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**Peripheral reflex feedbacks in chronic heart failure: Is it time for a direct treatment?**

Giannoni A *et al.* Reflex feedbacks in heart failure

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**Abstract**

Despite repeated attempts to develop a unifying hypothesis that explains the clinical syndrome of heart failure (HF), no single conceptual paradigm for HF has withstood the test of time. The last model that have been developed, the neurohormonal model, had the great virtue of highlighting the role of the heart as an endocrine organ, as well as to shed some light on the key role on HF progression of neurohormones and peripheral organs and tissues beyond the heart itself. However, while survival in clinical trials based on neurohormonal antagonist drugs has improved, HF currently remains a lethal condition. At the borders of the neurohormonal model of HF, a partially unexplored path trough the maze of HF pathophysiology is represented by the feedback systems. There are several evidences, from both animal studies and humans reports, that the deregulation of baro-, ergo- and chemo-reflexes in HF patients elicits autonomic imbalance associated with parasympathetic withdrawal and increased adrenergic drive to the heart, thus fundamentally contributing to the evolution of the disease. Hence, on top of guideline-recommended medical therapy mainly based on neurohormonal antagonisms, all visceral feedbacks have been recently considered in HF patients as additional potential therapeutic targets.

**Key words**: Baroreflex; Chemoreflex; Ergoreflex; Heart failure; Sympathetic system; Neurohormones

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**Core tip:** At the borders of the neurohormonal model of heart failure (HF), a partially unexplored path trough the maze of HF pathophysiology is represented by the feedback systems. There are several evidences, from both animal studies and humans reports, that the deregulation of baro-, ergo- and chemo-reflexes in HF patients elicits autonomic imbalance associated with parasympathetic withdrawal and increased adrenergic drive to the heart, thus fundamentally contributing to the evolution of the disease. Hence, on top of guideline-recommended medical therapy mainly based on neurohormonal antagonisms, all visceral feedbacks have been recently considered in HF patients as additional potential therapeutic targets.

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Heart failure (HF), a pathophysiological state in which the heart is unable to pump blood at a rate commensurate with the metabolizing tissues requirements, or can do so only with elevated filling pressures[1], is currently a real epidemic in western countries, affecting more than 20 million people in the world, with massive socio-sanitary costs[2].

Despite repeated attempts to develop a unifying hypothesis that explains the clinical syndrome of HF, no single conceptual paradigm for HF has withstood the test of time. The last model that have been developed, after the cardiorenal and the cardiocirculatory models focusing respectively on salt-water retention and low cardiac output/peripheral vasoconstriction, is the neurohormonal model[3]. This model had the great virtue of highlighting the role of the heart as an endocrine organ, as well as to shed some light on the key role on HF progression of neurohormones and peripheral organs and tissues beyond the heart itself. However, while survival in clinical trials based on neurohormonal antagonist drugs has improved, HF currently remains a lethal condition, with 50% mortality within 5 years of diagnosis and less than 15% survival after 10 years[2,4].

At the borders of the neurohormonal model, a partially unexplored path trough the maze of HF pathophysiology is represented by the feedback systems (Figure 1). There are indeed several evidences, from both animal studies and humans reports, that the deregulation of baro-, ergo- and chemo-reflexes in HF patients elicits autonomic imbalance associated with parasympathetic withdrawal and increased adrenergic drive to the heart, thus fundamentally contributing to worsening arrhythmias and haemodynamics. Hence, on top of guideline-recommended medical therapy mainly based on neurohormonal antagonisms, all visceral feedbacks have been recently considered in HF patients as additional therapeutic targets: baroreflex activation therapy for baroreceptors[5], physical training for muscle metaboreceptors[6],and carotid body (CB) denervation for chemoreceptors[7].

