factor (Hall et al., 1998).

The effects of \(\beta_2\)-AR G\(_i\) stimulation are not identical to them obtained from \(\beta_1\)-AR stimulation. However, similarly to \(\beta_1\)-AR, \(\beta_2\)-AR-G\(_i\) stimulation increases the c-AMP and PKA activity. Recent studies have demonstrated that the effect of \(\beta_2\)-AR c-AMP/PKA stimulation is limited to the subsurface membrane of the L-type Ca\textsuperscript{++} channels without cellular signal transmission. As a result of this mechanism it is observed a positive inotropic effect without influence on the intracellular Ca\textsuperscript{++} transient decay time, changes in myofilaments sensitivity to Ca\textsuperscript{++} and increase SR Ca\textsuperscript{++} uptake. Obviously \(\beta_2\)-AR stimulation does not affect the relaxation time (lusitropic effect) as \(\beta_1\)-AR does (Kuschel et al., 1999; Xiao et al., 1999).

Surprisingly the Ca\textsuperscript{++} influx, the PKA and the c-AMP levels apparently do not show significant association after \(\beta_2\)-AR stimulation in studies using adult rat and canine models. Even if the reason of this dissociation remains unclear, the role of c-AMP modulation in the contractility of cardiac muscle is well established. In heart animal models forskolin induces c-AMP levels augmentation increasing the inotropic effect. Moreover, it has well demonstrated that the Ca\textsuperscript{++} influx mechanism is exclusively mediated by c-AMP pathway (Xiao et al., 1994).

On the other hand, the G\(_{\alpha_i}\) protein subunit inhibits the adenylcyclase enzyme activity. The \(\beta_2\)-AR G\(_i\) signaling inhibits the c-AMP synthesis and has negative effects on the PKA activation. Persistent activation of \(\beta_2\)-AR-G\(_{\alpha_i}\) signaling activates in turn the phosphoinositol3- Kinase (PI3-K), an important downstream messenger that triggers the antiapoptotic factor Akt and seems to have a cardioprotective role. (Zhu et al., 2001; Cannavo et al., 2013).

The model of \(\beta_2\)-AR dual coupling of to multiple G protein (G\(_i\) and G\(_s\) protein), is not well clarified. Several evidence indicate that \(\beta_2\)-AR–G\(_i\) signaling compartmentalizes the \(\beta_2\)-AR-G\(_i\)-c-AMP signaling. Disrupting the G\(_i\) functioning by a potent G\(_i\) inhibitor like Pertussis Toxine (PTX) induces an enhance in the phosphorylation of PLB and an increased inotropic effect after \(\beta_2\)-AR stimulation. In this occasion the \(\beta_2\)-AR signaling is comparative to the \(\beta_1\)-AR signaling\textsuperscript{31} (Xiao et al., 1995).

The \(\beta_2\)-AR phosphorylation by PKA and GRK2 switches the \(\beta_2\)-AR receptor coupling from G\(_i\) to G\(_s\). As demonstrated in several studies, the \(\beta_2\)-AR-G\(_s\) coupling is suppressed after GRK2 activity inhibition. In a near future, it may be possible to prevent important structural changes, like myocardial stiffness, reactive fibrosis and remodeling present in the aging and failing heart, modifying the GRK2 and the G\(_s\) activated or inhibited status. Another candidate mechanism, underlying the compartmentalization of \(\beta_2\)-AR–G\(_i\)-c-AMP-PKA signaling in response to the \(\beta_2\)-AR-Gi coupling, is the structural restriction of PKA diffusion by muscle specific protein A kinase anchovy proteins (AKAP) (Enns et al., 2009). The phosphorylation of AKAP plays multiple roles including: ions influx, contraction, transcription of different genes, phosphorylation of multiple intracellular targets in cardiac myocytes including the L-type Ca\textsuperscript{++} channel in the sarcosome, the ryandone receptor (RyR2), and phospholamban in the SR. Deficiencies in this pathway have been linked to cardiomyopathy in humans, due to reduced phosphorylation of downstream targets such as cardiac troponin (McConnachie et al., 2006). Moreover, in genetically manipulated models it is obtained an increased positive inotropic effect after disrupting the APK kinase anchovy protein (Marshall, 1995; Spindler et al., 2013).

