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Mitochondrial respiratory states and rates: Building blocks of mitochondrial physiology Part 1

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49 Patel HH, Pavlova N, Pecina P, Pelnena D, Pereira da Silva Grilo da Silva F, Perez Valencia JA, Pesta D, Petit PX, Pettersen IKN, Pichaud N, Piel S, Pietka TA, Pino MF, Pirkmajer S, Porter C, Porter RK, 50 Pranger F, Prochownik EV, Pulinilkunnil T, Puskarich MA, Puurand M, Quijano C, Radenkovic F, 51 52 Radi R, Ramzan R, Rattan S, Reboredo P, Rich PR, Renner-Sattler K, Rial E, Robinson MM, Roden 53 M, Rodríguez-Enriquez S, Rohlena J, Rolo AP, Ropelle ER, Røsland GV, Rossignol R, Rossiter HB, 54 Rubelj I, Rybacka-Mossakowska J, Saada A, Safaei Z, Salin K, Salvadego D, Sandi C, Sanz A, 55 Sazanov LA, Scatena R, Schartner M, Scheibye-Knudsen M, Schilling JM, Schlattner U, Schönfeld P, 56 Schots P, Schulz R, Schwarzer C, Scott GR, Shabalina IG, Sharma P, Sharma V, Shevchuk I, Siewiera K, Silber AM, Silva AM, Sims CA, Singer D, Skolik R, Smenes BT, Smith J, Soares FAA, Sobotka 57 58 O, Sokolova I, Sonkar VK, Sowton AP, Sparagna GC, Sparks LM, Spinazzi M, Stankova P, Starr J, Stary C, Stelfa G, Stiban J, Stier A, Stocker R, Storder J, Sumbalova Z, Suravajhala P, Svalbe B, 59 60 Swerdlow RH, Swiniuch D, Szabo I, Szewczyk A, Szibor M, Tanaka M, Tandler B, Tarnopolsky MA, Tausan D, Tavernarakis N, Tepp K, Thakkar H, Thyfault JP, Tomar D, Torp MK, Towheed A, Tretter 61 62 L, Trifunovic A, Trivigno C, Tronstad KJ, Trougakos IP, Truu L, Tuncay E, Turan B, Tyrrell DJ, 63 Urban T, Valentine JM, Vella J, Vendelin M, Vercesi AE, Victor VM, Vieira Ligo Teixeira C, Viel C, 64 Vieyra A, Vilks K, Villena JA, Vincent V, Vinogradov AD, Viscomi C, Vitorino RMP, Vogt S, Volani C, Volska K, Votion DM, Vujacic-Mirski K, Wagner BA, Ward ML, Warnsmann V, 65 66 Wasserman DH, Watala C, Wei YH, Wickert A, Wieckowski MR, Wiesner RJ, Williams C, Winwood-Smith H, Wohlgemuth SE, Wohlwend M, Wolff J, Wüst RCI, Yokota T, Zablocki K, 67 Zaugg K, Zaugg M, Zdrazilova L, Zhang Y, Zhang YZ, Zíková A, Zischka H, Zorzano A, Zvejniece L 68 69 70 **Updates and discussion:**

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132 **Abstract** As the knowledge base and importance of mitochondrial physiology to human health expands, 133 the necessity for harmonizing the terminology concerning mitochondrial respiratory states and rates has become increasingly apparent. The chemiosmotic theory establishes the mechanism of energy 134 135 transformation and coupling in oxidative phosphorylation. The unifying concept of the protonmotive 136 force provides the framework for developing a consistent theoretical foundation of mitochondrial physiology and bioenergetics. We follow IUPAC guidelines on terminology in physical chemistry, 137 extended by considerations on open systems and thermodynamics of irreversible processes. The 138 139 concept-driven constructive terminology incorporates the meaning of each quantity and aligns concepts 140 and symbols to the nomenclature of classical bioenergetics. We endeavour to provide a balanced view 141 on mitochondrial respiratory control and a critical discussion on reporting data of mitochondrial 142 respiration in terms of metabolic flows and fluxes. Uniform standards for evaluation of respiratory states 143 and rates will ultimately contribute to reproducibility between laboratories and thus support the 144 development of databases of mitochondrial respiratory function in species, tissues, and cells. Clarity of 145 concept and consistency of nomenclature facilitate effective transdisciplinary communication, 146 education, and ultimately further discovery.