**BAROREFLEX**

The baroceptors are mechanoreceptors located in the sinus caroticus and in the aortic arch, where terminal nerve endings are endowed in the wall of these vessels and activated by blood pressure-induced wall stretch. Information deriving from these sites travel along a path constituted by the nerve of hering, that merges with the fibres of the glossopharingeal nerve; those travelling from the aortic arch take the path of the afferent fibres of the *vagus* nerve. Inputs hence travel towards the principal centre of integration of information regarding the cardiovascular system, that is the nucleus tractus solitarii (NTS) in the dorsal area of its medial and lateral divisions. Here signal are processed and integrated with information ascending from the periphery and descending from central nervous system and given back to the heart and peripheral arterial vessels *via* the vagus nerve[8]. The response is a vagally-mediated change in heart rate and a sympathetic modulation of vasomotion, in order to preserve blood pressure stability over time and avoid fluctuations[9].Altered baroreflex sensitivity (BRS) has been demonstrated to independently contribute to worsen prognosis in HF, mainly by failing to counteract the adrenergic activation with consequent electrical instability and arrhythmic sequelae, in both the pre- and post-betablocker era[10,11].

The baroreflex has been the first neurovegetative feedback to be clinically targeted in HF. BRS activation was first indirectly attempted by vagal nerve stimulation (VNS). After the first safety and tolerability reports on VNS (side effects: hoarseness, cough and sensation of electrical stimulation) some preliminary studies also showed amelioration of symptoms and indexes of LV remodelling[12,13]. These observations led to a phase III sham-controlled trial. The neural cardiac therapy for HF (NECTAR-HF) trial enrolled 87 patients with systolic HF (LVEF < 35%) who underwent device implantation and randomization to device in ON or OFF modality, but failed to demonstrate any effect of VNS on both primary (LV end systolic diameter) and secondary endpoints (LV end systolic volumes, LV ejection fraction, oxygen consumption and natriuretic peptide levels)[14].

Baroreceptor stimulation could also be reached by directly stimulating carotid sinus by subcutaneously implanted device: this approach is known as baroreflex activation therapy (BAT). The first promising results obtained in an animal model of HF (dog with HF induced by microembolization) in terms of reverse remodelling, improved systolic function and amelioration of neurohormonal profile (reduced adrenergic activity), where secondarily confirmed also in a proof of concept study performed in humans, where an amelioration of symptoms was also observed[15]. Few on-going randomized studies are currently addressing the efficacy and therapeutic potential of baroreflex activation therapy in HF; in particular, the CVRx® Rheos® Diastolic Heart Failure Trial (clinicaltrial.org: NCT00718939) and the Rheos® HOPE4HF Trial (NCT00957073) will address the impact of BAT on diastolic HF (LVEF > 40%), whereas in systolic HF patients, the only ongoing randomized trial is the Barostim HOPE4HF (Hope for Heart Failure) study (NCT01720160).

**CHEMOREFLEX**

The chemoreflex is physiologically in charge of proportionally modulating ventilation in response to a change in the respiratory gases, namely oxygen (O2) and carbon dioxide (CO2), in order to keep pH constant for enzymatic processes. Classical physiology indicates two separate chemoreceptor groups: peripheral chemoreceptors (PC) located in carotid-aortic bodies and sensitive both to hypoxia and hypercapnia/acidosis, and central chemoreceptors (CC) located in different regions of the brainstem, cerebellum, hypothalamus and glia and considered to be sensitive only to hypercapnia/acidosis.

Chemoreceptors seem to act as primary inputs in HF. Several studies indicate that both PC and CC are hyperactive in HF[16-19].The increased activity of chemoreceptors is commonly considered the main determinants of Cheyne-Stokes respiration[16-19], a detrimental respiratory pattern (with prognostic significance) characterized by alternating cycles of hyperventilation and apneas, with unfavourable oxygen desaturation. Furthermore, PC/CC hypersensitivity also negatively impact on respiration kinetics during exercise with ventilatory inefficiency and dyspnoea on effort in HF patients[18,19].The hyperactivity of PC/CC, both directly (baseline tonic activity and phasic stimulation during O2/CO2 changes)[20] and indirectly, *via* Cheyne-Stokes respiration (CSR) occurrence[21]. Alongside baroreflex deactivation, chemoreflex activation is also responsible of increased adrenergic drive and arrhythmias in HF patients[17-19]. Finally, increased chemosensitivity to both hypoxia[16] and hypercapnia[19] was found to be an independent prognostic marker in HF.