**AGE-INDUCED CHANGES IN THE \(\beta\)-AR SIGNALING**

Adrenergic signaling is a very important for cardiovascular physiology. In conditions involving physical or psychological stress high levels of cathecolamines like norepinephrine and epinephrine are released from the adrenal medulla. It is well known that the action of cathecolamines is mediated by adrenergic receptors and the effects on cardiovascular system include: increased heart rate and myocardial contractility force and relaxation, increased cardiac output, reduced left ventricular afterload, a diversion of blood flow from the skin and splanchnic vessels to those supplying skeletal muscles, bronchial dilatation and a decline in metabolic activity (Young and Landsberg, 1998). Generally the age-related decrease in \(\beta\)-adrenoceptor response has...
been explained by a mechanism called “β-adrenoceptor desensitization.” It is a process characterized by β-AR molecular changes: phosphorylation of receptor structures enhanced by an agonist-receptor bind state, that induces the reduction of receptors density and their internalization. This process is well-described also in the heart failure (Rengo et al., 2012a). By aging a post-synaptic “β-adrenoceptor desensitization” responsible for a reduction in the autonomic modulation of the cardiac system, especially during physical exercise (Ehsani, 1987; Scarpace et al., 1994), is observed. During exercise the stroke volume increase is similar in young and older, but the mechanism of this increase is different. The elderly tends to augment stroke volume more through cardiac dilatation with an end-diastolic volume increase, while the young shows an increase in the ejection fraction without cardiac dilatation. Moreover, during exercise the older has a lower increase in heart rate and a greater raise in blood pressure (Ferrara et al., 2006; Corbi et al., 2012b). In particular in the aging human heart the maintaining of cardiac output during exercise is supported more heavily by the Franck-Starling mechanism and less by sympatho-stimulation demonstrating the presence of an age-related change in the adrenergic modulation. It is interesting to point out the similarity observed in terms of hemodynamic profile between younger β-blocked subjects and healthy elderly without β-blockers treatment in human models without HF (Fleg et al., 1994). Younger β-blocked subjects showed during exercise reduced heart rate and contractility index, and increased left ventricular volume, in short terms they apparently “looked older” but the results of β-AR blockade were greater in the young subjects compared to older ones, suggesting that β-AR responsiveness reduces with aging.

Interestingly, Leosco et al. (2007) demonstrated that exercise training alone as well as metoprolol alone or in combined therapy (exercise + metoprolol) improved the β-AR signaling in the aged heart suggesting a similar effect on β-AR signaling of chronic treatment with β-blockers and chronic exercise training. Furthermore they found an increased β-AR density inducing a reversible level of β-AR desensitization. The overall reduction in cardiac reserve is responsible for the decreased exercise tolerance and for the impairment in cardiac response at exercise in terms of reduced ejection fraction at peak and heart rate responsiveness during dynamic exercise. The responsibility of this response could be related to increased vascular afterload, arterial-ventricular load mismatching, reduced intrinsic myocardial contractility, physical deconditioning and impaired autonomic regulation (so called “β-adrenergic desensitization”). The mechanism of this impaired autonomic regulation induced by age is not completely clarified, although it is hypothesized that the increase in catecholamines levels plays an important role in the “β-adrenergic desensitization” in aging and failing heart. In both conditions, as a result of a reduced plasma clearance and an increased spillover from the tissues, the level of circulating catecholamines further rises. This prolonged catecholamines action seems to be also related to the age-dependent reduction of the catecholamines re-uptake transporter localized in the sympathetic nerve terminals (Leineweber et al., 2002). However, controversial opinions exist in the present literature about the changes in the systemic norepinephrine levels with age and the relationship between the age-related increased levels of catecholamines and the suggested mechanisms of age-related “β-adrenergic desensitization” (Folkow and Svanborg, 1993; Esler et al., 1995).