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Keywords: Mitochondrial respiratory control, coupling control, mitochondrial preparations,
protonmotive force, uncoupling, oxidative phosphorylation, OXPHOS, efficiency, electron transfer, ET;
electron transfer system, ETS; proton leak, LEAK, residual oxygen consumption, ROX, State 2, State
3, State 4, normalization, flow, flux, O₂

153 Executive summary

155 In view of the broad implications for health care, mitochondrial researchers face an increasing 156 responsibility to disseminate their fundamental knowledge and novel discoveries to a wide range of 157 stakeholders and scientists beyond the group of specialists. This requires implementation of a commonly 158 accepted terminology within the discipline and standardization in the translational context. Authors, 159 reviewers, journal editors, and lecturers are challenged to collaborate with the aim to harmonize the 160 nomenclature in the growing field of mitochondrial physiology and bioenergetics, from evolutionary biology and comparative physiology to mitochondrial medicine. In the present communication we focus 161 162 on the following concepts in mitochondrial physiology:

- 1. Aerobic respiration depends on the coupling of phosphorylation (ADP \rightarrow ATP) to O₂ flux in 163 catabolic reactions. Coupling in oxidative phosphorylation is mediated by the translocation of 164 165 protons across the inner mitochondrial membrane through proton pumps generating or 166 utilizing the protonmotive force, that is maintained between the mitochondrial matrix and intermembrane compartment or outer mitochondrial space. Compartmental coupling 167 168 distinguishes this vectorial component of oxidative phosphorylation from glycolytic fermentation as the counterpart of cellular core energy metabolism (Figure 1). Cell respiration 169 is distinguished from fermentation: (1) Electron acceptors are supplied by external respiration 170 171 for the maintenance of redox balance, whereas fermentation is characterized by an internal electron acceptor produced in intermediary metabolism. In aerobic cell respiration, redox 172 173 balance is maintained by O_2 as the electron acceptor. (2) Compartmental coupling in vectorial oxidative phosphorylation contrasts to exclusively scalar substrate-level phosphorylation in 174 fermentation. 175
- 2. When measuring mitochondrial metabolism, the contribution of fermentation and other cytosolic 176 177 interactions must be excluded from analysis by disrupting the barrier function of the plasma membrane. Selective removal or permeabilization of the plasma membrane yields 178 179 mitochondrial preparations-including isolated mitochondria, tissue and cellular 180 preparations—with structural and functional integrity. Subsequently, extra-mitochondrial 181 concentrations of fuel substrates, ADP, ATP, inorganic phosphate, and cations including H⁺ 182 can be controlled to determine mitochondrial function under a set of conditions defined as coupling control states. We strive to incorporate an easily recognized and understood, concept-183 184 driven terminology of bioenergetics with explicit terms and symbols that define the nature of respiratory states. 185
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 3. Mitochondrial coupling states are defined according to the control of respiratory oxygen flux by
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 the protonmotive force. Capacities of oxidative phosphorylation and electron transfer are

measured at kinetically saturating concentrations of fuel substrates, ADP and inorganic phosphate, and O_2 , or at optimal uncoupler concentrations, respectively, in the absence of Complex IV inhibitors such as NO, CO, or H₂S. Respiratory capacity is a measure of the upper bound of the rate of respiration; it depends on the substrate type undergoing oxidation, and

provides reference values for the diagnosis of health and disease, and for evaluation of the effects of Evolutionary background, Age, Gender and sex, Lifestyle and Environment.

195 Figure 1. Internal and external respiration

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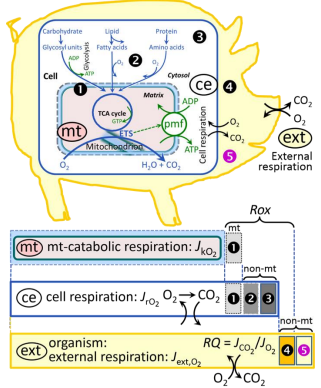
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196 Mitochondrial respiration is the oxidation of fuel 197 substrates (electron donors) and reduction of O₂ catalysed by the electron transfer system, ETS: 198 199 (mt) mitochondrial catabolic respiration; (ce) 200 total cellular O₂ consumption; and (ext) external 201 respiration. All chemical reactions, r, that 202 consume O_2 in the cells of an organism, 203 contribute to cell respiration, J_{rO2} . In addition to 204 mitochondrial catabolic respiration, O_2 is 205 consumed by:

206 \bullet Mitochondrial residual O₂ consumption, *Rox*. 207 0 Non-mitochondrial O₂ consumption by catabolic reactions, particularly peroxisomal 208 209 oxidases and microsomal cytochrome P450 210 systems. ³ Non-mitochondrial *Rox* by reactions 211 unrelated to catabolism. 0 Extracellular *Rox*. 0212 Aerobic microbial respiration. Bars are not at a 213 quantitative scale.

