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The ATP synthase glycine zipper of the *c* subunits: from the structural to the functional role in mitochondrial biology of cardiovascular diseases.

The holo-ATP synthase has a membrane rotor engine embedded in the inner mitochondrial membrane (IMM) formed by the c-ring structure in the F₀ hydrophobic domain [1]. Packing several copies of the c subunits provides a palisade of hairpin-shaped lpha-helices. The stoichiometry of the c-ring is specie-dependent and leads to a different c-ring size. Each hydrophobic hairpin shaped c subunit constitutes the basic unit of the ring: the N- and C-terminal α-helices of each subunit flank the same terminal regions of another subunit. This assembly forms two concentric circles with the N- and C- terminal α-helices that form the tight and wide transmembrane α -helix packing of the c-ring, respectively (Fig. 1A). In eukaryotes, these two circles in each c subunit are linked by a loop region which faces the headgroup region of phospholipids on the matrix side of the IMM and forms the contact site of the feat of the central stalk (γ , δ , and ϵ subunits). The hole of the cring is occupied with phospholipids [2]. In detail, the recent cryo-EM structure of the entire mammalian ATP synthase [3] reports lipids into the central hole from both sides of the cylinder. In the c-ring at the positive side of IMM a lipid plug, probably a lyso-phosphatidylserine, is linked by the terminal lysine of e subunit, whereas at the matrix side the Arg³⁸ of the c subunit on the N-terminal α -helix coordinates electrostatically the phosphatidylserine. The two phospholipids are separated in the middle of the c-ring by Val¹⁶ of c subunits (Fig. 1B). The single acyl-chain of lyso-phosphatidylserine inside the c-ring hole has enough space around it. Conversely, the double acyl-chains of phosphatidylserine are a tight fit to the c subunits. This strongly suggests that the lyso-phosphatidylserine does not rotate with the c-ring, acting as a "lubricated" lipid plug on the positive side of iMM, while the matrix-side phosphatidylserine does [3]. It has been established that the glycine zipper domain of the c subunit (Fig. 1B) bundles the lipid acyl-chains into the cavity of the rotor, apart from playing a specific influence on the c-ring stoichiometry that fulfils the mechanism of biological energy conversion of each individual specie [4]. Moreover, the repetitive glycine residues ($^{20}GxGxGxG^{20}$) are important for the inner concentric circle of the c-ring that, by the c/c-subunit interface; contributes to the cring stability [5]. The degree of structural compliance in the assembly of the rotor and geometry relies on i) the hydrophobic and weak H-bonds mix interactions arise from glycine zipper and ii) limited steric hindrance of Gly side chain (hydrogen atom). Probably, for this feature, the ²⁰GxGxGxG²⁶ motif can play a central role in the permeability transition pore complex (PTPC) opening [6].

The activation of ATP synthase by an abrupt increase of Ca^{2+} in mitochondria can trigger the PTPC phenomenon [7]. The new insights on the ATP synthase functions reveal at the molecular level the biological feature of the enzyme complex PTPC formation [8] as hypothesized by the lipid-plug out c-ring release in the "bent-pull" model [9]. Accordingly, the PTPC opens when the ATP synthase assumes the disassembled conformation in the presence of Ca^{2+} bound to the catalytic sites of F_1 , the hydrophilic domain of the enzyme. In this state, the peripheral stalk that joins the F_1 to F_0 is twisted with a consistent structural difference compared to the conformation in the presence of the natural cofactor Mg^{2+} . The structural change is transduced in signal propagation along with the entire enzyme from the catalytic site to the e subunit. Lastly, the lyso-phosphatidylserine is expelled from the central hole inside the e-ring and opens the channel at the positive side of IMM. The other side of the e-ring is filled with phosphatidylserine that can block ion conductance until the lipid is pushed apart by the hole. Simultaneously, water molecules fill the e-ring destabilizing the phosphatidylserine on the matrix side and the e-ring and the PTPC opening [3].

In physiological conditions, the opening of PTPC is involved in the cardiac and neural development. The relevance of a modulation of PTPC activity in this context is supported by evidence that depletion of the main activator of the PTPC, cyclophilin D, causes cardiac myocyte differentiation to begin earlier [9].

In pathological conditions, it has been well assessed that PTPC opening behaves as a key driver during myocardial ischemia/reperfusion (I/R) injury [10]. During ischemia cardiomyocytes are exposed to Ca²⁺ overload, increased oxidative stress and inflammation. These factors are known to favour the susceptibility of the PTPC to opening. In addition, once reperfusion occurs, it restores a normal pH which induces increased PTPC activity with consequent severe mitochondrial impairment and disruption of energy production, leading to a massive cellular damage [11]. Accordingly, acidic pH reduces cell death in I/R, and this has been attributed to inhibition of the PTPC opening by acid pH. Of importance, there is evidence that inhibitors of the PTPC, *i.e.* cyclosporin A, may reduce infarct size following I/R in humans [12] although a robust demonstration of this phenomenon is still lacking.