The age-related “β-adrenergic desensitization,” a possible adaptive mechanism, has been observed in both animals and humans. An age-related effect on the maximum contractility response to isoproterenol has been demonstrated in arterially perfused interventricular septa from adult and senescent rats (Froehlich et al., 1978), and a reduced inotropic response of aged myocardium to catecholamines has been found in superfused trabeculae (Jiang et al., 1993). At molecular level, in particular, the decrease in β-adrenoceptor responsiveness has been related to changes in G-proteins and kinases activity even if differences in the level and extent of these changes exist among different species. Concerning the β-AR density, the first studies performed in circulating lymphocytes did not show any important age-related changes in β-AR density (Abrass and Scapace, 1981; Landmann et al., 1981). Also other data obtained from young and old rats did not report any changes in β-AR number (Bohm et al., 1993) did not find any changes in β-AR number but noticed a G_{i} increased content. This study hypothesized that G_{i} might serve as a age-related regulator of cardiac AC activity in the absence of β-adrenoceptor changes. Interestingly Gudmundsdottir et al. (1991) showed that dietary fat and age modified the density of Ca^{++} channels and reduced the β-AR number in the rats.

Cerbai et al. (1995) confirmed that β_{2}-AR as well as β_{1}-AR are both functionally present in rat hearts, but only the β_{1}-AR density was reduced with aging. In human aging heart White et al. (1994) found that the β_{1}-AR down regulation mechanism was linked to the reduced number of β_{1}-AR in a high affinity agonist binding state. In our experience, using a model of myocytes obtained from human failing (donors) and non-failing hearts (small biopsies from elderly and young patients undergoing coronary vein graft with preserve ventricular function), the contractile response to β-adrenoceptor stimulation has been found to be strongly reduced in single myocytes from failing human ventricle, but a part of this reduction was statistically related to the patient’s age (Dobson et al., 1990; Davies et al., 1996). At beginning, it was thought that the desensitization mechanism of β-AR in the elderly is related to the increased levels of adenosine because its clear antiadrenergic action in the heart, as suggested by studies performed in guinea pig aging hearts. However, utilizing single isolated left ventricular cardiac myocytes from hearts of animals at different age, other studies showed that the age-related contractility impairment during β-adrenergic stimulation was confined to the β-adrenergic pathway with reduced net production of c-AMP, that could not be explained by the increased adenosine stimulation (Ferrara et al., 1995).

Ferrara et al. (1995), in a study performed on myocytes isolated from the young and old guinea-pigs, observed that there was a pronounced age-related decrease in contractility of myocytes after both isoproterenol and high Ca^{++} concentration stimulation. However, the Ca^{++} influence was less significant concluding that a general diminished contractility of single myocytes occurs during the physiological aging and these effects were more marked for β-AR stimulation than for high Ca^{++}, suggesting a specific lesion in the AC related pathway. In a further
investigation, using the same experimental model, the same authors (Ferrara et al., 1997a) studied the role of the inhibitory G-proteins (G_i) in the decline of contractility related to β-stimulated activity. In these experiments they found that β-AR number was decreased by 27% in senescent animals and the G_i activity, detected by PTX-catalyzed ADP ribosylation, was significantly increased in aged animals, while immunodetectable level of G_i was not modified. The authors concluded that increased G_i activity contributes significantly to the decreased response to β-AR stimulation in myocytes in aged guinea-pigs.

In different species (rats) Xiao et al. (1998) found a clear age-related reduction in contractility in response to β-AR subtype stimulation associated to a non-selective decrease in the density of both β-AR subtypes and a reduction in membrane AC response to either β1-AR or β2-AR stimulation. Moreover, the activity of β-AR kinase GRKs, and of G_i did not significantly change with aging, suggesting that the marked decreased response to β-ARs stimulation with aging in rat ventricular myocytes was linked to a decrease in both β-AR subtypes densities and a reduction in membrane AC activity, while neither GRKs nor G_i proteins appear to play a role in this mechanism. This apparent contrast with the results from Ferrara et al. (1997a) could be referred to the different animal models used for the experiments.

As in the failing heart, the role of chronic sympathetic drive in the β-AR desensitization in the aging heart can be hypothesized. Because the role of GRKs in the aging heart could be defined controversial same studies have been planned to focus on the role of GRKs in the aged cardiovascular system.

Schutzer et al. (2001) examined the correlation of GRKs level to age-related modifications in aorta of old rats. In particular they studied the age-related changes in distribution of GRK subtypes 2, 3, or 5 and β-arrestin (cytoplasmic vs. crude membrane preparations), and demonstrated that GRKs are implicated in the reduction of β-AR-mediated vasorelaxation with advancing age, suggesting a strong evidence that increased GRK activity plays an important role in cardiovascular physiology of aging. However, it is important to underline that in this study was not performed detection of GRKs in the heart tissues as it was performed in the previous studies mentioned above, explaining this controversial results. On this basis Leosco et al. (2003), studying the effects of exercise, found a reduced β-AR expression in aged rat carotid arteries, and the exercise itself restored the age-associated blunted β-AR responsiveness. This could explain with the age-related reduced adaptation of cardiovascular system to different stressors.

