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Proton leak through the UCPs and ANT carriers and beyond: A breath for the electron transport chain

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8 and 2010 **120 and 2010 120 and 2010 120 and 2010 120 and 2010 120 and 2010** 9 Correspondence: <u>salavatore.nesci@unibo.it</u>

# <sup>12</sup> Abstract

 $\frac{13}{14}$  Mitochondria produce heat as a result of an ineffective H<sup>+</sup> cycling of mitochondria respiration across the inner mitochondrial membrane (IMM). This event present in all mitochondria, known as proton leak, can decrease 15 16 protonmotive force  $(\Delta p)$  and restore mitochondrial respiration by partially uncoupling the substrate oxidation  $^{17}$  from the ADP phosphorylation. During impaired conditions of ATP generation with F<sub>1</sub>F<sub>O</sub>-ATPase, the  $\Delta p$  $\frac{18}{10}$  increases and IMM is hyperpolarized. In this bioenergetic state, the respiratory complexes support H<sup>+</sup> transport  $\frac{19}{20}$  until the membrane potential stops the H<sup>+</sup> pump activity. Consequently, the electron transfer is stalled and the 21 reduced form of electron carriers of the respiratory chain can generate  $O_2$  triggering the cascade of ROS 22 formation and oxidative stress. The physiological function to attenuate the production of  $O_2$  by  $\Delta p$  dissipation  $\frac{23}{24}$  can be attributed to the proton leak supported by the translocases of IMM. 14 Mitochondria produce heat as a result 20 until the membrane potential stops the 24 CM commence to the pretention of

### 27  $\boldsymbol{V}$  and  $\boldsymbol{V}$  $28$  Reywords

29 proton leak; proton-motive force; electron transport chain; reactive oxygen species; proton conductance; 30 mitochondria<br>31 31

### 57  $\lambda$ 58 **ADDreviations**

59  $\Delta p$ , proton-motive force;  $\Delta \psi$ , membrane potential; ETC, electron transport chain; O<sub>2</sub>, superoxide anion; 60 ROS, reactive oxygen species; IMM, inner mitochondrial membrane;  $J_{H^+}$ , proton flux;  $G_{H^+}$ , proton <sup>61</sup> conductance of the inner mitochondrial membrane; ANT, adenine nucleotide translocase; UCP, uncoupling protein; PUFA, polyunsaturated fatty acids; BAT, brown adipose tissue. 62 63 Protein, Port, polyunsulatured rates

### 1. Introduction

The mitochondria in eukaryotic organisms accomplish their function by creating energy in the form of ATP, 2 <sup>3</sup> the universal biological energy currency while consuming oxygen. In mitochondria, the reducing equivalents  $\frac{4}{5}$  are transferred, during the tricarboxylic acid cycle, pyruvate oxidation, fatty acid and amino acid catabolism to the cofactors  $NAD^+$  and  $FAD^+$ . The reduced cofactors, *i.e.*, NADH and  $FADH_2$ , are oxidized with mitochondrial respiration and electrons are transferred through the respiratory chain to the final acceptor, O<sub>2</sub>.<br>
8 The movement of electrons through the carriers in the electron transport chain (ETC) is led by a reducti potential that increases gradually creating a negative  $\Delta G^{\circ}$  exploited by the respiratory complexes to transfer  $\frac{10}{11}$  H<sup>+</sup> from the matrix (negative side) to the space existing between the inner and outer mitochondrial membranes  $\frac{11}{12}$  (positive side) [1,2]. The catalytic activity of H<sup>+</sup> pumps of the ETC (complex I, complex III and complex IV) 13 forms an electrochemical gradient of H<sup>+</sup> ( $\Delta \mu$ <sub>H+</sub>), which in terms of proton-motive force (pmf or  $\Delta p$ ) is equal in 14 voltage units at  $-\Delta\mu_H+|F|$  (with F the Faraday constant). The  $\Delta p$  in mammalian mitochondria consists mostly of <sup>16</sup> pH difference,  $\Delta p$ H) [3,4]. The  $\Delta p$  is the main energy source in mitochondria to guide the re-entry of H<sup>+</sup> into the matrix for the synthesis of ATP through the  $F_1F_0$ -ATPase and to maintain the ionic homeostasis of the matrix for the synthesis of ATP through the  $F_1F_0$ -ATPase and to maintain the ionic homeostasis of metabolites supporting the activity of the inner mitochondrial membrane (IMM) carrier proteins. In the 19 chemiosmotic system, the energy released by oxidation reactions of substrates is coupled to mitochondrial 20  $^{21}$  ATP synthesis in a biological process known as oxidative phosphorylation (OXPHOS) [5] (Fig. 1).  $6$  to the collactors NAD and  $FAD$ . mitochondrial respiration and electrons are transferred through the respiratory chain to the final acceptor, O<sub>2</sub>.<br><sup>8</sup> The movement of electrons through the carriers in the electron transport chain (ETC) is led by a reduct  $12$  (positive side) [1,2]. The catalytic act <sup>15</sup> an electrical gradient (transmembrane potential,  $\Delta \psi$ ) and a small part of a chemical gradient (transmembrane 18 line maintain tor the synthesis of ATT 22 **100 PM Program Program Program Program** Program Pr

23 The OXPHOS mechanism can be depicted as a circuit in which the proton flux  $(J<sub>H<sup>+</sup></sub>)$  is comparable to the  $\frac{24}{25}$  current flowing in an electrical circuit powered by the respiratory complexes that constitute the electrical 26 "chargers". The  $F_1F_0$ -ATPase serves as an engine that p  $27$  whereas the circuit resistance includes every step of H<sup>+</sup> translocation through the bilayer of IMM [6]. In an <sup>28</sup> "ideal" situation the  $\Delta p$  is coupled entirely to mitochondrial ATP synthesis. However, the biology of 30 mitochondria provides that  $\Delta p$  retains a physiological mild uncoupling of OXPHOS. This is handled with the  $H^+$  conductance through the IMM ( $G_{H^+}$ ) that sustains the  $\Delta p$  dissipation responsible for driving the electrons  $\frac{32}{33}$  transport and respiration by the ETC in the absence of ATP synthesis [7–9] (Fig. 1). 25 Carlott howing in an electrical circle  $29$  areal situation the  $\Delta p$  is coupled 33 real of the responsibility the ETC

 $\frac{34}{35}$  All mitochondria possess a proton leak through the IMM whose identity and function are still not fully 36 understood [10–13]. Proton leak is a particular type of thermogenic process. The key physiological activities  $\frac{37}{20}$  of other various electrogenic conductances of the IMM involved in mitochondrial uncoupling include the control of mitochondrial-regulated cell death by the permeability transition pore (PTP) phenomenon [14]. The 40 endogenous proton leak is highly dependent on the  $\Delta p$  decreasing in state III (active respiration). On the <sup>41</sup> contrary,  $\Delta p$  increases in state IV (respiratory status of rest or controlled) when the mitochondria supplied with substrates convert all added ADP to ATP [15,16]. Therefore, the effective  $G_{H^+}$  may sustain mitochondrial substrates convert all added ADP to ATP [15,16]. Therefore, the effective  $G_{H^+}$  may sustain mitochondrial 14 respiration in state IV acting as an escape route for H<sup>+</sup> when the  $\Delta p$  increases [10,12]. In state IV, with ETC <sup>45</sup> overloaded by stalled electrons, would increase the risk of superoxide anion  $(O_2)$  formation in Complexes I  $\frac{46}{47}$  and III [12]. Moreover, other powerful 48 hydrogen peroxide (H<sub>2</sub>O<sub>2</sub>), and peroxynitrite (ONOO<sup>-</sup>) can be obtained from  $O_2$ <sup>--</sup>[17]. 35 All mnochonuma possess a proton 39 control of mitochondrial-regulated ce 43 SUDSTRIES CONVERT ALL ACCED ADP TO  $47$  and  $III$  [12]. Moreover, other powerfu

The production of ROS in mitochondria depends entirely on the state of coupling of OXPHOS. Conditions 50  $\frac{51}{22}$  that lead to obtaining a low rate of electron transfer [18] can increase ROS generation since a prolonged Formal reduced state of the respiratory carriers causes the electron leak [19]. The mild uncoupling of OXPHOS, 54 decreasing the value of the  $\Delta p$  as a consequence of raising the  $G_{H^+}$ , stimulates respiration and might reduce the  $\frac{55}{25}$  formation of ROS [20,21]. 53 reduced state of the respiratory carr 56 **Common of ROS**  $[20,21]$ 

### 2. Proton leak: fact or artefact

The proton leak causes an uncoupling between the oxidation of the substrate and the phosphorylation of ADP 2 <sup>3</sup> by allowing H<sup>+</sup> of  $\Delta p$  to return to the matrix. Proton leak in mitochondria is demonstrated with  $\Delta p$  increase in the presence of the  $F_1F_0$ -ATPase inhibitor oligomycin [13]. Moreover, mitochondrial translocation of monovalent cations and anions by simple diffusion in the absence of energized IMM is the factor that affects 6  $\frac{7}{6}$  the passive osmotic swelling of mitochondria driven by a pH-dependent mechanism of ionophores (Fig. 2). Conversely, the phenomenon is counteracted by proton leak events  $[22-24]$ . As already deduced by  $\frac{8}{9}$ 10 Krishnamoorthy and Hinkle [18], the H<sup>+</sup> or OH<sup>-</sup> flux through the IMM shows non-linear progress as  $\Delta \psi$  varies, <sup>11</sup> whereas H<sup>+</sup> or OH<sup>-</sup> flow across the membranes is controlled by pH following Fick's Law [25]. Therefore, proton leak is directly proportional to  $\Delta pH$  [26]. Otherwise, this non-specific membrane permeability to H<sup>+</sup> at  $\Delta pH$  [26]. 14 high values of  $\Delta p$  violates Ohm's Law. Indeed, the non-ohmic proton leak is revealed by an exponential  $\frac{15}{16}$  behaviour of the current-to-voltage (I/V) ratio (Fig. 3) [6].  $5$  line presence of the  $\mathbf{r}_1 \mathbf{r}_0$ -Alpase in 9 Conversely, the phenomenon is cou 13 **proton leak is directly proportional to** 16 **Condition** of the early to voltage (1)

 $\frac{17}{10}$  The nonlinear curve, shown in Figure 3, may suggest that the permeability coefficient increases with the 19 membrane voltage,  $\Delta \psi$ , denoting a non-constant resistance for the H<sup>+</sup> flux. The relative permeability to H<sup>+</sup> of IMM is a function of the electrical component of  $\Delta p$ .<br>
21 An alternative interpretation to explain the H<sup>+</sup> permeability through IMM (as a function of the  $\Delta p$ ) was 18 The nominear carve, shown in Figure 20 IMM is a function of the electrical component of  $\Delta p$ .<br>21 21