- 214 (mt) **Mitochondrial catabolic respiration**, J_{kO2} , 215 is the O₂ consumption by the mitochondrial 216 ETS excluding *Rox*.
- 217 (ce) Cell respiration, J_{rO2} , takes into account



internal O₂-consuming reactions, r, including catabolic respiration and *Rox*. Catabolic cell respiration is the O₂ consumption associated with catabolic pathways in the cell, including mitochondrial catabolism in addition to peroxisomal and microsomal oxidation reactions (2).

- 221 (ext) External respiration balances internal respiration at steady-state, including extracellular Rox (④) and aerobic respiration by the microbiome (\mathbf{S}). O₂ is transported from the environment across the 222 223 respiratory cascade, *i.e.*, circulation between tissues and diffusion across cell membranes, to the 224 intracellular compartment. The respiratory quotient, RQ, is the molar CO_2/O_2 exchange ratio; when 225 combined with the respiratory nitrogen quotient, N/O_2 (mol N given off per mol O_2 consumed), the 226 RQ reflects the proportion of carbohydrate, lipid and protein utilized in cell respiration during 227 aerobically balanced steady-states. Bicarbonate and CO_2 are transported in reverse to the 228 extracellular mileu and the organismic environment. Hemoglobin provides the molecular paradigm 229 for the combination of O_2 and CO_2 exchange, as do lungs and gills on the morphological level. 230
- 231 4. Incomplete tightness of coupling, *i.e.*, some degree of uncoupling relative to the substratedependent coupling stoichiometry, is a characteristic of energy-transformations across 232 membranes. Uncoupling is caused by a variety of physiological, pathological, toxicological, 233 234 pharmacological and environmental conditions that exert an influence not only on the proton leak and cation cycling, but also on proton slip within the proton pumps and the structural 235 236 integrity of the mitochondria. A more loosely coupled state is induced by stimulation of mitochondrial superoxide formation and the bypass of proton pumps. In addition, the use of 237 238 protonophores represents an experimental uncoupling intervention to asses the transition from 239 a well-coupled to a noncoupled state of mitochondrial respiration.
- 5. Respiratory oxygen consumption rates have to be carefully normalized to enable meta-analytic
 studies beyond the question of a particular experiment. Therefore, all raw data on rates and
 variables for normalization should be published in an open access data repository.
 Normalization of rates for: (1) the number of objects (cells, organisms); (2) the volume or

mass of the experimental sample; and (3) the concentration of mitochondrial markers in the experimental chamber are sample-specific normalizations, which are distinguished from system-specific normalization for the volume of the chamber (the measuring system).

- 6. The consistent use of terms and symbols will facilitate transdisciplinary communication and support the further development of a collaborative database on bioenergetics and mitochondrial physiology. The present considerations are focused on studies with mitochondrial preparations. These will be extended in a series of reports on pathway control of mitochondrial respiration, respiratory states in intact cells, and harmonization of experimental procedures.
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Box 1: In brief – Mitochondria and Bioblasts

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'For the physiologist, mitochondria afforded the first opportunity for an experimental approach to structure-function relationships, in particular those involved in active transport, vectorial metabolism, and metabolic control mechanisms on a subcellular level' (Ernster and Schatz 1981).

Mitochondria are oxygen-consuming electrochemical generators that evolved from the endosymbiotic
alphaproteobacteria which integrated into a host cell related to Asgard Archaea (Margulis 1970; Lane
2005; Roger *et al.* 2017). They were described by Richard Altmann (1894) as 'bioblasts', which include
not only the mitochondria as presently defined, but also symbiotic and free-living bacteria. The word
'mitochondria' (Greek mitos: thread; chondros: granule) was introduced by Carl Benda (1898).