For the first time Leinweber et al. (2003) studied the possible GRKs activity alterations related to aging in humans. The study evaluated the cytosolic and membranous levels of GRKs in right atria from children with congenital heart disease undergoing cardiac surgery, elderly with coronary heart disease but not suffering from heart failure undergoing coronary artery by-pass grafting, and from a small group of elderly patients with heart failure also undergoing cardiac surgery. The main result was that neither cytosolic nor membranous GRKs activity were modified in elderly compared to children, while, as confirmed in previous studies (Ungerer et al., 1993; Ping et al., 1997; Vinge et al., 2001; Rockman et al., 2002; Iaccarino et al., 2005) there was a notable up-regulation in the GRKs activity in the failing heart. The reason because the β-AR desensitization in the aging heart is not associated to GRKs up-regulation is not well clarified at present. It can be hypothesized that the GRKs levels are more affected by acute triggers as in heart failure happens and not influenced by a gradual chronic “aging” process of β-adrenergic drive, or at least during this life-long process GRKs levels are associated to other adaptive mechanism that may influence their up-regulation. In Table 1 the main general age-related changes in β-adrenergic signaling are described.

**CLINICAL IMPLICATIONS OF ALTERATIONS IN β-ADRENOCEPTOR MECHANISMS IN THE AGING HEART. RELATIONSHIP WITH HEART FAILURE**

Heart failure and associated clinical implication induced by alterations in β-adrenocceptor mechanisms are a central problem for elderly population (Ferrara et al., 1997b). Epidemiological studies showed high prevalence and incidence of heart failure in the elderly (Corbi et al., 2008; Roger et al., 2011) due to an increased longevity. The incidence doubles with each decade of life and prevalence rises almost 10% of those older than 80 years (Cacciatore et al., 1998). About 50% of all heart failure cases is found in patients older than 70 (Roger et al., 2011). Although nowadays an important decrease in the mortality for heart disease, cardiovascular disease is the most frequent single cause of death in the elderly population. The prognosis of heart failure among elderly is poor with a 4 year survival of only around 50% (Cacciatore et al., 1998) heart failure is also one of the most important causes of comorbidity and hospitalization rising health costs.

It is important to underline that heart failure in the elderly appears when cardiovascular structural and functional age-related changes are already evident. Moreover, the age-associated changes in cardiovascular structure/function are involved in the increased risk for heart failure in older people. An age-dependent increase in left ventricular wall thickness in men and women without hypertension has been described in the Framingham Heart Study and Baltimore Longitudinal Study on Aging (Ho et al., 1993). These modifications in structure of heart are characterized by an increase in cardiomyocyte size and myocardial thickness modifying the contractile efficiency and increasing left ventricular mass (Olivetti et al., 1991; Khouri et al., 2005). As a result of age-related chronic stress, the myocardium can undergo cardiomyocyte death including necrosis, apoptosis or autophagy. This process induces initially a compensatory remodeling characterized by the alterations of extracellular matrix (ECM) composition involving the synthesis of myofibroblasts, the degradation of collagen through TGF-B signaling. These alterations lead to the hypertrophy of the remaining cells and to pathologic remodeling, with consequent reactive fibrosis that increases cardiac stiffness and reduces the cardiac compliance (Boyle et al., 2011).