 $\frac{23}{24}$  proposed by Pietrobon [27] suggesting the ph 25 mitochondrial H<sup>+</sup> pumps in which the H<sup>+</sup> permeability coefficient remains constant, but there is a decrease in <sup>26</sup> the stoichiometric ratio H<sup>+</sup>/e<sup>-</sup> in ETC at high values of  $\Delta p$  [27–29]. Reducing a half-molecule of oxygen, the <sup>27</sup> cytochrome c oxidase (Complex IV) couples to each electron transferred to the binuclear haem a<sub>3</sub>/Cu<sub>B</sub> the 29 pumping of H<sup>+</sup> from the negative side to the positive side of the IMM. The stoichiometry becomes 1:1 for the <sup>30</sup> ratio H<sup>+</sup>/e<sup>-</sup>. In addition, a "chemical" transport of H<sup>+</sup> by Complex IV from the matrix to *intracristae* space is  $\frac{31}{32}$  responsible for the  $\Delta \psi$  generation sustained by H<sub>2</sub>O production [30,31]. During the intrinsic uncoupling in 33 Complex IV, the electrons can reduce oxygen to  $H_2O$  bypassing the haem "a" and decoupling  $H^+$  pump to the positive side of the IMM [28]. In this situation, the ratio  $H^+ / e^-$  decreases since  $H^+$  flow is not driven by the  $\frac{35}{36}$  redox potential of the respiratory chain [29,32]. The slip of OXPHOS was observed only in Complex IV but not in Complex I or Complex III. A fundamental possibility of slip is also assumed for Complex III decoupled 37  $\frac{38}{2}$  by DCCD [33] or by cyclic transport of H<sup>+</sup> within the complex sustained with the "rescue pathways" of  $\alpha$ , $\omega$ - $\frac{39}{40}$  dioic acids to protect against ROS generated in mitochondria [34].  $24$  proposed by Fieldboll [27] suggest 28 Cytochronic c oxidase (Complex TV)  $32$  responsible for the  $\Delta \psi$  generation st 36 redox potential of the respiratory cha 40 along actus to protect against ROS ge.

<sup>41</sup> The existence of proton leak events in mitochondria supports different functions: *i*) thermogenesis, a way for 43 the avoidance of dielectric breakdown of IMM at excessive  $\Delta p$ ; *ii*) the improved capacity to regulate the <sup>44</sup> oxidative energy metabolism; *iii*) the ability to continue mitochondrial catabolism when the cellular ATP  $\frac{45}{46}$  demand is low maintaining a high ratio of NAD<sup>+</sup>/NADH suitable for the catabolism; *iv*) regulation of body mass; v) attenuation of ROS production [7,35]. 47 42 The existence of proton leak events in 46 demand is low maintaining a night ra

The proton leak occurs in endothermic cells as well as in ectothermic cells. Therefore, non-shivering 49  $\frac{50}{2}$  thermogenesis is a mechanism in which mitochondrial respiration is exploited to produce heat without statemining and through an inducible H<sup>+</sup> conductance sustained by uncoupling protein-1 (UCP1) in brown synthesizing ATP through an inducible H<sup>+</sup> conductance sustained by uncoupling protein-1 (UCP1) in brown adipose tissue (BAT) [9,36,37]. However, non-shivering thermogenesis also dwells in other organs of 53 mammals and birds, and in particular in skeletal muscles [38]. The proton leak appear in the mitochondria of many different species including mammals, reptiles, amphibians and molluscs. The true in the mitochondria of many different species including mammals, reptiles, amphibians and molluscs. The true function(s) seems to be so important for those living organisms that are ready to pay the high price of energy 57 <sup>58</sup> dissipation to afford the proton leak [7]. These considerations suggest that the energy cost supported by the  $\frac{1}{60}$  mitochondria in this futile cycle of H<sup>+</sup> must be counterbalanced by a high benefit for the cells that interest a great variety of organisms (endotherms and ectotherms). Among the proposed functions, it is also counted the 61 52 synthesizing ATP through an inducit 56 in the mitochondria of many different 59 assipation to arror the proton reak  $62$ 

attenuation of ROS production that is associated with ageing and protection against damage to cellular components. This key role seems to justify the energy cost imposed by the proton leak to the living organism [7]. The components. This key role seems to justify the energy cost imposed by the proton leak to the living organism 1 <sup>2</sup> [7]. The loss of energy, attributed to the dissipation of the  $\Delta p$ , is probably necessary for mitochondrial biology  $_4$  to contain oxidative stress [6].  $\frac{1}{3}$   $\frac{1}{3}$ 

### $\frac{8}{3}$  3. Basal and inducible proton leak 9

 $\frac{10}{11}$  The proton leak is thought to be the result of two processes: basal H<sup>+</sup> conductance, which is not regulated, and 12 an inducible H<sup>+</sup> conductance catalyzed by regulated membrane proteins of the IMM [11,39–42]. The  $13$  physiological relevance of the "basal proton leak" in mammals participates significantly in the basal metabolic The thermogenesis of the body [13]. The basal proton leak is attributed to the  $G_{H+}$  across the lipid rate for the thermogenesis of the body [13]. The basal proton leak is attributed to the  $G_{H+}$  across the lipid bilayer and it has been suggested that the FAs composition of the membrane can modulate it. IMM with a high 16 <sup>17</sup> level of unsaturation index and n-3 polyunsaturated fatty acids (PUFA) and a low level of linoleic acid (18:2  $n-6$ ) has a significant proton leak. Indeed, docosahexaenoic acid (22:6 n-3) is correlated with a high H<sup>+</sup> conductance. Therefore, one might expect that n-3 PUFAs are responsible for the increased permeability of 20 <sup>21</sup> the lipid bilayer [43]. However, it cannot be excluded that this effect is additionally the result of peroxidation  $\frac{22}{23}$  of n-3 FAs and/or the action of products of their decomposition, which act as powerful activators of the anionic carriers [44]. The increase of the proton leak in membranes rich in PUFAs [45] could be also correlated 24 <sup>25</sup> indirectly with an elevated mitochondrial metabolism that requires more fluid membranes to ensure the  $\frac{26}{27}$  appropriate catalytic activity of membrane proteins. The relationship between the presence of PUFAs and the 28 raising of the H<sup>+</sup> conductance would correspond to a basal proton leak assigned to adenine nucleotide translocase (ANT) [39]. 29 11 The proton reak is thought to be the R 15 and for the thermogenesis of the bo- $19$   $\mu$ -0) has a significant proton ican. 23 OF IPS TAS and/or the action of product 27 appropriate catarytic activity of filem  $30 \left( \frac{1}{2} \right)$ 

 $^{31}$  The loss of H<sup>+</sup> through the membrane, at the interface between proteins and lipids, may be responsible for most  $\frac{32}{33}$  of the basal proton leak [13]. It was calculated that more than half of H<sup>+</sup> conductance is dependent on the abundance of membrane-embedded proteins of IMM, specifically, anion carriers of the family SLC25 as ANT 34  $\frac{35}{25}$  [39,46] and UCP1 in BAT [40]. Genetic manipulation of the amount of ANT embedded in the IMM causes a  $\frac{36}{37}$  substantial change in the  $G_{H^+}$  [39]. The basal H<sup>+</sup> conductance may be an inevitable consequence of the structure and abundance of ANT although other anionic carriers can participate in the basal proton leak. Since ANT is 38  $\frac{39}{10}$  the most abundant translocase in the IMM, the role of other carriers may be irrelevant [39]. Contrariwise, in BAT mitochondria, the UCP1 is expressed at equal concentrations of ANT. Studies conducted in conditions 40 of the limited presence of endogenous FAs or with inhibitors of UCP1 and ANT, as well as in conditions with 42 <sup>43</sup> UCP1 knockout, have led Parker and colleagues to consider the UCP1 implicated in basal proton leak [40]. However, not all proteins of the IMM are implicated in basal proton leak. The nicotinamide nucleotide discussion of the IMM are implicated in basal proton leak. The nicotinamide nucleotide transhydrogenase which is up to 2% of total mitochondrial proteins does not affect the basal proton leak. 46 Although it is not exhaustive as absolute proof, available results suggest that the basal  $H^+$  conductance is one way only perpetrated by members of the family of mitochondrial anion carriers and not by other proteins of 50 the IMM. The idea is that the most significant contribution comes from ANT and UCP1 [40].  $33$  of the basal proton reak [15]. It was  $37$  substantial change in the  $G_{H^+}[39]$ . In 41 BAT muochondria, the UCPT is exp 45 However, not all proteins of the IN 49 way only perpetrated by members of

On the contrary, the inducible proton leak is dependent on the activation of ANT [40], UCPs [9,13,37] or the 52 <sup>53</sup> phosphate carrier [47], the aspartate/glutamate carrier [48], and the dicarboxylate carrier [49]. In brown/beige fat, the mitochondrial proton leak and thermogenesis are verified to be caused by UCP1, whereas the inducible 55  $56$  proton leak in extra brown fat tissues was not easily detectable as  $H^+$  conductance activity carrier(s)-dependent [50].  $54$  phosphate carrier  $\left[17\right]$ , are asparately  $57 \frac{1}{1501}$  $58$   $1^{50}$ .

 $\frac{59}{60}$  The turnover of ANT, as demonstrated in the presence and absence of a potent inhibitor carboxyatractylate 61 (CAT), is responsible for the CAT-sensitive inducible H<sup>+</sup> conductance that is catalysed by the translocase in  $60$  The turnover of Alv1, as demonstrat

the presence of FAs, AMP or alkenals [11,39,51]. The proton leak catalyzed by ANT can be inhibited not only by classical inhibitors (bongkrekic acid and CAT) but also by ATP, ADP and the GDP. The latter may partly 1  $\frac{2}{3}$  contribute to inhibiting the H<sup>+</sup> conductance binding weakly to ANT in a non-competitive site of adenine substrates without hindering their transport [11]. The molecular identity of the transport protein(s) driving the substrates without hindering their transport [11]. The molecular identity of the transport protein(s) drivin thermogenic proton leak across the IMM of mitochondria in muscle tissues remained enigmatic for decades, 5  $\frac{6}{7}$  even though proton leak in brown/beige fat is a crucial component of mitochondrial physiology in nonshivering thermogenesis [50]. Recently, studies show that the mitochondria of extra adipose tissues respond to 8 <sup>9</sup> an FA-induced proton leak mediated by ANT [52] and the molecular mechanism generating the H<sup>+</sup> current is  $\frac{10}{11}$  similar to the proton leak of UCP1. Proton leak negatively regulated by ADP/ATP exchange via ANT is reliant on cellular control of ATP synthesis and conseistently, cellular ATP requirement may be used to dynamically 12 control proton leak and mitochondrial uncoupling. [51,53,54]. However, ANT is also considered the main 13  $\frac{14}{15}$  component of low conductance supported by PTP [55]. PTP is sensitive to FAs and uncouples mitochondria 16 by H<sup>+</sup> flow through the IMM [56]. ANT might cause mitochondrial uncoupling by proton leak, (non)selective <sup>17</sup> PTP, or both. 4 substrates without hindering their trans 7 CVCII though proton four in crown of  $11$   $11$   $11$   $11$ 15 Component of Tow conductance supp-18