264 Contrary to current textbook dogma, mitochondria form dynamic networks within eukaryotic 265 cells. Mitochondrial movement is supported by microtubules and morphology can change in response 266 to energy requirements of the cell via processes known as fusion and fission; these interactions allow mitochondria to communicate within a network (Chan 2006). Mitochondria can even traverse cell 267 268 boundaries in a process known as horizontal mitochondrial transfer (Torralba et al. 2016). Another 269 defining characteristic of mitochondria is the double membrane. The mitochondrial inner membrane 270 (mtIM) forms dynamic tubular to disk-shaped cristae that separate the mitochondrial matrix, *i.e.*, the negatively charged internal mitochondrial compartment, from the intermembrane space; the latter being 271 272 enclosed by the mitochondrial outer membrane (mtOM) and positively charged with respect to the 273 matrix. The mtIM contains the non-bilayer phospholipid cardiolipin, which is not present in any other 274 eukaryotic cellular membrane. Cardiolipin has many regulatory functions (Oemer et al. 2018); in particular, it stabilizes and promotes the formation of respiratory supercomplexes (SC I_nIII_nIV_n), which 275 276 are supramolecular assemblies based upon specific and dynamic interactions between individual 277 respiratory complexes (Greggio et al. 2017; Lenaz et al. 2017). The fluidity of the mitochondrial 278 membrane is plastic and exerts an influence on the functional properties of proteins incorporated in 279 membranes (Waczulikova et al. 2007). Intracellular stress factors may cause shrinking or swelling of 280 the mitochondrial matrix, that can ultimately result in permeability transition.

Mitochondria are the structural and functional elementary components of cell respiration. 281 282 Mitochondrial respiration is the reduction of molecular oxygen by electron transfer coupled to 283 electrochemical proton translocation across the mtIM. In the process of oxidative phosphorylation 284 (OXPHOS), the catabolic reaction of oxygen consumption is electrochemically coupled to the transformation of energy in the form of adenosine triphosphate (ATP; Mitchell 1961, 2011). 285 286 Mitochondria are the powerhouses of the cell which contain the machinery of the OXPHOS-pathways, 287 including transmembrane respiratory complexes (proton pumps with FMN, Fe-S and cytochrome b, c, 288 *aa*₃ redox systems); alternative dehydrogenases and oxidases; the coenzyme ubiquinone (Q); F-ATPase 289 or ATP synthase; the enzymes of the tricarboxylic acid cycle (TCA), fatty acid and amino acid oxidation; 290 transporters of ions, metabolites and co-factors; iron/sulphur cluster synthesis; and mitochondrial 291 kinases related to catabolic pathways. The mitochondrial proteome comprises over 1,200 proteins 292 (Calvo et al. 2015; 2017), mostly encoded by nuclear DNA (nDNA), with a variety of functions, many 293 of which are relatively well known (e.g., proteins regulating mitochondrial biogenesis or apoptosis), while others are still under investigation, or need to be identified (e.g., permeability transition pore, 294 295 alanine transporter). Only recently has it been possible to use the mammalian mitochondrial proteome 296 to discover and characterize the genetic basis of mitochondrial diseases (Williams et al. 2016; Palmfeldt 297 and Bross 2017).

Numerous cellular processes are orchestrated by a constant crosstalk between mitochondria andother cellular components. For example, the crosstalk between mitochondria and the endoplasmic

300 reticulum is involved in the regulation of calcium homeostasis, cell division, autophagy, differentiation, 301 and anti-viral signaling (Murley and Nunnari 2016). Mitochondria contribute to the formation of peroxisomes, which are hybrids of mitochondrial and ER-derived precursors (Sugiura et al. 2017). 302 303 Cellular mitochondrial homeostasis (mitostasis) is maintained through regulation at transcriptional, 304 post-translational and epigenetic levels. Cell signalling modules contribute to homeostatic regulation 305 throughout the cell cycle or even cell death by activating proteostatic modules (e.g., the ubiquitinproteasome and autophagy-lysosome/vacuole pathways; specific proteases like LON) and genome 306 307 stability modules in response to varying energy demands and stress cues (Quiros et al. 2016). Several 308 post-translational modifications, including acetylation and bitrosylation, are also capable of influencing 309 the bioenergetic response, with clinically significant implications for health and disease (Carrico et al. 310 2018).