It has been reported that the process of cardiac aging, as well as the progressive heart failure, is also characterized by the impaired Ca^{++} reuptake and the decreased SR Ca^{++} storage. In some studies this decline has been explained by the modification of SERCA2 protein. This impaired Ca^{++} reuptake is responsible for the delays in the ventricular relaxation (Sucharov et al., 2006). The myocardial remodeling, myocardial stiffening and the
delayed ventricular relaxation compromise the left ventricular filling in early diastole. In order to maintain an adequate left ventricular filling in elderly the atrial contraction (A wave) is further increased leading to the atrial hypertrophy increasing so the risk for atrial fibrillation. However, these data have not been confirmed utilizing other epidemiological and autopsy-based studies (?) in subjects free from coronary artery disease and hypertension. In fact, these studies have demonstrated no significant changes in cardiac mass in women and a decrease in left ventricular mass in men. Based on these and other studies it has been shown that there is no significant changes in left ventricular mass in the elderly women, but there is a reduction of cardiac mass in men. Based on these and other studies it has been shown that there is no significant changes in left ventricular mass in the elderly women, but there is a reduction of cardiac mass in men (cardiac sarcopenia). Regarding to this Lin et al. (2008) demonstrated, combining experimental and mathematical models, in mice an age-related cardiac sarcopenia and that LV remodeling due to increased end diastolic pressure could be an underlying mechanism for age-related LV dysfunction. Senescence is characterized by a reduction of myocytes cell number. In order to maintain the cardiac function, some compensatory mechanisms are induced such as: increase of individual myocyte size, increase of intracellular glycogen storage and reactive fibrosis. This changes induce a depressed cardiac function in elderly. Moreover, asymmetric increase in the intraventricular septum does not influence the total cardiac mass. Nevertheless these changes do not modify significantly the systolic cardiac function at rest as demonstrated by echocardiographic and radionuclide studies in normotensive healthy population (Hess et al., 2002; Khouri et al., 2005). Instead the pattern of diastolic function changes with aging and plays a pivotal role in increased prevalence of diastolic heart failure in the elderly. There are many similarities between the aging and failing heart. For example, morphologically myocytes hypertrophy and cardiac fibrosis occur in both aging and failing heart that exhibit decreased diastolic function and increased ventricular mass. In both there is a decreased functional reserve and defective cardiac energetics. The age-related changes in β-AR behavior are quite similar to those related to failing heart and these changes may be induced by the compensatory adrenergic drive activation. The stimulation of adrenergic receptors by catecholamines is the most important regulatory mechanism for cardiovascular performance. It is well known that the levels of catecholamines are directly correlated with mortality and functional class of heart failure (Cohn et al., 1984). The cardiac β-receptors, in particular the β1 subtype, are down-regulated in heart failure and the heart failure-associated reduced cardiac adrenergic responsiveness are related to an increase in Gαi activity and in activity of the adrenoceptor kinases. Failing heart-associated β-AR down-regulation seems to effect the β1-AR but not the β2-AR density and this abnormality is also related to GRK2 and Gβ up-regulation (Bristow et al., 1986). Physical performance as well as other stressors stimulate the adrenergic system increasing heart rate, myocardial contractility and relaxation, reducing left ventricular afterload and redistributing blood to working muscle. As above described, this pattern modifies during aging. A possible explanation of this behavior is in relationship with the chronic and gradual sympathetic hyperactivity. The association of failing to aging heart physiologic changes is not important only for the worldwide health care system but also for explaining the worst of quality of life in these patients. For these reasons since about 30 years the research studies about the relationship between adrenergic system and heart failure are interestingly increasing. In the early 1982, Bristow et al. (1982) studying the myocardium contractility “in vivo,” examined the β-AR pathway in failing hearts obtained from human subjects undergoing cardiac-transplantation. It was noticed an important reduction in the contractile response to isoproterenol in these “in vivo” cardiac cells and also an evident reduction of the β-AR density. The same author (Bristow et al., 1986) after a couple of years, developed a radiolig and biopsy method for identifying β-AR in human ventricular myocardium. This method it is thought to be useful in the direct analyses in the β-AR density

Table 1 | Cardiovascular system: the main general age-related changes in β-adrenergic signaling.