The archetypal uncoupling protein, UCP1 [36], with its abundance in BAT, carries out the physiological role 19  $\frac{20}{21}$  in non-shivering thermogenesis importing H<sup>+</sup> or, in some models, transport of FA anions from the inner to the outer leaflet of the IMM [57]. UCP1 is activated by FAs and inhibited with nucleotides [58]. Therefore, the 22 <sup>23</sup> proton leak through the IMM of BAT is primarily physiologically regulated by GDP or by free FAs in the  $\frac{24}{25}$  opposite way. Indeed, the UCP1-dependent proton conductance is physiologically activated by FAs that are 26 released by intracellular triacylglycerol under β-adrenergic stimulation in response to cold, or strongly inhibits  $\frac{27}{10}$  by the purine nucleotides [59–61]. UCP1 has four different states of conductance and depends on the presence  $\frac{28}{29}$  of UCP1 regulatory molecules [37]. The absence of purine nucleotides promotes a state of catalytic activity of UCP1 that is greatly improved by the presence of FAs. There is some debate in the literature on the mechanism 30  $^{31}$  related to the role of FAs to induce transport activity since UCP1 can also conduct H<sup>+</sup> in their absence [37,62].  $\frac{32}{33}$  Contrariwise, the purine nucleotides inhibit the activity of UCP1 both experimentally and physiologically [37]. Interestingly, UCP1 is homologous to the ANT whose structure had previously been found [63] and six 34 35 predicted transmembrane helixes can be arranged into three homologous repeats of two helixes each. The molecular and structural characterization of UCP1 contributes to the elucidation of the mechanistic grounds of  $\frac{36}{37}$ its purine nucleotide inhibition. However, one of the most disputed issues in the field of bioenergetics is how 38  $^{39}$  UCP1 supports H<sup>+</sup> transport in the presence of free long-chain FAs [64]. The increasing interest in human and the comparison of th to understand the mechanisms of non-shivering adaptive thermogenesis [65]. 42 21 In non-sinvering thermogenesis impo 25 opposite way. mueed, the OCF 1-uep 29 of OCFT regulatory molecules [37]. 33 Contrariwise, the purine nucleonues i 37 Indictural and structural characterization 41 metabolic disorders related to obesity

<sup>44</sup> The common criterion for H<sup>+</sup> transport by UCP1 and ANT considers the important role of long-chain FAs  $^{45}$  (lcFAs) containing more than 12 carbon atoms. In the presence of lcFAs, UCP1 acts as an H<sup>+</sup> uniporter and 16<br>
17 **LeFAs are lodged within UCP1**. The hydrophobic tails of lcFAs establish hydrophobic interactions with UCP1 48 acting as a cofactor for H<sup>+</sup> transport [66]. Indeed, a single lcFA can guide H<sup>+</sup> transport via UCP1 and facilitate conformation changes between c- and m-state. The protonatable headgroup of lcFA serves as a missing<br>  $\frac{49}{50}$  "stepping stope" for H<sup>+</sup> translocation via UCP1 [67]  $t_{\text{50}}$  such that the state. The problemation is the state of 47 **ICFAS are lodged within UCPI.** The r 51 Stepping stone for H transfocation

Moreover, elevated levels of calcium uptake in mitochondria by mitochondrial calcium uniporter (MCU) 54 stimulate the Krebs cycle and supply more protons, promoting uncoupled respiration and acting as a  $\frac{55}{25}$  thermogenic uniporter. Upon adrenergic stimulation, MCU recruits UCP1 through the essential MCU regulator (EMRE) to form an MCU-EMRE-57  $58$  enhanced H<sup>+</sup> supply for UCP1 operation in the thermogenesis of brown and beige adipose tissue [68]. 52 53 Moreover, elevated levels of calcium  $56$  and  $\frac{1}{2}$  and  $\$ 59

 $60$  In contrast to UCP1, FAs do not induce the  $c-m$  conformational change in ANT. Conversely, conformational  $\frac{61}{62}$  change happens during the adenine nucleotides translocation mechanism [69], whereas an increase in the ANT-62 Change happens during the ademne in

mediated H<sup>+</sup> translocation action is induced by FAs [50,54,70]. LcFAs must bind to the positive side of the 1 IMM in order to trigger the H<sup>+</sup> conductance via ANT, and ANT can be in either the  $c$ - or *m*-state to drive the  $\frac{2}{2}$  proton leak. Moreover, lcFAs anion bound to ANT may induce a mild conformational change allowing H<sup>+</sup> to 4 move via a narrow translocation pathway of ANT [52]. 3 Proton read, Moreover, terms amon or

Noteworthy, DNP, FCCP, SF6847, and BAM15, which are mitochondrial uncouplers that induce <sup>7</sup> pharmacological proton leak across the IMM can activate ANT or UCP1 with a protein-independent  $\frac{8}{9}$  protonophoric mechanism emulating the physiological FA-induced mitochondrial proton leak [71].  $5 - 5$  $6$  Noteworthy, DNP, FCCP, SF6847, 9 Protonophore incenditism emulating to

 $\frac{10}{11}$  The strategies of how nucleotides block proton leak via UCP1 and ANT differ from how carriers are activated with FAs. Nucleotides are transported by ANT, whereas they are inhibitors of UCP1. In contrast to ANT, 12 <sup>13</sup> whose nucleotide-binding site alternately opens to both sides of the IMM, UCP1 has a nucleotide-binding site  $\frac{14}{15}$  located on the cytosolic face of the IMM [64,72]. Purine nucleotide binding on the positive side of IMM blocks 16 the H<sup>+</sup> translocation of UCP1 [64]. The nucleotide antiport and the FA-dependent H<sup>+</sup> translocation in the ANT <sup>17</sup> reveal a close relationship between mitochondrial ATP and heat generation merging two transport modalities that control mitochondrial ATP production or non-shivering thermogenesis in the bioenergetics process [50]. 11 The strategies of now much offices of 15 rotated on the eyes one race of the live 19 liidu control innochonumat ATT produ

20<br>21 The "new" uncoupling proteins (nUCPs), UCP2 and UCP3, with a widespread distribution in different tissues, 22 are a field of interest in the analysis of variation in  $H^+$  conductance in mitochondria treated under artificial  $^{23}$  conditions [37]. Then, UCP4 and UCP5 are mitochondrial carriers widely distributed in the brain but perform  $\frac{24}{25}$  similar conformational and H<sup>+</sup> transport activities of UCP1 UCP3 [73]. Due to the possible neuroprotective effects of the UCP-dependent decrease of ROS production in the nervous system, UCP4 and UCP5 might play 26  $\frac{27}{20}$  a significant role to prevent neurological disorders [74]. Due to their ubiquitous expression, UCP2 and UCP3 28<br>
may be able to mediate mitochondrial uncoupling in tissues other than brown fat. The nUCPs can transport the 30 H<sup>+</sup> under activation by specific agents, whereas the H<sup>+</sup> conductance is inhibited with purine nucleotides  $^{31}$  [75,76]. nUCPs catalyze an inducible H<sup>+</sup> conductance in the presence of specific activators, which include the products of lipid peroxidation [9]. How occurs the catalysis of proton leak through UCP2 and UCP3 in the products of lipid peroxidation [9]. How occurs the catalysis of proton leak through UCP2 and UCP3 in the presence of physiological concentrations of ATP and ADP in the cell remains to be understood [37]. It is 34 assumed that the inhibition by purine nucleotides is relieved by the FAs as proposed for the UCP1. Nevertheless, there are no results that consider the nUCPs responsible for a fraction of the proton leak in Nevertheless, there are no results that consider the nUCPs responsible for a fraction of the proton leak in mitochondria. In the absence of UCP2 and UCP3, mitochondria do not show improved coupling status of 38 <sup>39</sup> OXPHOS [77] and nUCPs are not involved in controlling body weight or adaptive non-shivering <sup>40</sup><br><sup>41</sup> thermogenesis [78]. However, UCP3 knockout mice have increased mitochondrial respiration coupling in skeletal muscle mitochondria [79]. Endogenous expression of UCP3 has uncoupling activity and its absence 42 may result in increased ROS production [79] as well as a thermogenic response in skeletal muscle induced by MDMA (ecstasy) [80]. In addition to this, UCP3 contributes to the export of mitochondrial FA anions, preventing mitochondrial damage brought on by lipid peroxidation [81]. 46 21 The new uncoupling proteins (not 25 SIMINAL COMOTINATIONAL AND **17** Transp 29 may be able to mediate infloction and 33 products of liptu peroxidation [9]. H 37 Nevertheless, there are no results the 41 thermogenesis [78]. However, UCP. 45 MDMA (ecstasy) [80]. In addition

Thus, it is difficult to distinguish between an inducible proton leak by nUCPs observed experimentally and 48 what occurs in the cell under physiological conditions [52]. The purported mild uncoupling activity of UCP2 has been reassessed highlighting its biochemical role in mitochondrial oxidation of glucose, glutamine 51 Exporting out of mitochondria, and the exchange of four-carbon dicarboxylate Krebs cycle intermediate (e.g., oxaloacetate and malate) for phosphate plus an H<sup>+</sup> from opposite sides of the membrane [82]. In cell bioenergetics, UCP2 reveals a novel regulatory mechanism in cellular metabolic demand or substrate 55 utilization. The UCP2 activity may promote the switch of glucose metabolism to fatty acid metabolism 56  $\frac{57}{50}$  controlling the interaction between UCP2 and ANT [83]. Therefore, H<sup>+</sup> conductance or four-carbon metabolite transport via UCP2 may be influenced by ANT [83]. 59 50 What occurs in the cent and epitysion 52 exporting out of mitochondria, and the exchange of four-carbon dicarboxylate Krebs cycle intermediate (e.g., 54 **Conditional California** malately for phosp 58 controlling the interaction between  $\sigma$ 

