Human heart as a shock organ in anaphylaxis

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Summary
Anaphylaxis is a potentially fatal, immediate hypersensitivity reaction. Mast cells and basophils, by elaborating vasoactive mediators and cytokines, are the main primary effector cells of anaphylaxis. Mast cells have been identified in human heart between myocardial fibers, perivascularly, in the adventitia, and in the arterial intima. Mast cells isolated from human heart tissue (HHMC) of patients undergoing cardiac transplantation express high affinity immunoglobulin E (IgE) receptors (FcεRI), C3a, C5a, and kit receptors (KIT). Anti-IgE, anti-FcεRI, and immunoglobulin superallergens induce in vitro secretion of preformed mediators (histamine, tryptase, chymase, and renin) and the de novo synthesis of cysteinyl leukotriene C4 (LTC4) and prostaglandin D2 (PGD2) from HHMC. Complement is activated and anaphylatoxin forms during anaphylaxis. C5a and C3a cause the in vitro release of histamine and tryptase from HHMC. Therapeutic (general anesthetics, protamine, etc.) and diagnostic agents (radio contrast media, etc.), which can cause anaphylactoid reactions, activate HHMC in vitro. Low concentrations of histamine and cysteinyl leukotrienes given to subjects undergoing diagnostic catheterisation caused significant systemic and coronary hemodynamic effects. These data indicate that human heart mast cells and their mediators play a role in severe anaphylactic reactions.

Key words
Anaphylaxis – heart – histamine – leukotrienes – mast cells – tryptase

Introduction
Anaphylaxis is the most dramatic and potentially fatal manifestation of immediate hypersensitivity, accounting for more than 500 deaths annually [1, 2]. Anaphylaxis is characterized by laryngeal edema, bronchoconstriction, hypotension, and vascular leakage. It is well known that the heart is directly and/or indirectly involved in severe human anaphylaxis [3, 4, 5, 6, 7, 8, 9]. Cardiac anaphylaxis in humans is characterized by coronary spasm, decreased left ventricular contractility, and/or arrhythmias [10, 11, 12, 13]. Myocardial lesions might be the anatomical basis for the irreversible cardiac failure occasionally associated with systemic anaphylaxis [14]. Coronary spasm or myocardial infarction can occur after insect stings [15, 16] and as a consequence of idiopathic anaphylaxis [17]. Moreover, patients with anaphylaxis may present profound myocardial depression presumably due to the negative inotropic effects of mast cell-derived mediators [18].

Abbreviations
C3aR C3a receptor
C5aR C5a receptor
CH Control heart
CBF Coronary blood flow
CVR Coronary vascular resistance
CysLT2 Cysteinyl leukotriene receptor 2
CysLT1 Cysteinyl leukotriene receptor 1
DCM Dilated cardiomyopathy
ECP Eosinophil cationic protein
FcεRI High affinity immunoglobulin E receptors
HHMC Mast cells isolated from human heart tissue
ICM Ischemic cardiomyopathy
IgE Immunoglobulin E
LTC4 Cysteinyl leukotriene C4
LTD4 Cysteinyl leukotriene D4
LV Left ventricular
MBP Major basic protein
PAF Platelet-activating factor
PGD2 Prostaglandin D2
SCF Stem cell factor

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In this review we draw together evidence that cardiac mast cells and their mediators play a role in systemic and cardiac human anaphylaxis.

**Mast cells are present in human heart**

Several years ago Vincenzo Patella, in collaboration with Bärbel Lamparter-Schummert and Monika Adt, established in our laboratory an elegant technique to isolate and purify mast cells from heart tissue of patients undergoing heart transplantation [19, 20, 21]. Mast cells are present in normal and atherosclerotic human intima of coronary arteries [12, 13, 22, 23, 24]. We have identified and characterized mast cells around coronary blood vessels and between myocardial fibers by electron microscopy in all sections of human hearts [19, 20, 25]. Tab. 1 summarizes the three different localizations of mast cells in human cardiac tissue we have found examining sections of more than 50 hearts [19, 20]. A small percentage (about 5%) of cardiac mast cells appears partially degranulated [23, 19].