| Author                      | Constituent | Species                          | Changes  |
|-----------------------------|-------------|----------------------------------|----------|
| Abrass and Scapace, 1981    | β-AR density| Human (circulating lymphocytes)  | Unchanged|
| Landmann et al., 1981       | β-AR density| Human (circulating lymphocytes)  | Unchanged|
| Gudmundsdottir et al., 1991 | β-AR density| Rat (heart)                      | Decreased|
| Bohm et al., 1993           | β-AR density| Rat (heart)                      | Unchanged|
| White et al., 1994          | Gβ1 content | Rat (heart)                      | Increased|
| Xiao et al., 1998           | β-AR density| Rat (heart)                      | Decreased|
| Cerbai et al., 1995         | β-AR density| Guine-pig (heart)                | Decreased|
| Ferrara et al., 1997a,b     | Gβ1, β-AR   | Guine-pig (heart)                | Unchanged|
| Ferrara et al., 1997a,b     | Gβ1 activity| Giuine-pig (heart)               | Increased|
| Xiao et al., 1998           | β1, β-AR    | Rat (heart)                      | Decreased|
| Xiao et al., 1998           | Gβ1 activity| Rat (heart)                      | Unchanged|
| Schutzer et al., 2001       | β-AR        | Rat (aorta)                      | Decreased|
| Schutzer et al., 2001       | GRK2, GRK3, β-arrestin | Rat (aorta) | Increased|
| Leinweber et al., 2003      | GRK         | Human (heart)                    | Unchanged|
and in the study of down-regulation. The development of this new method opened a new window in the molecular pharmacology of adrenergic system. Brodde et al. (1989) concluded that the β-AR density level reduction is associated to the severity of congestive heart failure, and in valvular heart failure exist a decrease in either β1-AR and β2-AR density. β-AR desensitization in the failing heart is proved from many authors, but this is not the one and only mechanism involved in a complex disease like heart failure. Feldman et al. (1988) observed the increased activity of α G40 complex in the human failing heart and considered as a possible new marker for the severity of failing heart. Fu et al. (1992) also concluded that the role of Gβ protein is crucial in the failing heart but it was observed an increase in functional activity rather than in its amount. In 1997 Ping et al. (1997) during the pacing induced congestive heart failure (CHF) examined the alteration of β-ARs, AC activity and GRKs. They concluded that in the advanced stages of CHF, β-ARs and AC are down-regulated while there is an increase in the GRKs levels since the early stages of CHF. Ungerer et al. (1993) studying the involvement of β-AR in human heart failure models found for the first time an inverse correlation between β-AR density and β-ARKs. The β-AR density is reduced but also the remaining receptors are less effective. Initially the catecholamines released from sympathetic nerves try to restore the cardiac functioning by acting on the β-AR. In long term the enhanced sympathethic drive causes the β-AR desensitization. At least part of this adaptative mechanism should protect heart from further sympathetic activation, but the overall cardiac performance is depressed.

The genetic engineering and the need to establish a quite similar model of human disease have driven the scientific research to develop transgenic animal models. These model are produced by using DNA microinjections in animals or by using another methodology like the gene knock-out models. In 1996 Rockman et al. (1996) in order to investigate the β-AR alteration and contribution in heart failure, used a model of transgenic mice with overexpression of β-ARK1 inhibitors or β2-AR overexpression. It resulted that overexpressed β-ARK1 inhibitors increased the myocardial contractility and prevented the development of heart failure. A couple of years later Harding et al. (2001) developed a transgenic mice model overexpressing β-ARK1 inhibitor (β-ARK1CT) and calcsequestrin protein. In this model a treatment with β-blockers led to an improvement of cardiac contractile functioning and an longer survival.

The development of animal surgery techniques also helped to obtain similar rat models of heart failure. Vinge et al. (2001) studied the involvement of GRKs in postinfarction heart failure model in rats undergoing ligation of left coronary artery. They concluded that GRK2 and β-arrestin1 are primary regulators in the endothelial function in the heart failure, while the GRK3 and GRK5 play a very important role in the cardiac myocyte functioning. The increased levels of GRK2 was related to the infarction induction suggesting that this protein may precipitate the development of acute heart failure. Iaccarino et al. (2005) found an inverse correlation between the β-ARs expression and the levels of GRK2. This inverse correlation can be explained by the phosphorylation of β-ARs by GRKs and furthermore their internalization and desensitization process. Interestingly the level of GRK3 is associated to

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### Table 2 | Failing heart: the main general changes in β-adrenergic signaling.