63 64

The physiological differences between the UCPs and  $F_1F_0$ -ATPase activities during the  $\Delta p$  dissipation are the uncoupling or coupling of respiration and ADP phosphorylation in mitochondria, respectively (Fig. 1) [21,8 uncoupling or coupling of respiration and ADP phosphorylation in mitochondria, respectively (Fig. 1) [21,84]. 1 <sup>2</sup> Therefore, if ATP synthesis by  $F_1F_0$ -ATPase was dissipated manipulating cellular energy expenditure, the  $\frac{1}{4}$  result would be a mechanism involving direct H<sup>+</sup> recycling to override respiratory control reflecting an UCPindependent thermogenic mechanism based on dissipative hydrolysis of ATP in beige and brown adipose tissue 5  $\frac{6}{7}$  [41]. The suggestion for this bioenergetic phenomenon of non-shivering thermogenesis is attributed to a futile <sup>8</sup> creatine cycle [42]. Mitochondrial phosphocreatine(PCr)/creatine (Cr) circuit is sustained by mitochondrial <sup>9</sup> creatine kinase using mitochondrial ATP in the interconversion of Cr to PCr and liberation of ADP.  $\frac{10}{11}$  Contrariwise, a phosphatase might replenish the Cr pool by hydrolyzing PCr. The substrate ATP and the product ADP of creatine cycle are exchanged by ANT increasing the rate of mitochondrial respiration driven 12 13 by ATP synthesis of  $F_1F_0$ -ATPase [42]. Therefore, substrate oxidation during mitochondrial respiration driven  $\frac{14}{15}$  by ATP expenditure by futile creatine cycle cause a noncanonical UCP1-independent, but ATP-dependent, non-shivering thermogenesis. This process can counter obesity and glucose dysregulation in pre-clinical 16  $17 \text{ models} [85-87].$  $\frac{3}{3}$   $\frac{1000000}{3}$   $\frac{111}{3}$   $\frac{9}{3}$   $\frac{1000000}{3}$   $\frac{9}{3}$   $\frac{110}{3}$ 7 [11]. The suggestion for this ordering 11 Communisty, a prosphanol might  $15$  by  $111$  experience by Tattle erealize

Generally, the inducible proton leak can be alleviated by the addition of bovine serum albumin, which removes 19  $\frac{20}{21}$  FAs and derivatives of reactive alkenals considered endogenous activators of anion carriers of IMM [46]. The 22 endogenous activation of  $G_H$  seems to be directly proportional to the energy state of the IMM and the effect 23 is not dependent on the redox state of ETC but dependent on the  $\Delta \psi$ . The  $\Delta \psi$  could change the conformation  $\frac{24}{25}$  of the anion carriers by exposing the binding sites to the activator molecules. The IMM energization dependent on mitochondrial uncoupling, whatever the mechanism involved or the activators who participate, could have 26 27 its importance for the cell because it limits the  $\Delta p$  during the state IV and as a consequence decreases ROS  $\frac{28}{29}$  production [11]. 21 TAS and derivatives of reactive anche 25 Of the amon carriers by exposing the t  $29$  production [11].

### 4. Proton leak: a biological formula of prevention against ROS 33

 $\frac{35}{25}$  The formation of superoxide anion depends on the redox potential of the electron donor (respiratory carriers),  $\frac{36}{37}$  the concentration of O<sub>2</sub> (the acceptor) and the second-order rate constant for the reaction between them. The 38 standard reduction potential ( $E^{\circ}$ ) to transfer an electron to  $O_2$  to form  $O_2$  is -160 mV [88]. By considering  $O_2$  pK<sub>a</sub> value of 4.7 [88], the E<sup>o</sup> does not vary in the range of physiological pH of living organisms [89]. Since the reduction potential (E) is determined as the product of  $E^{\circ}$  and the ratio  $[O_2]/[O_2^-]$ , according to  $41$  $\mu$  possible O<sub>2</sub><sup>-</sup> concentrations that might be obtained in the matrix by assuming a low [O<sub>2</sub>] of 1  $\mu$ M, which is <sup>43</sup> enough lower than the 3-30  $\mu$ M range measured *in vivo*, mitochondria can thermodynamically support the  $\frac{44}{45}$  reduction of O<sub>2</sub> to O<sub>2</sub><sup>-</sup> [28,89].  $37$  the concentration of  $O_2$  (the acceptor  $41$  Since the reduction potential (E) is  $45$  reduction of  $O_2$  to  $O_2$ . [28,89].

<sup>46</sup> The sites of O<sub>2</sub><sup> $-$ </sup> formation in the ETC are Complexes I or III, especially in resting conditions with decreased 48 ATP production, slow respiration, high ratio of NADH/NAD<sup>+</sup>, and high concentration of reduced coenzyme  $Q$  (QH<sub>2</sub>) associated with high  $\Delta p$  [90]. Under conditions of low energy demand, the accumulation of NADH in the mitochondria creates a fully reduced FMN in Complex I and consequently the formation of  $O_2$  [89,91]. 52 Seo and colleagues showed in mammalian mitochondria, independently of the overexpression of NADH  $^{53}_{54}$  dehydrogenase of Saccharomyces cerevisiae, an NADH/NAD<sup>+</sup> ratio associated with reduced production of  $O_2$  [92]. Complex I can generate  $O_2$  during reverse electron transport that occurs if the electron flow towards 56 O<sub>2</sub> reduction in Complex IV is blocked and there is a high QH<sub>2</sub> pool and  $\Delta p$  [93–96]. In this situation, the  $^{57}_{50}$  electrons of QH<sub>2</sub> are driven by the thermodynamic strength of the  $\Delta p$  to return to Complex I where the 59 production of  $O_2$  is extremely powerful. However, it can be abolished by decreasing the  $\Delta \psi$  [89,97,98]. This 60 conclusion is based on observations that the addition of uncoupling agents, which reduce the  $\Delta p$ , decreases the For the of production of O<sub>2</sub><sup>-</sup>, although it is the collapse of  $\Delta pH$  that limits the ROS generation in Complex I [98].  $47$  I he sites of  $O_2$  formation in the E1  $50 \times (112)$  associated with light  $2p$  [20 54 acrystrogenase of Baccharomyces et  $58$  creations of Q112 are driven by the  $62$  and or production of  $O_2$ , annough it

<sup>7</sup> The rate of  $O_2$ <sup>-</sup> generation can be very slow if an  $\frac{8}{9}$  too far away from O<sub>2</sub>. In the biological system, the transfer of electrons is supported by the existence of electron [99,100]. The production  $11$  of O<sub>2</sub> anion-sensitive to mild uncoupling can occur only by accessing the sites in which the electrons can be  $\frac{12}{12}$  lost and received by O<sub>2</sub>. Distances greater than those allowed for a fast transfer between O<sub>2</sub> and respiratory 14 carriers can minimize the formation of ROS [89]. The  $O_2$  is impermeable to the IMM and has a very short <sup>15</sup> life-time being rapidly converted to  $H_2O_2$  by Mn-superoxide dismutase (Mn-SOD or SOD-2). The  $H_2O_2$ formed is degraded by the enzymes glutathione peroxidase and peroxiredoxin III to H<sub>2</sub>O. The H<sub>2</sub>O<sub>2</sub> can react 18 alternately through two chemical reactions: with metal ions (such as  $Fe^{2+}$ ) in the known Fenton reaction or 19 with another molecule of  $O_2$ <sup>-</sup> in the Haber-Weiss reaction produces the highly toxic  $\cdot$ OH that, unlike  $O_2$ <sup>-</sup> and  $^{20}_{21}$  H<sub>2</sub>O<sub>2</sub>, can extract the first hydrogen atom from a methylene (-CH<sub>2</sub>-) group of PUFA to start the lipid 22 peroxidation process [9]. The  $O_2$  and lipid peroxidation products are potent activators of H<sup>+</sup> conductance by <sup>23</sup> UCPs in mitochondria. The idea that the O<sub>2</sub> activates UCPs arises from the results of lipid peroxidation  $\frac{24}{25}$  products such as 4-hydroxy-trans-2-nonenal (HNE) inducing the proton leak through UCPs [9,20,101]. 26 Malingriaux and colleagues suggest that aldehyde does not directly activate UCP1 or UCP2. HNE, on the other <sup>27</sup> hand, significantly increased the membrane conductance mediated by different lcFAs in both UCP-containing 28 and UCP-free membranes [102]. Moreover, the PTP may mediate a portion of the proton leak effect of HNE on brown-fat mitochondria [103,104]. 30  $9$  we far away from  $\sigma_2$ . In the orongical 10 tunnels with maximum distances bet  $13$  Tost and received by  $O_2$ . Distances  $g$ 17 **IDITION** IS degraded by the chargines g  $21$   $1202$ , can exitate the first hydroge  $25$  products such as  $4$ -hydroxy-trails- $2$ 29 and OCT-free memoranes [102]. NO.

The catalytic activity of UCPs can decrease the ROS concentration within the mitochondria [20,105]. These 32 33 observations suggest the hypothesis for the functions of the UCPs. OXPHOS mild uncoupling UCPsdependent decreases the  $\Delta p$  and relieves the production of  $O_2$ . Accordingly, cells are protected from oxidative damage at the cost of a slight lowering of the efficiency of OXPHOS [61,77,101]. The relationship between 36  $^{37}$  UCPs and ROS has been demonstrated in both isolated mitochondria and intact cells by the existence of a mild  $\frac{38}{39}$  uncoupling UCP-specific during the production of ROS. Indeed, the decrease in O<sub>2</sub> production is reverted by 40 a GDP-sensitive H<sup>+</sup> conductance [101,105]. Therefore, the proton leak can attenuate the O<sub>2</sub> generation <sup>41</sup> suggesting a system controlled by a feedback loop in which the ROS induces an H<sup>+</sup> conductance sustained by 12<br>
UCPs able to decrease ROS production [20,106]. Considering the uncoupling function of UCPs and the adverse action on mitochondrial energy production, in mitochondrial biology, the apparent ambiguous nature of UCPs 44  $^{45}_{46}$  has been selected for the protective effect against oxidative stress exploiting a mild uncoupling of OXPHOS [61].  $35$  dependent decreases the  $\Delta p$  and reflex 39 uncoupling OCP-specific during the p 43 UCPs able to decrease ROS production 46 Fell Legal Communication and protective 47 [01].