**Fig. 1** shows the electron micrograph of a human cardiac mast cell in close contact with the capillary vessel wall. The presence of mast cells close to coronary vessels suggests that circulating antigens, superallergens, autoantibodies (anti-IgE, anti-FcεRI, etc.), complement (C3a and C5a), drugs (general anesthetics, protamine, etc.), and diagnostic agents (radio contrast media, etc.) can easily reach perivascular HHMC. Activated mast cells can in turn release vasoactive substances (histamine, cysteinyl leukotrienes, PGD₂, platelet-activating factor; PAF, etc.) that can affect blood vessels [26].

Cardiac mast cell density and the histamine and tryptase contents of mast cells isolated from failing hearts from patients with idiopathic dilated cardiomyopathy (DCM) and ischemic cardiomyopathy (ICM) undergoing heart transplantation were higher than in control hearts (CH) without cardiovascular disease [27]. Immunologic activation of HHMC induced a significantly greater release of mediators (histamine, tryptase, and LTC₄) in patients with failing hearts than in CH. The latter observation suggests that underlying cardiovascular diseases might represent a risk factor for severe anaphylaxis.

**Mediators synthesized by HHMC**

HHMC contain histamine (≈ 3 pg/cell), tryptase (≈ 24 µg/10⁶ cells) and IgE-mediated activation of HHMC caused tryptase release parallel to histamine secretion [19]. By using the immunogold technique we showed that HHMC contain chymase as well as tryptase [19]. Human chymase generates angiotensin II from angiotensin I, acting as an angiotensin-converting enzyme [28]. Interestingly, supernatants of HHMC challenged in vitro with anti-IgE can convert angiotensin I into angiotensin II, suggesting that chymase released from HHMC play a role in the homeostatic control of blood pressure. Therefore, the activation of HHMC and release of chymase may control blood pressure during anaphylaxis.

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**Table 1**

| Location of mast cells in human heart tissue | Close proximity to myocytes | Increased density in ischemic and idiopathic cardiomyopathy |
|--------------------------------------------|----------------------------|----------------------------------------------------------|
| Interstitial mast cells                     | Close proximity to myocytes | Increased density in ischemic and idiopathic cardiomyopathy |
| Perivascular and adventitial mast cells     | Increased density in coronary spasm and thrombosis | Few mast cells in the arterial media |
| Intimal mast cells                          | Present in the human arterial intima, at the site of atherogenesis | Preferentially located in the shoulder region of atheromas |
|                                            | Increased density of activated mast cells in human coronary atheroma |
We have extended the seminal observation by Roberto Levi’s and Randi B. Silver’s groups [29, 30] by demonstrating that HHMC also contain renin [Marone G., unpublished information]. Thus, there is the possibility that renin released from activated cardiac mast cells can trigger local formation of angiotensin II.

Activation of HHMC with anti-IgE or anti-FcεRI induces the de novo synthesis of PGD₂ (≈ 18 ng/10⁶ cells) and LTC₄ (≈ 18 ng/10⁶ cells) [19, 20]. This is a relevant observation because both PGD₂ and LTC₄ have several cardiovascular effects [31, 32], meaning that the release of PGD₂ from HHMC can cause coronary vasoconstriction in man.

PAF is an important mediator of anaphylaxis produced by human mast cells [33]. There is preliminary evidence that immunologically activated HHMC synthesize PAF (Triggiani M., unpublished information). Importantly, serum PAF levels are directly correlated and PAF acetylhydrolase levels are inversely correlated with the severity of anaphylaxis [34]. Fig. 2 summarizes the principal receptors present on the surface of HHMC and the mediators released by their activation.

**Cytokines synthesized by HHMC**

Cytokines have been implicated in the pathogenesis of a variety of cardiovascular diseases [35]. There is growing evidence of the possible roles of cardiac mast cell-derived cytokines in anaphylaxis. Mast cells in human coronary atheromas contain TNF-α [36]. We have demonstrated the presence of granule-associated stem cell factor (SCF), the principal growth factor for human mast cells [37], in HHMC of patients with dilated cardiomyopathy [27]. Thus, there is the possibility that SCF released by HHMC is an autocrine factor that contributes to mast cell hyperplasia observed in dilated cardiomyopathy [27]. Additional studies are necessary to clarify the roles of other cytokines in anaphylaxis and the contribution of cardiac mast cells to their production.