The link between inhibition of the PTPC opening and cardioprotection has been demonstrated in cells, isolated hearts, and *in-vivo* models [10]. Apart from the I/R and the myocardial infarct conditions, a role of PTPC opening has been hypothesized, but not yet definitively demonstrated, in heart failure [10].

Recent data gained on the glycine-rich region of *c* subunit provide useful information for a better understanding of the molecular mechanism that leads to the PTPC phenomenon and of its involvement in cardiovascular diseases. Indeed, glycine zipper of *c* subunits can be a critical factor in exacerbating PTPC opening under pathological conditions [13]. Gly residue is the only amino acid without a side chain. The substitution of Gly with other amino acids necessarily produces a steric hindrance and much more interactions than the H atom alone. Increased channel conductance of mutated glycine zipper suggests decreased packing and a large central pore formation [14]. Different substitution of Gly²⁶ would exhibit a structural deformation at the level of the N-terminal helix that differently modulates the PTPC opening [6]. Moreover, Gly²⁶ mutation is more sensitive to form PTPC than mutation on Gly²⁴. Gly²², and Gly²⁰ and more than wild type *c* subunit. In particular, the Gly²⁶Glu moration within the glycine-rich region of *c* subunit causes the alteration of the side chain length and the addition of negatively charged residue that could favour the instability of phosphatidylserine in the *c*-ring. Therefore, when the ATP synthase triggers the PTPC formation, the mutant *c*-subunit may prefer a hydrophilic hole into caring for the passage of water.

In this context, the pathological role of the naturally occurring mutation ATP5G1^{G87E} within the c subunit has been recently unraveled by the elegant work of Morciano $et\ al$. [13]. The authors of the study showed that carriers of this mutation suffered from a greater myocardial damage at the time of reperfusion following ST-elevation myocardial infarction (STEMI). A significant inverse correlation was detected between PTPC activity and reperfusion damage in this condition, in particular, $Gly^{26}Glu$ substitution exhibited hyperresponsive PTPC opening compared with wild type c subunit by aggravating the PTPC-mediated hypoxia/reoxygenation cell death in human cardiomyocytes [13]. These effects are consistent with the abovementioned notion that PTPC remains closed during myocardial ischemia and re-opens at the time of reperfusion. The human evidence is corroborated by the consequences of genetic manipulations of c subunit expression levels by siRNA in HeLa cells. Herein, the *in-vitro* data confirmed that c subunit is required for PTPC driven mitochondrial fragmentation and cell death triggered by cytosolic Ca^{2+} overload and oxidative stress [14,15].

The evidence so far gained on the pathogenic implications of c subunit in ischemic heart disease raises another interesting issue, that is its potential therapeutic implications to improve treatment of this common disease and to reduce its dramatic consequences. In fact, several efforts have been already made in the attempt to reduce the tissue damage at the time of reperfusion, but none turned out successful. The discovery that the ATP5G1^{G87E} mutation, within the highly conserved glycine-rich domain of the c subunit, amplifies the reperfusion injury by further enhancing PTPC opening reinforces the hypothesis that a timely intervention with inhibitors of PTPC activity may serve as a valid therapeutic approach to reduce the hypoxia/reoxygenation damage at the time of revascularization of an occluded coronary artery. In particular, the strategy based on the selective targeting of c subunit, to reduce PTPC opening, may reveal the most suitable approach during coronary angioplasty following myocardial infarction. Moreover, by reducing the

myocardial damage dependent from coronary occlusion and particularly from the subsequent revascularization, a decrease of the cardiac remodelling process and a reduced progression of ischemic cardiomyopathy toward heart failure development would be expected.

The emerging link of the ATP synthase *c* subunit with cardiovascular diseases deserves to be fully dissected out through future experimental and human studies.

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Eigure caption

Figure 1. Structure representation of c-ring. A) Cryo-EM of c-ring ATP synthase, 8 c subunits, PDB ID codes: 6TT7. Lipid-plug int the c-ring viewed from the positive side of IMM (left panel) and from matrix side (right panel). B) Cross section of the c-ring. Phosphatidylserine (PS), as negatively charged lipid, at the matrix side is coordinated by the positive charge of Arg³⁸ residue of c subunits. Lyso-phospatidylserine (LPS) at the positive side of IMM is bound with its polar head to the Lys⁷¹ of the e subunit (not shown). Hole lipid chambers are separated by the annular of Val¹⁶ residues. In the boxes the structure of both wild-type (left panel) and mutant (right panel) glycine zipper are shown. PS and LPS are drawn as stick models, whereas amino acid residues are drawn as ball-and-stick.