<sup>48</sup> The  $\Delta p$  has a special role also in the supramolecular organization of respiratory complexes that constitute the system of OXPHOS [107,108]. Indeed, the association and the structural organization in respiratory 50  $\frac{51}{2}$  supercomplex can be modulated by  $\Delta p$  [109,110]. The dissociation of supercomplexes in individual units 53 occurs at high values of  $\Delta p$  [111] causing the O<sub>2</sub><sup>-</sup> production sensitive to  $\Delta p$  suggesting a link between the two events [110]. Consistently, supercomplexes can hide auto-oxidizable prosthetic groups preventing their 54  $\frac{55}{25}$  reaction with O<sub>2</sub> [112]. This could be a possible explanation for the role of the supercomplex organization to limit ROS formation [97,107]. Moreover, a direct channelling of substrates in supercomplexes increases the 57 58 flux of electrons and the respiratory chain remains oxidized [97,112,113]. The proton leak might <sup>59</sup> valve" dissipating the  $\Delta p$  slowing the disassembly of supercomplexes and preventing the formation of O<sub>2</sub>.  $49$  I he  $\Delta p$  has a special role also in the s 52 supercomptex can be modulated by  $56$  reaction with  $O_2$  [112]. This could be  $60$  valve dissipating the  $\Delta p$  slowing the

### 5. Conclusions

2 During mitochondrial respiration, a cross-connection between the  $J_{H^+}$  and the redox reactions ensures the  $\frac{3}{4}$  mitochondrial H<sup>+</sup> circuit coupled to electrons transfer in the ETC. Consistently, the resting state should promote the reduced steady state of respiratory carriers without ATP synthesis. Physiological oxygen 6 consumption in the mitochondria independent of ADP phosphorylation can exploit the non-ohmic H<sup>+</sup> conductance stimulated by  $\Delta p$  values above the inherent state IV. The decrease of protonic backpressure on the respiratory chain is also related to UCPs and ANT induction of proton leak. Proton leak, a key component  $\frac{8}{9}$ of mitochondrial energy production, is a feature of mitochondria occurring in all tissues. The dissipation of the 10 <sup>11</sup>/<sub>12</sub> excessive H<sup>+</sup> potential by proton leak can prevent the formation of O<sub>2</sub><sup>-</sup> by OXPHOS mild uncoupling, thus ensuring a system of prevention of damage caused to ROS production. In IMM the increase of the proton leak ensuring a system of prevention of damage caused to ROS production. In IMM the increase of the proton leak is probably a safer way for the welfare of aerobic living organisms. 14 5 promote the reduced steady state of 9 the respiratory chain is also related to 13 ensuring a system of prevention of da

### 20 **Figure captions**

22 Figure 1. Proton circuit of the mitochondrial chemiosmotic system. Reduced substrates are oxidized by the respiratory complexes (CI, CII, CIII, and CIV) to create the proton gradient (arrows with solid or dashed lines) 23  $\frac{24}{25}$  dissipated by F<sub>1</sub>F<sub>O</sub>-ATPase (CV) and by the proton conductance through the IMM and/or the mitochondrial  $\frac{25}{26}$  carriers, in particular UCPs (uncoupling proteins) and adenine nucleotide translocase (ANT). Q<sub>10</sub>, coenzyme  $Q_{10}$ ; Cyt c, cytochrome c; IMM, Inner mitochondrial membrane. Figure created with BioRender (BioRender.com). 28 26 Carriers, in particular OCFs (uncoupl

Figure 2. Conditions for welling of non-respiring mitochondria. Passive osmotic swelling of mitochondria in potassium acetate (KAc) (A) in the presence of valinomycin (Val) plus FCCP or (B) in the presence of 33 nigericin (Nig). The decrease of light-scattering, identify as a decrease of optical density (O.D.) of 34  $\frac{35}{26}$  mitochondria spectrophotometrically measured to 540 nm, is related to mitochondrial swelling depending on the nature of the IMM permeability induced by the ionophores. Figure created with BioRender (BioRender.com). 38 31 32 Figure 2. Conditions for welling of 37 line nature of the livity permeable

<sup>41</sup> Figure 3. Relationship current-voltage (I/V) for ohmic and non-ohmic circuits. Linear ratio I/V that <sup>44</sup> non-ohmic trend is typically observed in mitochondria by proton leak (Figure adapted from [13]).  $42$  rigure 3. Relationship current-vo observes Ohm's Law (straight line); exponential curve (dash line) that does not comply with Ohm's Law. The<br>
44 45 honorando de cypreary observed

### $\frac{51}{52}$  Acknowledgements 52 ACKHOWIEGGEMENTS

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### <sup>61</sup> References

- $[1]$ M. Saraste, Oxidative phosphorylation at the fin de siècle, Science. 283 (1999) 1488-1493. 1 https://doi.org/10.1126/science.283.5407.1488.
- 2 [2] B.E. Schultz, S.I. Chan, Structures and proton-pumping strategies of mitochondrial respiratory <sup>3</sup> enzymes, Annu Rev Biophys Biomol Struct. 30 (2001) 23–65.  $\frac{4}{1000}$  https://doi.org/10.1146/oppur
- $5$  mups.//wol.org/10.1140/annum  $\zeta$  [3] P. Mitchell, Proton current flow in mitochondrial systems, Nature. 214 (1967) 1327–1328. 7 https://doi.org/10.1038/2141327a0.
- 8 [4] D.G. Nicholls, S.J. Ferguson, 4 The Chemiosmotic Proton Circuit in Isolated Organelles: Theory and Practice, in: Bioenergetics (Fourth Edition), Academic Press, Boston, 2013: pp. 53–87.  $10$ <br>https://doi.org/10.1016/B978-0-12-388425-1.00004-X.
- 11  $12$   $\phantom{0}$   $\phantom{0}$  13 of mechanism, Nature. 191 (1961) 144-148.
- [6] D.G. Nicholls, Forty years of Mitchell's proton circuit: From little grey books to little grey cells,<br>15 Biochim Bionbys Acta 1777 (2008) 550–556 https://doi.org/10.1016/i.bhabio.2008.03.014 Biochim Biophys Acta. 1777 (2008) 550-556. https://doi.org/10.1016/j.bbabio.2008.03.014.
- 16 [7] M.D. Brand, Uncoupling to survive? The role of mitochondrial inefficiency in ageing, Exp Gerontol. 35  $17$  [1]  $10000 \, 011 \, 020 \, \text{https://www.14e6.070}$  $18$  (2000) 011-020.  $\frac{18}{100}$
- $\frac{1}{19}$  [8] C.E. Amara, E.G. Shankland, S.A. Jubrias, D.J. Marcinek, M.J. Kushmerick, K.E. Conley, Mild 20 mitochondrial uncoupling impacts cellular aging in human muscles in vivo, Proc Natl Acad Sci U S A. 21 104 (2007) 1057-1062. https://doi.org/10.1073/pnas.0610131104.
- 22 [9] K.S. Echtay, Mitochondrial uncoupling proteins--what is their physiological role?, Free Radic Biol 23<br>Med. 43 (2007) 1351–1371. https://doi.org/10.1016/j.freeradbiomed.2007.08.011.  $24$  Micu.  $\rightarrow$  (2007) 1991 1971.
- $25$  [10] D.G. Nicholls, The effective pr 26 adipose tissue. Dependency on proton electrochemical potential gradient, Eur J Biochem. 77 (1977) 27 349-356. https://doi.org/10.1111/j.1432-1033.1977.tb11674.x.
- <sup>28</sup> [11] N. Parker, C. Affourtit, A. Vidal-Puig, M.D. Brand, Energization-dependent endogenous activation of 29 proton conductance in skeletal muscle mitochondria, Biochem J. 412 (2008) 131–139. 30 Proton conductance in site can  $31$  11ttps.//doi.org/10.1042/bJ20
- 32 [12] D.G. Nicholls, Mitochondrial ion circuits, Essays Biochem. 47 (2010) 25-35. 33 https://doi.org/10.1042/bse0470025.
- 34 [13] M. Jastroch, A.S. Divakaruni, S. Mookerjee, J.R. Treberg, M.D. Brand, Mitochondrial proton and <sup>35</sup> electron leaks, Essays Biochem. 47 (2010) 53–67. https://doi.org/10.1042/bse0470053.
- 36 [14] C Negei E Trembetti V Vent  $37$   $14$  J. Nesci, it is non-term in the set  $38$  mitochondrial permeability transition pore: an overview, Biochimie. 152 (2018) 85–93. 39 https://doi.org/10.1016/j.biochi.2018.06.022.
- 40 [15] M.D. Brand, D.G. Nicholls, Assessing mitochondrial dysfunction in cells, Biochem. J. 435 (2011) 297– 41 312. https://doi.org/10.1042/BJ20110162.
- $\frac{42}{161}$  VN Samartsoy A A Somonov 43 **10** v.w. summitted, A.A. sements  $\frac{13}{44}$  Uncouplers and Decoupling Agents as Inducers of Free Respiration in Mitochondria in States 3 and 4: 45 Theoretical and Experimental Approaches, Cell Biochem Biophys. 78 (2020) 203-216. 46 https://doi.org/10.1007/s12013-020-00914-5.
- <sup>47</sup> [17] S.I. Liochev, I. Fridovich, Superoxide and iron: partners in crime, IUBMB Life. 48 (1999) 157–161.  $^{48}$  https://doi.org/10.1080/713803492.
- $\frac{49}{40}$   $\frac{100}{100}$   $\frac{100}{100}$   $\frac{100}{100}$   $\frac{100}{100}$   $\frac{100}{100}$   $\frac{100}{100}$   $\frac{100}{100}$   $\frac{100}{100}$  $50$  [10] v.f. skuldchev, nole of uncou 51 oxygen and its one-electron reductants, Q Rev Biophys. 29 (1996) 169-202. 52 https://doi.org/10.1017/s0033583500005795.
- 53 [19] G. Lenaz, The mitochondrial production of reactive oxygen species: mechanisms and implications in 54 human pathology, IUBMB Life. 52 (2001) 159-164. https://doi.org/10.1080/15216540152845957.
- 55 [30] K.C. Febtov, M.D. Prand, 4 by  $56$  [20] K.J. Ethiay, M.D. Dianu, 4-iiyo  $\frac{57}{57}$  of mitochondrial ROS production, Redox Rep. 12 (2007) 26–29. 58 https://doi.org/10.1179/135100007X162158.
- 59 [21] D.G. Nicholls, Mitochondrial proton leaks and uncoupling proteins, Biochim Biophys Acta Bioenerg. 60 1862 (2021) 148428. https://doi.org/10.1016/j.bbabio.2021.148428. 61