**Stimuli that activate HHMC in vitro**

Immunologic activation of HHMC can be induced by IgE-mediated stimuli (antigens, immunoglobulin superantigens, anti-IgE or anti-FcεRI) [19, 20, 38, 39]. Activation of HHMC by anti-IgE and by a monoclonal antibody against an epitope of the α-chain of FcεRI may be clinically important. Histamine-releasing autoantibodies against IgE or against the α-subunit of FcεRI are present in the circulation of some patients with various allergic disorders [40, 41].

Anaphylatoxins (C3a and C5a) are formed when complement is activated during cardiac [42] and systemic anaphylaxis [9]. C5a causes cardiovascular derangement directly or through the release of vasoactive mediators [43, 44]. We demonstrated that C5a and C3a cause rapid histamine release from HHMC [19]. Interestingly, C3a causes cardiac dysfunction in the Langendorff-perfused guinea pig heart system [44]. SCF, which is present in the secretory granules of HHMC [27] and is released after their immunologic activation [45], activates HHMC through the engagement of kit receptor. Eosinophil granule proteins (ECP and MBP) are selective activators of HHMC [20, 46, 47]. The latter observation suggests that eosinophil-mast cell interactions might be clinically relevant in patients with hypereosinophilia.

A variety of non-immunological stimuli can activate HHMC. Some of clinical relevance because they might explain adverse effects observed in vivo when these compounds are used for diagnostic (contrast media, etc.) [3, 7] or therapeutic purposes (general anesthetics, protamine, etc.) [20]. For example, protamine and certain general anesthetics (propofol and atracurium) can cause histamine release from HHMC [20, 48]. Radio contrast media, injected into the coronary arteries for diagnostic purposes, can also activate HHMC in vitro [20]. The proximity of HHMC to coronary blood vessels [12, 13] and the presence of mast cells in human coronary atheromas [23] suggest that intracoronary injection of high doses of contrast media can activate mast cells and induce in vivo release of vasoactive mediators. In fact, in a multicenter study of patients who experienced immediate reactions to the injection of contrast media, the concentrations of plasma histamine and tryptase rose [49].

**HHMC in systemic and cardiac anaphylaxis**

Roberto Levi’s group has provided extensive evidence that the heart is involved in experimental anaphylaxis [29, 30, 43, 50] through the release of chemical mediators from cardiac mast cells. There is increasing evidence of cardiac involvement in human anaphylaxis too [8, 9, 12, 13]. Initially this has been attributed to mediators originating from the lung and reaching the heart. However, it is likely that the local release of vasoactive mediators by mast cells around coronary arteries and in human coronary atheromas, particularly in patients with ischemic heart disease [23], can contribute to cardiovascular derangements during anaphylaxis. In addition, C5a formation has been documented during anaphylaxis in man [9]. HHMC possess FcεRI, C3a receptor (C3aR), and C5a receptor (C5aR) bound to their membrane surface. Therefore, it is likely that IgE-, C5a-, and C3a-mediated activation of these cells could play a role in systemic and cardiac anaphylaxis in humans [9, 12, 13].

In addition, several agents injected intravenously (general anesthetics, protamine, radio contrast me-
Cardiovascular effects of histamine in man

Several years ago, in collaboration with our colleagues of the Division of Cardiology at the University of Naples Federico II, we have investigated the effects of histamine on peripheral and coronary hemodynamics in man. Low doses of histamine (0.4 µg/kg/min) infused in patients with normal left ventricular (LV) function undergoing diagnostic catheterisation induced significant drops in systolic, diastolic, and mean aortic pressure, systemic vascular resistance, LV end-diastolic pressure, and stroke index [51]. Hemodynamic changes started 1–2 min after the infusion started and reverted to normal within 5 min after the infusion. In one subject there was a transient progression from first to third degree atrioventricular block, with prompt recovery of atroioventricular conduction at the end of the infusion. Thus, exogenous histamine in man causes significant transient hemodynamic changes, mainly systemic hypotension, tachycardia, and increased LV performance. These changes can be partly attributed to the related increase in sympathoadrenergic activity, although it cannot be excluded that histamine has some direct cardiac effect.