311 Mitochondria of higher eukaryotes typically maintain several copies of their own circular genome 312 known as mitochondrial DNA (mtDNA; hundred to thousands per cell; Cummins 1998), which is 313 maternally inherited in humans. Biparental mitochondrial inheritance is documented in mammals, birds, 314 fish, reptiles and invertebrate groups, and is even the norm in some bivalve taxonomic groups (Breton 315 et al. 2007; White et al. 2008). The mitochondrial genome of the angiosperm Amborella contains a 316 record of six mitochondrial genome equivalents aquired by horizontal transfer of entire genomes, two 317 from angiosperms, three from algae and one from mosses (Rice et al. 2016). In unicellular organisms (*i.e.*, protists) the structural organization of mitochondrial genomes is highly variable and includes 318 319 circular and linear DNA (Zikova et al. 2016). While some of the free-living flagellates exhibit the largest 320 known gene coding capacity (e.g. jakobid Andalucia godoyi mitochondrial DNA codes for 106 genes; 321 Burger et al. 2013), some protist groups (e.g. alveolates) possess mitochondrial genomes with only three protein-coding genes and two rRNAs (Feagin et al. 2012). The complete loss of mitochondrial genome 322 323 is observed in highly reduced mitochondria of Cryptosporidium species (Liu et al. 2016). Reaching the final extreme, the microbial eukaryote, oxymonad Monocercomonoides, has no mitochondrion 324 325 whatsoever and lacks all typical nuclear-encoded mitochondrial proteins demonstrating that while in 326 99% of organisms mitochondria play a vital role, this organelle is not indispensable (Karnkowska et al. 327 2016).

328 In vertebrates but not all invertebrates, mtDNA is compact (16.5 kB in humans) and encodes 13 329 protein subunits of the transmembrane respiratory Complexes CI, CIII, CIV and ATP synthase (F-330 ATPase), 22 tRNAs, and two RNAs. Additional gene content has been suggested to include microRNAs, 331 piRNA, smithRNAs, repeat associated RNA, and even additional proteins (Duarte et al. 2014; Lee et 332 al. 2015; Cobb et al. 2016). The mitochondrial genome requires nuclear-encoded mitochondrially 333 targeted proteins, e.g., TFAM, for its maintenance and expression (Rackham et al. 2012). Both genomes 334 encode peptides of the membrane spanning redox pumps (CI, CIII and CIV) and F-ATPase, leading to strong constraints in the coevolution of both genomes (Blier et al. 2001). 335

Given the multiple roles of mitochondria, it is perhaps not surprising that mitochondrial dysfunction is associated with a wide variety of genetic and degenerative diseases. Robust mitochondrial function is supported by physical exercise and caloric balance, and is central for sustained metabolic health throughout life. Therefore, a more consistent set of definitions for mitochondrial physiology will increase our understanding of the etiology of disease and improve the diagnostic repertoire of mitochondrial medicine with a focus on protective medicine, lifestyle and healthy aging.

342 Mitochondrion is singular and mitochondria is plural. Abbreviation: mt, as generally used in343 mtDNA.

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347 **1. Introduction**348

Mitochondria are the powerhouses of the cell with numerous physiological, molecular, and genetic functions (**Box 1**). Every study of mitochondrial health and disease faces Evolution, Age, Gender and sex, Lifestyle, and Environment (MitoEAGLE) as essential background conditions intrinsic to the individual person or cohort, species, tissue and to some extent even cell line. As a large and coordinated group of laboratories and researchers, the mission of the global MitoEAGLE Network is to generate the necessary scale, type, and quality of consistent data sets and conditions to address this intrinsic complexity. Harmonization of experimental protocols and implementation of a quality control and data management system are required to interrelate results gathered across a spectrum of studies and to generate a rigorously monitored database focused on mitochondrial respiratory function. In this way, researchers from a variety of disciplines can compare their findings using clearly defined and accepted international standards.

360 With an emphasis on quality of research, published data can be useful far beyond the specific 361 question of a particular experiment. For example, collaborative data sets support the development of 362 open-access databases such as those for National Institutes of Health sponsored research in genetics, 363 proteomics, and metabolomics. Indeed, enabling meta-analysis is the most economic way of providing robust answers to biological questions (Cooper et al. 2009). However, the reproducibility of quantitative 364 365 results and databases depend on accurate measurements under strictly-defined conditions. Likewise, meaningful interpretation and comparability of experimental outcomes requires standardisation of 366 367 protocols between research groups at different institutes. In addition to quality control, a conceptual 368 framework is also required to standardise and homogenise terminology and methodology. Vague or 369 ambiguous jargon can lead to confusion and may relegate valuable signals to wasteful noise. For this 370 reason, measured values must be expressed in standard units for each parameter used to define 371 mitochondrial respiratory function. A consensus on fundamental nomenclature and conceptual 372 coherence, however, are missing in the expanding field of mitochondrial physiology. To fill this gap, 373 the present communication provides an in-depth review on harmonization of nomenclature and 374 definition of technical terms, which are essential to improve the awareness of the intricate meaning of 375 current and past scientific vocabulary. This is important for documentation and integration into 376 databases in general, and quantitative modelling in particular (Beard 2005).