- $[22]$ G.P. Brierley, M. Jurkowitz, K.M. Scott, A.J. Merola, Ion transport by heart mitochondria. XX. Factors 1 affecting passive osmotic swelling of isolated mitochondria, J Biol Chem. 245 (1970) 5404–5411.
- 2 [23] K.D. Garlid, R.A. Nakashima, Studies on the mechanism of uncoupling by amine local anesthetics.  $\frac{3}{2}$  Evidence for mitochondrial pro  $\frac{4}{7074,7090}$  $\frac{5}{5}$   $\frac{1514}{1500}$ ,  $\frac{1}{5000}$ ,  $\frac{1}{5000}$
- $6$  [24] L.B. Popova, E.S. Nosikova, E.A. 7 Antonenko, Protonophoric action of triclosan causes calcium efflux from mitochondria, plasma 8 membrane depolarization and bursts of miniature end-plate potentials, Biochim Biophys Acta 9 Biomembr. 1860 (2018) 1000-1007. https://doi.org/10.1016/j.bbamem.2018.01.008.
- 10 [25] P.D. Davis, G.D. Parbrook, G.N.C. Kenny, CHAPTER 7 Diffusion and Osmosis, in: P.D. Davis, G.D. 11  $12$  Particular, G.N.C. Kenny (Eqs.) 13 Butterworth-Heinemann, 1995: pp. 89-102. https://doi.org/10.1016/B978-0-7506-1713-0.50012-3.
- [26] G. Krishnamoorthy, P.C. Hinkle, Non-ohmic proton conductance of mitochondria and liposomes,<br>15 Biochemistry 23 (1984) 1640–1645 https://doi.org/10 1021/bi00303a009 Biochemistry. 23 (1984) 1640-1645. https://doi.org/10.1021/bi00303a009.
- 16 [27] D. Pietrobon, M. Zoratti, G.F. Azzone, Molecular slipping in redox and ATPase H+ pumps, Biochim  $17$  Pinetropoly m. Editity on 1. 18 **DIOPHYS ACTA** . 72J (1903) 317-
- $\frac{1}{19}$  [28] S. Papa, V.P. Skulachev, Reactive oxygen species, mitochondria, apoptosis and aging, Mol Cell 20 Biochem. 174 (1997) 305-319.
- 21 [29] B. Kadenbach, Intrinsic and extrinsic uncoupling of oxidative phosphorylation, Biochim Biophys Acta. 22 1604 (2003) 77-94. https://doi.org/10.1016/s0005-2728(03)00027-6.
- $\frac{23}{24}$  [30] K. Faxén, G. Gilderson, P. Adelroth, P. Brzezinski, A mechanistic principle for proton pumping by 24  $[30]$  K. Paken, G. Ghaerson, P. Aug. 25 **Cylochrome c oxidase, Nature**
- 26 [31] I. Belevich, M.I. Verkhovsky, M. Wikström, Proton-coupled electron transfer drives the proton pump 27 of cytochrome c oxidase, Nature. 440 (2006) 829-832. https://doi.org/10.1038/nature04619.
- <sup>28</sup> [32] G.F. Azzone, M. Zoratti, V. Petronilli, D. Pietrobon, The stoichiometry of H+ pumping in cytochrome 29 oxidase and the mechanism of uncoupling, J Inorg Biochem. 23 (1985) 349–356. 30 bitter //dxi sus/10.1016/0169  $31$   $\frac{1}{2}$   $\frac{1}{$
- 32 [33] B.D. Price, M.D. Brand, Proton translocation by the mitochondrial cytochrome b-c1 complex is 33 inhibited by NN'-dicyclohexylcarbodi-imide, Biochem J. 206 (1982) 419-421. 34 https://doi.org/10.1042/bj2060419.
- $\frac{35}{2}$  [34] A.A. Semenova, V.N. Samartsev, M.V. Dubinin, The stimulation of succinate-fueled respiration of rat 36 [ev.j. film a computer both film computer both A k  $37$  $38$  of the inner membrane. Intrinsic uncoupling of the bc1 complex, Biochimie. 181 (2021) 215–225. 39 https://doi.org/10.1016/j.biochi.2020.12.021.
- 10 [35] D.F. Rolfe, M.D. Brand, The physiological significance of mitochondrial proton leak in animal cells and tissues, Biosci Rep. 17 (1997) 9-16. https://doi.org/10.1023/a:1027327015957.
- [36] D.G. Nicholls, E. Rial, A history of the first uncoupling protein, UCP1, J Bioenerg Biomembr. 31 (1999) 43  $44$  399–400. https://doi.org/10.1
- 45 [37] D.G. Nicholls, The physiological regulation of uncoupling proteins, Biochim. Biophys. Acta. 1757 46 (2006) 459-466. https://doi.org/10.1016/j.bbabio.2006.02.005.
- <sup>47</sup> [38] N.C. Bal, S. Singh, F.C.G. Reis, S.K. Maurya, S. Pani, L.A. Rowland, M. Periasamy, Both brown adipose tissue and skeletal muscle thermogenesis processes are activated during mild to severe cold 49 adentation in miss LDial Cha  $50$  audplation in thice, J biot Che
- 51 https://doi.org/10.1074/jbc.M117.790451.<br>52 [39] M.D. Brand, J.L. Pakay, A. Ocloo, J. Kokoszk. 52 [39] M.D. Brand, J.L. Pakay, A. Ocloo, J. Kokoszka, D.C. Wallace, P.S. Brookes, E.J. Cornwall, The basal 53 proton conductance of mitochondria depends on adenine nucleotide translocase content, Biochem 54 J. 392 (2005) 353-362. https://doi.org/10.1042/BJ20050890.
- 55 [40] N Darker B.C. Criebton A.L.  $56$   $140$   $11.1$  and  $1.0.$  Chemon, A.J. v  $\frac{57}{57}$  basal proton conductance of brown adipose tissue mitochondria, J Bioenerg Biomembr. 41 (2009) 58 335-342. https://doi.org/10.1007/s10863-009-9232-8.
- 59 [41] D.G. Nicholls, M.D. Brand, A critical assessment of the role of creatine in brown adipose tissue <sup>60</sup> thermogenesis, Nat Metab. 5 (2023) 21-28. https://doi.org/10.1038/s42255-022-00718-2. 61

- $[42]$ L. Kazak, E.T. Chouchani, M.P. Jedrychowski, B.K. Erickson, K. Shinoda, P. Cohen, R. Vetrivelan, G.Z. 1 Lu, D. Laznik-Bogoslavski, S.C. Hasenfuss, S. Kajimura, S.P. Gygi, B.M. Spiegelman, A creatine-driven 2 substrate cycle enhances energy expenditure and thermogenesis in beige fat, Cell. 163 (2015) 643- $\frac{3}{2}$  655. https://doi.org/10.1016/j.cell.2015.09.035.
- $\frac{4}{100}$  DK Derter A Littlebert MD F  $\frac{1}{5}$   $\frac{1}{5}$   $\frac{1}{100}$   $\frac{1}{100}$   $\frac{1}{100}$   $\frac{1}{100}$   $\frac{1}{100}$   $\frac{1}{100}$  $6$  membrane surface area and fatty acid composition, Am J Physiol. 271 (1996) R1550-1560. 7 https://doi.org/10.1152/ajpregu.1996.271.6.R1550.
- 8 [44] N. Parker, A. Vidal-Puig, M.D. Brand, Stimulation of mitochondrial proton conductance by <sup>9</sup> hydroxynonenal requires a high membrane potential, Biosci Rep. 28 (2008) 83–88.  $10$  https://doi.org/10.1042/BSR20080002.
- $11$   $[45]$   $[36]$   $[38]$   $[38]$   $[39]$  $12$  [45] E.IVI. FOITLAILLE, IVI. IVIOUSSA, A. 13 polyunsaturated fatty acids deficiency on oxidative phosphorylation in rat liver mitochondria, Biochim Biophys Acta. 1276 (1996) 181–187. https://doi.org/10.1016/0005-2728(96)00075-8.<br>15 [16] E.N. Mokhova J.S. Khailova, Involvement of mitochondrial inner membrane anion carriers in t
- [46] E.N. Mokhova, L.S. Khailova, Involvement of mitochondrial inner membrane anion carriers in the 16 uncoupling effect of fatty acids, Biochemistry (Mosc). 70 (2005) 159-163. 17 ancoupling creat of latty acts  $18$  mtps.//woi.org/10.1007/3103
- $\frac{1}{19}$  [47] M. Zácková, R. Krämer, P. Jezek, Interaction of mitochondrial phosphate carrier with fatty acids and 20 hydrophobic phosphate analogs, Int J Biochem Cell Biol. 32 (2000) 499-508. 21 https://doi.org/10.1016/s1357-2725(00)00006-6.
- [48] V.N. Samartsey, A.V. Smirnov, I.P. Zeldi, O.V. Markova, E.N. Mokhova, V.P. Skulachev, Involvement of 23<br>aspartate/glutamate antiporter in fatty acid-induced uncoupling of liver mitochondria, Biochim  $24$  asparance and and  $24$  $25$  Biophys Acta. 1319 (1997) 25.
- 26 [49] M.R. Wieckowski, L. Wojtczak, Involvement of the dicarboxylate carrier in the protonophoric action 27 of long-chain fatty acids in mitochondria, Biochem Biophys Res Commun. 232 (1997) 414-417. 28 https://doi.org/10.1006/bbrc.1997.6298.
- [50] A.M. Bertholet, Y. Kirichok, Mitochondrial H+ Leak and Thermogenesis, Annu Rev Physiol. 84 (2022)  $30$  [30]  $7000$  Berthold,  $1000$ ,  $1000$  $31$  301-407.  $31$
- 32 [51] null Andreyev AYu, T.O. Bondareva, V.I. Dedukhova, E.N. Mokhova, V.P. Skulachev, N.I. Volkov, 33 Carboxyatractylate inhibits the uncoupling effect of free fatty acids, FEBS Lett. 226 (1988) 265-269. 34 https://doi.org/10.1016/0014-5793(88)81436-4.
- $\frac{35}{2}$  [52] A.M. Bertholet, E.T. Chouchani, L. Kazak, A. Angelin, A. Fedorenko, J.Z. Long, S. Vidoni, R. Garrity, J. 36 [04] Amm Bermeier, Erromanna.  $37$  CIIO, IV. Teraud, D.C. Wallace,  $\frac{38}{38}$  mitochondrial ADP/ATP carrier, Nature. 571 (2019) 515–520. https://doi.org/10.1038/s41586-019-39 1400-3.
- 40 [53] A.S. Divakaruni, M.D. Brand, The regulation and physiology of mitochondrial proton leak, Physiology 41 (Bethesda). 26 (2011) 192-205. https://doi.org/10.1152/physiol.00046.2010.
- $\frac{42}{1541}$  VD Skulachov Eatty acid circle  $43$   $197$   $\cdot \cdot \cdot$  Skalactic  $\cdot \cdot \cdot$  atty acta circ
- phosphorylation, FEBS Lett. 294 (1991) 158–162. https://doi.org/10.1016/0014-5793(91)80658-p.<br>45 [55] M.A. Neginskaya, M.E. Solesio, E.V. Berezhnaya, G.F. Amodeo, N. Mnatsakanyan, E.A. Jonas, E.V. M.A. Neginskaya, M.E. Solesio, E.V. Berezhnaya, G.F. Amodeo, N. Mnatsakanyan, E.A. Jonas, E.V. 46 Pavlov, ATP Synthase C-Subunit-Deficient Mitochondria Have a Small Cyclosporine A-Sensitive 47 Channel, but Lack the Permeability Transition Pore, Cell Rep. 26 (2019) 11-17.e2.  $^{48}$  https://doi.org/10.1016/j.celrep.2018.12.033.
- 49 ELCL MD Missionali L. Meitard  $50$  [30] IVI.N. WIECKOWSKI, L. WOJICZAK  $51$  due to opening of the mitochondrial permeability transition pore, FEBS Lett. 423 (1998) 339–342. 52 https://doi.org/10.1016/s0014-5793(98)00118-5.
- 53 [57] V.P. Skulachev, Anion carriers in fatty acid-mediated physiological uncoupling, J Bioenerg Biomembr. 54 31 (1999) 431-445. https://doi.org/10.1023/a:1005492205984.
- $55$   $[FO]$   $E \text{ Piol } E \text{ Aquirreacite } L \text{ line}$  $56$  [JO] L. Mai, L. Aguiregonia, J. Jine  $\frac{57}{57}$  protein UCP1: implications for the transport mechanism, Biochim Biophys Acta. 1608 (2004) 122– 58 130. https://doi.org/10.1016/j.bbabio.2003.11.001.
- 59 [59] D.G. Nicholls, Hamster brown-adipose-tissue mitochondria. Purine nucleotide control of the ion <sup>60</sup> conductance of the inner membrane, the nature of the nucleotide binding site, Eur J Biochem. 62  $61$  (1976) 223-228 https://doi.org  $62$  (1576) 225 226. https://doi.o