In a subsequent study, the effects of selective activation of histamine H₁ receptors on coronary hemodynamics were examined in two groups: patients with atypical chest pain and normal coronary arteries and patients with vasospastic angina [52, 53]. H₁ receptor stimulation was achieved by infusing histamine intravenously for 5 min after pretreatment with cimetidine to antagonize the H₂ receptors. Heart rate was kept constant by coronary sinus pacing. In the first group mean aortic pressure and coronary vascular resistance (CVR) dropped, while coronary blood flow (CBF) and myocardial oxygen consumption remained unchanged during histamine infusion. No patient in this group developed angina during histamine infusion. By contrast, a percentage of the second group developed angina during histamine infusion (~40%), with a decrease in CBF and an increase in CVR. These findings suggest that stimulation of the H₁ receptor in subjects with normal coronary arteries reduces CVR, probably because of vasodilation of small coronary resistance vessels. However, in some patients with vasospastic angina, H₁ receptor activation can cause vasoconstriction of large-capacitance coronary arteries.

The results of these studies support the hypothesis that the endogenous release of histamine during anaphylactic reactions may precipitate coronary spasm in a subset of patients with vasospastic angina. Interestingly, it has been reported that premedication with an H₁ receptor antagonist increases the risk of heart block in patients who develop anaphylaxis [54].

Studies have now started to clarify the role of H₁ receptors in the cardiovascular system. Roberto Levi and collaborators have identified H₁ receptors as inhibitory heteroreceptors in cardiac adrenergic nerve endings [55, 56] suggesting a mechanism by which endogenous histamine can activate norepinephrine release in normal and ischemic conditions [57]. The presence of H₁ receptors in human heart [55] suggests that these receptors might be directly and/or indirectly involved in anaphylactic reactions.

Cardiovascular effects of cysteinyl leukotrienes in man

In collaboration with cardiologists we have evaluated the effects of cysteinyl leukotriene D₄ (LTD₄) on systemic and coronary hemodynamic in patients with normal coronary arteries [17]. LTD₄ induced an early (20 s) transient fall in mean arterial pressure paralleled by rises in heart rate and plasma levels of catecholamines which returned to baseline after 10 min. CVR rose at 10 and 15 min. Thus, small doses of cysteinyl leukotrienes may induce both an early transient fall in mean arterial pressure, with secondary sympatho-adrenergic activation, and a
later increase in small coronary arteriolar resistance. Although high levels of cysteinyl leukotriene receptor CysLT₁ mRNA were detected in the human heart and coronary artery, while cysteinyl leukotriene receptor CysLT₂ mRNA was barely detectable [58], it is not clear which receptor was involved in these cardiovascular effects.

**Conclusion**

Within human cardiac tissue, mast cells are located between myocytes and are in close contact with blood vessels [12, 13, 19, 20, 22, 45]. They are also found in the coronary adventitia and in the shoulder regions of coronary atheroma [22, 23, 24].

Circulating antigens, superantigens, autoantibodies (anti-IgE, anti-FceRI, etc.), eosinophil granule proteins (major basic protein, MBP, and eosinophil cationic protein ECP), therapeutic (e.g. protamine, general anesthetics) or diagnostic substances (e.g. radio-contrast media) can induce the release of mediators from HHMC. These cells express FceRI, C5aR, and C3aR, which could explain the involvement of cardiac mast cells in systemic and cardiac anaphylaxis.

The increases in cardiac mast cell density and release of vasoactive mediators in patients with cardiomyopathy might also have clinical implications given the marked cardiovascular effects of histamine, cysteinyl leukotrienes, and PGD₂ [31, 32, 51, 52, 53].

Pioneering studies have started to shed light on the cardiovascular effects in humans caused by such mast cell-derived mediators as histamine [51, 52, 53] and leukotrienes [12]. Interestingly, these studies provided the important information that the hemodynamic effects of mediators depend on both the underlying cardiovascular conditions and pharmacologic treatment (e.g., H₂ blockade). Using specific antagonists of CysLT₁ and CysLT₂ [58] it will be possible to assess the each receptor’s contribution to the cardiovascular effects of these vasoactive mediators. Immunologically activated human mast cells synthesize PAF [33], which can play a role in human [34] and experimental anaphylaxis [59].

In conclusion, there is extensive experimental and clinical evidence that the human heart can be viewed as both a site and a target in anaphylaxis. Pharmacological targeting of cardiac mast cells might offer novel therapeutic opportunities to prevent and/or modulate the dramatic and potentially fatal manifestations of anaphylaxis.
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