A partial inhibitory effect on PC was shown in HF patients with both transient hyperoxia, and drugs, such as dihydrocodeine or acetazolamide. In HF patients, dihydrocodeine mediated PC inhibition was only associated with improved exercise performance[22].In the same setting, acetazolamide[23] and hyperoxia[24] were instead associated with about 50% reduction of CSR severity, translating in the case of hyperoxia also with reduced sympathetic activity. Denervation of the PC chemoreceptors by carotid bodies (CB) ablation in animals with experimentally induced HF has recently emerged as a very promising option. CB ablation is indeed able to normalize the chemoreflex sensitivity in HF animals, with reduction of both adrenergic activity and disappearance of central apneas[7,25].This was confirmed also by pharmacologic attenuation of CB activity with an inhibitor of hydrogen sulfide[26].Interestingly, in a model of HF induced by coronary ligation in rats, CB also reduced the amount of myocardial fibrosis unrelated to myocardial infarction, with positive effect on left ventricular systolic function and, more importantly on short term survival[25].A single report in a patient with HF has testified the feasibility in humans[27]. Differently from these still preliminary, but intriguing results on PC modulation, currently no studies have tested the possibility to directly act on CC, maybe due to the multiplicity of CC centers in the central nervous system, the complexity of their interlink, and the difficulty to directly and selectively act on these receptors.

**ERGOREFLEX**

The ergoreflex is the neural mechanism enabling to modulate ventilation and sympathetic outflow according to the intensity of physical activity[28]. Its components are the metaboreflex, activated by the accumulation of metabolites in the exercising muscles, and the mechanoreflex, responsive to muscle tension during exercise[29-31].

HF patients frequently develop a skeletal myopathy ascribable to deconditioning, reduced perfusion of the muscles, inflammation, and a systemic catabolic state[29,30,32].In 1994, a “muscle hypothesis” of HF was formulated, suggesting that ergoreceptor contribution to the autonomic, hemodynamic, and respiratory responses to exercise would be enhanced in CHF patients[33]. Two years later, ergoreflex overactivity was first found in HF patients compared with healthy subjects[6]. These results were corroborated by subsequent studies, which correlated increased ergoreceptor sensitivity to lower lean body mass, reduced exercise tolerance, decreased left ventricular function, and worse New York Heart Association functional class[30].Interestingly, in HF patients with preserved exercise capacity, ergoreflex overactivity has been also associated with increased central and peripheral chemoreceptor sensitivity, and depressed baroreceptor sensitivity[30].

Currently the only acknowledged treatment for modulating ergoreflex overactivity is represented by exercise training. The effects of training on ergoreflex sensitivity have been evaluated mostly in animal models[34]. In humans, six weeks of forearm training were able to markedly reduce metaboreceptor sensitivity, while six weeks of detraining brought the situation back to baseline[29].A positive effect on muscle structure and function has been after confirmed in other studies, still in HF patients[35,36].It is reasonable to assume that the positive impact of exercise training on HF patients (in terms of increased exercise tolerance, quality of life, cardiac function, neuro-hormonal activation and overall prognosis)[37-40] partially relies upon reduced ergoreflex overactivity, as confirmed by a recent study[41].

**CONCLUSION**

The lessons learned from failures (*e.g*., inotropic drugs) and the successes (*e.g*., neurohormonal antagonist drugs) in treating HF indicate that the development of innovative treatments for HF should to take into account the complex pathophysiology of the disease: in particular, new treatments should target the pathways involved in the evolution of the disease. As outlined above, peripheral reflexes are deeply involved in the pathophysiology of HF and represent a potential target of therapy. Despite, some preliminary data in animals and humans are promising, more studies enrolling a large number of patients are clearly needed to reinforce the rationale of treating the peripheral reflex feedbacks and to disclose the prognostic value of these interventions.

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**Figure 1 Schematic representation of the reflex feedbacks involved in heart failure. Arrows indicate direct effects/influences.** Dotted arrows link established or potential therapeutic interventions with targets. CNS: Central nervoius system; O2: Oxygen; CO2: Carbon dioxide; RIC: Remote ischemic conditioning; CSR: Cheyne-Stokes respiration; NIMV: Noninvasive mechanical ventilation; LV: Left ventricular; HF: Heart failure; BAT: Baroreceptor activation therapy.