- $[60]$ G.M. Heaton, D.G. Nicholis, Hamster brown-adipose-tissue mitochondria. The role of fatty acids in 1 the control of the proton conductance of the inner membrane, Eur J Biochem. 67 (1976) 511-517. 2 https://doi.org/10.1111/j.1432-1033.1976.tb10717.x.
- $\frac{3}{2}$  [61] Z.B. Andrews, S. Diano, T.L. Horvath, Mitochondrial uncoupling proteins in the CNS: in support of  $\frac{4}{4}$  function and survival Not Boy I  $\frac{5}{5}$  and  $\frac{1}{5}$  a
- $6$  [62] K.D. Garlid, M. Jabůrek, P. Jezek, The mechanism of proton transport mediated by mitochondrial<br>438 (1998) 10–14. https://doi.org/10.1016/s0014-5793(98)01246 uncoupling proteins, FEBS Lett. 438 (1998) 10-14. https://doi.org/10.1016/s0014-5793(98)01246-0.
- 8 [63] E. Pebay-Peyroula, C. Dahout-Gonzalez, R. Kahn, V. Trézéguet, G.J.-M. Lauquin, G. Brandolin, 9 Structure of mitochondrial ADP/ATP carrier in complex with carboxyatractyloside, Nature. 426  $10$  (2003) 39–44. https://doi.org/10.1038/nature02056.
- $11$  [64] M Klinesshops 11604 Assal  $12$  [04] IVI. Nilligenberg, OCPT - A sopi 13 https://doi.org/10.1016/j.biochi.2016.10.012.
- [65] A. Pfeifer, L.S. Hoffmann, Brown, beige, and white: the new color code of fat and its pharmacological<br>15 magnetications Annu Bey Pharmacol Toxicol, 55 (2015) 207-227 https://doi.org/10.1146/annurey. implications, Annu Rev Pharmacol Toxicol. 55 (2015) 207-227. https://doi.org/10.1146/annurev-16 pharmtox-010814-124346.
- 17 FCCL M Klingenhers C.C. Huggs C  $18$  [OU] IVI. Killigerinerg, J.O. Hualig, J  $\frac{10}{19}$  tissue, Biochim Biophys Acta. 1415 (1999) 271–296. https://doi.org/10.1016/s0005-2736(98)00232-20 **6. 6.**
- 21 [67] A. Fedorenko, P.V. Lishko, Y. Kirichok, Mechanism of fatty-acid-dependent UCP1 uncoupling in 22 brown fat mitochondria, Cell. 151 (2012) 400-413. https://doi.org/10.1016/j.cell.2012.09.010.
- 23  $[60]$  K Yuo D Wu V Wang V 7h  $24$  [bo] K. Auc, D. Wu, T. Wung, T. Znd  $\frac{25}{25}$  mitochondrial calcium uniporter engages UCP1 to form a thermoporter that promotes 26 thermogenesis, Cell Metab. 34 (2022) 1325-1341.e6. https://doi.org/10.1016/j.cmet.2022.07.011.
- 27 [69] J.J. Ruprecht, M.S. King, T. Zögg, A.A. Aleksandrova, E. Pardon, P.G. Crichton, J. Steyaert, E.R.S. Kunji, <sup>28</sup> The Molecular Mechanism of Transport by the Mitochondrial ADP/ATP Carrier, Cell. 176 (2019) 435-29 447.e15. https://doi.org/10.1016/j.cell.2018.11.025.
- 30 117.013. https://www.org/10.12  $31$  [70] N. DIUSLOVELSKY, IVI. KIIIIgeline 32 free fatty acids, which is further stimulated by mersalyl, J Biol Chem. 269 (1994) 27329–27336.
- 33 [71] A.M. Bertholet, A.M. Natale, P. Bisignano, J. Suzuki, A. Fedorenko, J. Hamilton, T. Brustovetsky, L. 34 Kazak, R. Garrity, E.T. Chouchani, N. Brustovetsky, M. Grabe, Y. Kirichok, Mitochondrial uncouplers  $\frac{35}{25}$  induce proton leak by activating AAC and UCP1, Nature. 606 (2022) 180–187. 36 https://doi.org/10.1029/c415
- $37$   $100.018$   $10.1030$   $3413$  $\frac{3}{38}$  [72] M. Klingenberg, The ADP and ATP transport in mitochondria and its carrier, Biochim Biophys Acta. 39 1778 (2008) 1978-2021. https://doi.org/10.1016/j.bbamem.2008.04.011.
- 40 [73] T. Hoang, M.D. Smith, M. Jelokhani-Niaraki, Toward understanding the mechanism of ion transport activity of neuronal uncoupling proteins UCP2, UCP4, and UCP5, Biochemistry. 51 (2012) 4004-4014.  $\frac{42}{42}$  https://doi.org/10.1021/hi300
- $43$   $[74]$   $R$   $7$   $7$   $6$   $1$   $7$   $1$   $7$  $\frac{12}{44}$  [74] R.-Z. Zhao, S. Jiang, L. Zhang, Z.-B. Yu, Mitochondrial electron transport chain, ROS generation and 45 uncoupling, Int J Mol Med. 44 (2019) 3-15. https://doi.org/10.3892/ijmm.2019.4188.
- 46 [75] T.C. Esteves, M.D. Brand, The reactions catalysed by the mitochondrial uncoupling proteins UCP2 and UCP3, Biochim Biophys Acta. 1709 (2005) 35-44. https://doi.org/10.1016/j.bbabio.2005.06.002.
- 18 [76] M.D. Brand, T.C. Esteves, Physiological functions of the mitochondrial uncoupling proteins UCP2 and  $\frac{49}{1000}$   $\frac{1000}{24}$   $\frac{611}{24}$   $\frac{1000}{2000}$  $50$  OCF3, Cell Ivietab.  $2(2003)$  os
- $\frac{51}{51}$  [77] J. Nedergaard, B. Cannon, The "novel" "uncoupling" proteins UCP2 and UCP3: what do they really 52 do? Pros and cons for suggested functions, Exp Physiol. 88 (2003) 65-84. 53 https://doi.org/10.1113/eph8802502.
- <sup>54</sup> [78] V. Azzu, M.D. Brand, The on-off switches of the mitochondrial uncoupling proteins, Trends in 55 Ricchemical Sciences 25 (201  $56$  Blochemical Sciences. 33 (201
- $\frac{8}{57}$  [79] A.J. Vidal-Puig, D. Grujic, C.Y. Zhang, T. Hagen, O. Boss, Y. Ido, A. Szczepanik, J. Wade, V. Mootha, R. 58 Cortright, D.M. Muoio, B.B. Lowell, Energy metabolism in uncoupling protein 3 gene knockout mice, 59 J Biol Chem. 275 (2000) 16258-16266. https://doi.org/10.1074/jbc.M910179199.
- $60$  [80] E.M. Mills, M.L. Banks, J.E. Sprague, T. Finkel, Pharmacology: uncoupling the agony from ecstasy,  $61$  Nature 426 (2003) 403-404  $62$  Mature:  $420 (2005)$   $405$   $404$ .