| Author            | Constituent                  | Species                  | Changes          |
|-------------------|------------------------------|--------------------------|------------------|
| Bristow et al., 1982 | β-AR density                | Human (heart)            | Decreased        |
| Bristow et al., 1986 | β1-AR density               | Human (heart)            | Decreased        |
| Brodde et al., 1989 | β1, 2-AR density            | Human (heart)            | Decreased        |
| Feldman et al., 1988 | α G40 activity              | Human (heart)            | Decreased        |
| Ungerer et al., 1993 | β-ARK                     | Human (heart)            | Up-regulated     |
| Fu et al., 1992   | Gβ                           | Human (heart)            | Increased        |
| Ping et al., 1997 | β-AR                        | Human (heart)            | Down-regulated   |
| Ping et al., 1997 | AC                          | Human (heart)            | Down-regulated   |
| Ping et al., 1997 | GRKs                        | Human (heart)            | Increased        |
| Rockman et al., 1996 | β-ARK1 Inhibitors (overexpression) | Transgenic mouse (heart) | Increased contractility |
| Harding et al., 2001 | β-ARKct (overexpression)    | Transgenic mouse (heart) | Increased contractility |
| Vinge et al., 2001 | GRK2                        | Rat (heart)              | Increased        |
| Vinge et al., 2001 | GRK3                        | Rat (heart)              | Increased        |
| Vinge et al., 2001 | GRK4                        | Human (heart)            | Up-regulated     |
| Iaccarino et al., 2005 | GRK2                      | Rat (adrenal medulla)    | Down-regulated   |
| Lymeropoulos et al., 2007 | a2-AR                     | Transgenic mouse (adrenal medulla) | Down-regulated |
| Lymeropoulos et al., 2007 | a2-AR                     | Rat (adrenal medulla)    | Up-regulated     |
| Lymeropoulos et al., 2007 | GRK2                      | Transgenic mouse (adrenal medulla) | Up-regulated |
| Lymeropoulos et al., 2007 | GRK3                      | Rat (adrenal medulla)    | Down-regulated   |
| Rengo et al., 2012a,b,c | a2-AR density               | Rat (adrenal medulla)    | Increased        |
| Rengo et al., 2012a,b,c | GRK2                      | Human (lymphocytes)      | Increase         |
| Rengo et al., 2014 | GRK2                        | Human (lymphocytes)      | Increase         |
the degree of heart failure, suggesting an importance of GRKs as a heart failure progression biomarker. In 2007, for the first time, Lymeropoulos et al. (2007) studied the involvement of adrenergic signaling in the pathophysiology of heart failure. The study was performed in two different models, in transgenic mice with cardiac overexpression of SR calcium binding protein calsequestrin (CSQ) and rat model undergoing myocardial infarction. It was noticed an α2-AR down-regulation independent of species and a GRK2 up-regulation. In 2012, in a post-infarction heart failure rat model Rengo et al. (2012b) studied the influence of adrenergic blockade in the in GRK2 and α2-AR adrenal medulla dysregulation. It has been confirmed that there is an up-regulation of adrenal medulla GRK2 and a down-regulation of α2-AR and that their blockade with β-blockers normalizes the level of GRK2. The same author (Rengo et al., 2012c) in 2012 investigated the role of β2-AR in a post-myocardial infarction heart failure rat model. This model underwent an adenosinal-mediated overexpression of β2-AR after 4 weeks of surgery procedure. It was observed that the overexpression of β2-AR improves the angiogenesis process and enhanced the coronary reserve and myocardial blood flow. This pro-angiogenic characteristic of β2-AR was associated to the activation of pro-angiogenic vascular endothelial growth factor, protein kinase B, endothelial nitric oxide synthase VEGF/PKB/eNOS pathway.

Recently returning to the well-known method of detection GRKs levels from the lymphocytes, Rengo et al. (2014) performed a prospective study in a group of heart failure patients who underwent exercise training. It was shown for the first time that the subjects after physical performance presented reduced levels of GRKs and also predicted survival. In Table 2 the main general failing heart-related changes in β-adrenergic signaling are described.

CONCLUSIONS
Cardiovascular diseases are the most common cause of death in elderly population. Increased longevity is associated to an increased heart failure morbidity with poor prognosis among elderly. For this reasons it is very important to understand and clarify the pathophysiological mechanisms that underlay the aging heart process. From the present literature it seems that “β-adrenoceptor desensitization/down-regulation” is a general and common mechanism which explains age- and heart failure-related decrease in β-adrenoceptor response to agonists. In particular, in both aging and failing hearts the decrease in β-adrenoceptor responsiveness is related to changes in G-proteins and kinases activity, although there are differences in the level and the extent of these changes among different studied species. These molecular alterations are responsible for the most structural and functional changes in aging heart. The improved understanding of the inactivation of irinimide could help in the future to develop new therapy approach and ameliorate life quality among elderly population.

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