- $[81]$ P. Schrauwen, M.K.C. Hesselink, The role of uncoupling protein 3 in fatty acid metabolism: 1 protection against lipotoxicity?, Proc Nutr Soc. 63 (2004) 287-292. 2 https://doi.org/10.1079/PNS2003336.
- $\frac{3}{3}$  [82] A. Vozza, G. Parisi, F. De Leonardis, F.M. Lasorsa, A. Castegna, D. Amorese, R. Marmo, V.M. 4 Colognile L. Delmieri, D. Biogu  $5$  Calcagnite, L. Familien, D. Nicqu  $6$  UCP2 transports C4 metabolites out of mitochondria, regulating glucose and glutamine oxidation, 7 Proc Natl Acad Sci U S A. 111 (2014) 960-965. https://doi.org/10.1073/pnas.1317400111.
- 8 [83] T.A. Schiffer, L. Löf, R. Gallini, M. Kamali-Moghaddam, M. Carlström, F. Palm, Mitochondrial 866590. Respiration-Dependent ANT2-UCP2 Interaction, Front Physiol. 13 (2022) 866590.  $10$  https://doi.org/10.3389/fphys.2022.866590.
- $11$  [0.4] C Next A Perfect C Alti- $12$  [64] S. Nesci, A. Pagilarani, C. Aigie 13 functions revealed by the membrane-embedded FO structure, Crit. Rev. Biochem. Mol. Biol. 55 14 (2020) 309–321. https://doi.org/10.1080/10409238.2020.1784084.<br>15 1851 - Kazak LE Bahbani B Samborska G 7 Lu M B ledrychowski M.
- [85] L. Kazak, J.F. Rahbani, B. Samborska, G.Z. Lu, M.P. Jedrychowski, M. Lajoie, S. Zhang, L. Ramsay, F.Y. Dou, D. Tenen, E.T. Chouchani, P. Dzeja, I.R. Watson, L. Tsai, E.D. Rosen, B.M. Spiegelman, Ablation 17 **Superior** *Seconds exercise treasure* 18 **Draupocyte creating transpo** 1<sub>9</sub><br>19 1 (2019) 360–370. https://doi.org/10.1038/s42255-019-0035-x.<br>20 [86] J.F. Rahbani, C. Scholtes, D.M. Lagarde, M.F. Hussain, A. Roesler.
- 20 [86] J.F. Rahbani, C. Scholtes, D.M. Lagarde, M.F. Hussain, A. Roesler, C.B. Dykstra, J. Bunk, B. Samborska, 21 S.L. O'Brien, E. Tripp, A. Pacis, A.R. Angueira, O.S. Johansen, J. Cinkornpumin, I. Hossain, M.D. Lynes, 22<br>22 Y. Zhang, A.P. White, W.A. Pastor, M. Chondronikola, L. Sidossis, S. Klein, A. Kralli, A.M. Cypess, S.B. 23 Dedecen N Jessen V H Tre 24 reaction, m. sessen, n. n. ise  $25$  ADRA1A–Gαq signalling potentiates adipocyte thermogenesis through CKB and TNAP, Nat Metab. 4 26 (2022) 1459-1473. https://doi.org/10.1038/s42255-022-00667-w.
- 27 [87] L. Kazak, E.T. Chouchani, G.Z. Lu, M.P. Jedrychowski, C.J. Bare, A.I. Mina, M. Kumari, S. Zhang, I. <sup>28</sup> Vuckovic, D. Laznik-Bogoslavski, P. Dzeja, A.S. Banks, E.D. Rosen, B.M. Spiegelman, Genetic Depletion 29 of Adipocyte Creatine Metabolism Inhibits Diet-Induced Thermogenesis and Drives Obesity, Cell 30 **Matchelling 26 (2017) CCO CT**  $31$  IVIELADOIISIII. 20 $(2017)$ 000-07
- $32$  [88] D.T. Sawyer, J.S. Valentine, How super is superoxide?, Acc. Chem. Res. 14 (1981) 393-400. 33 https://doi.org/10.1021/ar00072a005.
- $34$  [89] M.P. Murphy, How mitochondria produce reactive oxygen species, Biochem J. 417 (2009) 1–13.  $\frac{35}{25}$  https://doi.org/10.1042/BJ20081386.
- $36$   $1001$  AV Androvev V.F. Kushnarov  $37$  [JO] A.I. Andreyev, i.e. Nushinarev
- 38 Biochemistry Mosc. 70 (2005) 200–214.<br>39 [91] J. Hirst, M.S. King, K.R. Pryde, The produ 39 [91] J. Hirst, M.S. King, K.R. Pryde, The production of reactive oxygen species by complex I, Biochem Soc 40 Trans. 36 (2008) 976-980. https://doi.org/10.1042/BST0360976.
- [92] B.B. Seo, M. Marella, T. Yagi, A. Matsuno-Yagi, The single subunit NADH dehydrogenase reduces 42 generation of reactive exygen 43 **SCHEMION OF REGISTER**
- <sup>13</sup><br>44 https://doi.org/10.1016/j.febslet.2006.10.008.<br>45 [93] E.T. Chouchani, V.R. Pell, E. Gaude, D. Aksentije E.T. Chouchani, V.R. Pell, E. Gaude, D. Aksentijević, S.Y. Sundier, E.L. Robb, A. Logan, S.M. Nadtochiy, 46 E.N.J. Ord, A.C. Smith, F. Eyassu, R. Shirley, C.-H. Hu, A.J. Dare, A.M. James, S. Rogatti, R.C. Hartley, S. 47 Eaton, A.S.H. Costa, P.S. Brookes, S.M. Davidson, M.R. Duchen, K. Saeb-Parsy, M.J. Shattock, A.J. Robinson, L.M. Work, C. Frezza, T. Krieg, M.P. Murphy, Ischaemic accumulation of succinate controls 49 nonostroion inium through ni  $\frac{1}{20}$  repertusion injury unough  $\frac{1}{20}$ 51 https://doi.org/10.1038/nature13909.<br>52 [94] S.S. Korshunov, V.P. Skulachev, A.A. Sta
- 52 [94] S.S. Korshunov, V.P. Skulachev, A.A. Starkov, High protonic potential actuates a mechanism of 53 production of reactive oxygen species in mitochondria, FEBS Lett. 416 (1997) 15-18. 54 https://doi.org/10.1016/s0014-5793(97)01159-9.
- 55 [OF] D Fate C Bergemini S Leoni  $56$  [33] K. Tato, C. Defgamin, S. Leoni  $\frac{57}{57}$  mitochondrial complex I: implications in neurodegeneration, Neurochem Res. 33 (2008) 2487–2501. 58 https://doi.org/10.1007/s11064-008-9747-0.
- 59 [96] E.L. Robb, A.R. Hall, T.A. Prime, S. Eaton, M. Szibor, C. Viscomi, A.M. James, M.P. Murphy, Control of 60 mitochondrial superoxide production by reverse electron transport at complex I, J Biol Chem. 293  $61$  (2018) Q860-0870 https://do  $62$  (2010) 5005 5015.  $m_{\rm p}$
- 63
- 64 65
- $[97]$ G. Lenaz, M.L. Genova, Structure and organization of mitochondrial respiratory complexes: a new 1 understanding of an old subject, Antioxid Redox Signal. 12 (2010) 961-1008. 2 https://doi.org/10.1089/ars.2009.2704.
- $\frac{3}{3}$  [98] A.J. Lambert, M.D. Brand, Superoxide production by NADH:ubiquinone oxidoreductase (complex I)  $\frac{4}{4}$  depends on the pH gradient as  $5$  adjacent according to the programmation of  $\frac{5}{2}$
- $6$  517. https://doi.org/10.1042/BJ20040485.<br>7 [99] C.C. Moser, T.A. Farid, S.E. Chobot, P.L. Dut [99] C.C. Moser, T.A. Farid, S.E. Chobot, P.L. Dutton, Electron tunneling chains of mitochondria,<br>8 Biochimica et Biophysica Acta (BBA) - Bioenergetics, 1757 (2006) 1096–1109. Biochimica et Biophysica Acta (BBA) - Bioenergetics. 1757 (2006) 1096-1109. <sup>9</sup> https://doi.org/10.1016/j.bbabio.2006.04.015.
- $10$  [100] CC Page CC Moser Y Cher  $11$   $1200$   $1200$   $1200$   $1200$   $1200$   $1200$   $1200$   $1200$   $1200$   $1200$   $1200$  $\frac{1}{12}$  biological oxidation-reduction, Nature. 402 (1999) 47–52. https://doi.org/10.1038/46972.
- 13 [101] M.D. Brand, C. Affourtit, T.C. Esteves, K. Green, A.J. Lambert, S. Miwa, J.L. Pakay, N. Parker, Mitochondrial superoxide: production, biological effects, and activation of uncoupling proteins, Free<br>15 Badic Biol Med. 37 (2004) 755–767, https://doi.org/10.1016/i freeradbiomed.2004 05.034 Radic Biol Med. 37 (2004) 755-767. https://doi.org/10.1016/j.freeradbiomed.2004.05.034.
- 16 [102] E.A. Malingriaux, A. Rupprecht, L. Gille, O. Jovanovic, P. Jezek, M. Jaburek, E.E. Pohl, Fatty acids are 17 Levin Abudrous 3 noncondum  $18$  Rey III 4-Hydroxy-z-Hollendi-H  $19$  e77786. https://doi.org/10.1371/journal.pone.0077786.
- 20 [103] K.S. Echtay, T.C. Esteves, J.L. Pakay, M.B. Jekabsons, A.J. Lambert, M. Portero-Otín, R. Pamplona, A.J. 21 Vidal-Puig, S. Wang, S.J. Roebuck, M.D. Brand, A signalling role for 4-hydroxy-2-nonenal in regulation  $^{22}$  of mitochondrial uncoupling, EMBO J. 22 (2003) 4103-4110. https://doi.org/10.1093/emboj/cdg412.
- $23$  [104] B Cannon LG Shabalina TV 24  $[10 + 1]$  D. Carmon, i.o. Shabanna, i.v  $\frac{25}{25}$  protection against reactive oxygen species--or not?, Biochim Biophys Acta. 1757 (2006) 449–458. 26 https://doi.org/10.1016/j.bbabio.2006.05.016.
- 27 [105] K.S. Echtay, D. Roussel, J. St-Pierre, M.B. Jekabsons, S. Cadenas, J.A. Stuart, J.A. Harper, S.J. Roebuck, 28 A. Morrison, S. Pickering, J.C. Clapham, M.D. Brand, Superoxide activates mitochondrial uncoupling 29<br>proteins, Nature. 415 (2002) 96–99. https://doi.org/10.1038/415096a.
- $30$  Proteins, Matthe. 123 (2002)  $31$  [100] L.J. Tollite, M.D. Brand, Uncor 32 mitochondria, Free Radic Biol Med. 49 (2010) 606-611. 33 https://doi.org/10.1016/j.freeradbiomed.2010.05.010.
- $34$  [107] G. Lenaz, M.L. Genova, Structural and functional organization of the mitochondrial respiratory chain:  $\frac{35}{2}$  a dynamic super-assembly, Int J Biochem Cell Biol. 41 (2009) 1750–1772.  $36$  a dynamic super assembly,  $\ldots$  $37$  mups.//wor.org/10.1010/j.blot
- $\frac{3}{38}$  [108] S. Nesci, G. Lenaz, The mitochondrial energy conversion involves cytochrome c diffusion into the 39 respiratory supercomplexes, Biochim Biophys Acta Bioenerg. 1862 (2021) 148394. 40 https://doi.org/10.1016/j.bbabio.2021.148394.
- $\frac{41}{12}$  [109] J.A. Letts, L.A. Sazanov, Clarifying the supercomplex: the higher-order organization of the 42 mitochondrial oloctron transp 43 mww.nomanarcicci.on transp  $\frac{13}{44}$  https://doi.org/10.1038/nsmb.3460.<br>45 [110] I. Vercellino, L.A. Sazanov, The assem
- [110] I. Vercellino, L.A. Sazanov, The assembly, regulation and function of the mitochondrial respiratory 46 chain, Nat Rev Mol Cell Biol. (2021). https://doi.org/10.1038/s41580-021-00415-0.
- 47 [111] C. Piccoli, R. Scrima, D. Boffoli, N. Capitanio, Control by cytochrome c oxidase of the cellular oxidative phosphorylation system depends on the mitochondrial energy state, Biochem J. 396 (2006)  $49$   $572, 592$   $545$   $542$   $542$  $\frac{50}{20}$  375-505.  $\frac{1}{20}$  375-61.  $\frac{1}{20}$
- $51$  [112] S. Nesci, C. Algieri, F. Trombetti, M. Fabbri, G. Lenaz, Two separate pathways underlie NADH and 52 succinate oxidation in swine heart mitochondria: Kinetic evidence on the mobile electron carriers, 53 Biochimica et Biophysica Acta (BBA) - Bioenergetics. (2023) 148977. 54 https://doi.org/10.1016/j.bbabio.2023.148977.
- 55 [112] C Noce F Trembetti A Doct  $56$  [115] 5. Nescrit From betti, A. Fagu  $\frac{57}{57}$  Supramolecular Structure of the Mitochondrial Oxidative Phosphorylation System: Implications for 58 Pathology, Life. 11 (2021) 242. https://doi.org/10.3390/life11030242.<br>59

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intermembrane space







 $\Delta\Psi$  (mV)

Proton current (nmol  $H_{+}$ , nin  $_{1}$ , ng<sup>1</sup>