In this review, we focus on coupling states and fluxes through metabolic pathways of aerobic energy transformation in mitochondrial preparations as a first step in the attempt to generate a conceptually-oriented nomenclature in bioenergetics and mitochondrial physiology. Respiratory control by fuel substrates and specific inhibitors of respiratory enzymes, coupling states of intact cells, and respiratory flux control ratios will be reviewed in subsequent communications, prepared in the frame of COST Action MitoEAGLE open to global bottom-up input.

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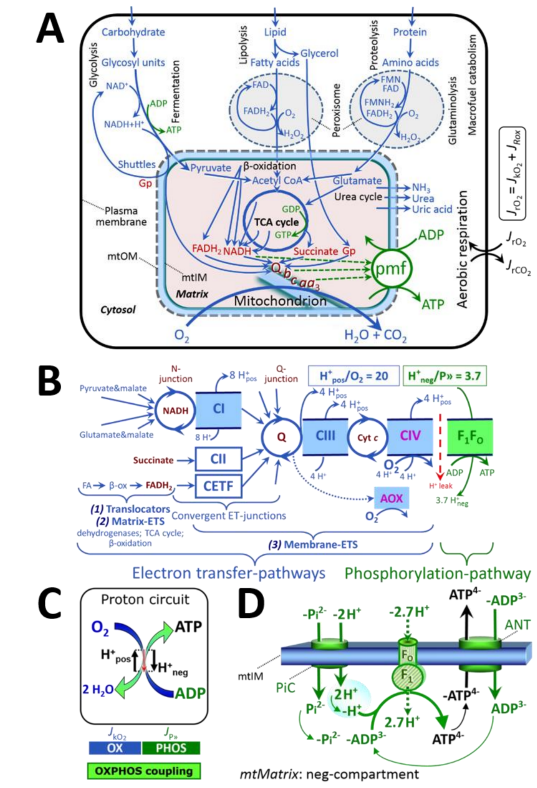
2. Coupling states and rates in mitochondrial preparations

'Every professional group develops its own technical jargon for talking about matters of critical concern ... People who know a word can share that idea with other members of their group, and a shared vocabulary is part of the glue that holds people together and allows them to create a shared culture' (Miller 1991).

391 2.1. Cellular and mitochondrial respiration392

393 2.1.1. Aerobic and anaerobic catabolism and ATP turnover: In respiration, electron transfer 394 is coupled to the phosphorylation of ADP to ATP, with energy transformation mediated by the 395 protonmotive force, pmf (Figure 2). Anabolic reactions are coupled to catabolism, both by ATP as the 396 intermediary energy currency and by small organic precursor molecules as building blocks for 397 biosynthesis. Glycolysis involves substrate-level phosphorylation of ADP to ATP in fermentation 398 without utilization of O₂, studied mainly in intact cells and organisms. Many cellular fuel substrates are 399 catabolized to acetyl-CoA or to glutamate, and further electron transfer reduces nicotinamide adenine 400 dinucleotide to NADH or flavin adenine dinucleotide to FADH₂. Subsequent mitochondrial electron transfer to O2 is coupled to proton translocation for the control of the protonmotive force and 401 phosphorylation of ADP (Figure 2B and 2C). In contrast, extra-mitochondrial oxidation of fatty acids 402 403 and amino acids proceeds partially in peroxisomes without coupling to ATP production: acyl-CoA 404 oxidase catalyzes the oxidation of FADH₂ with electron transfer to O₂; amino acid oxidases oxidize 405 flavin mononucleotide FMNH₂ or FADH₂ (Figure 2A).

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408 Figure 2. Cell respiration and oxidative phosphorylation (OXPHOS) 409 Mitochondrial respiration is the oxidation of fuel substrates (electron donors) with electron 410 transfer to O₂ as the electron acceptor. For explanation of symbols see also Figure 1. 411 (A) Respiration of intact cells: Extra-mitochondrial catabolism of macrofuels and uptake of small molecules by the cell provide the mitochondrial fuel substrates. Dashed arrows 412 indicate the connection between the redox proton pumps (respiratory Complexes CI, CIII 413 and CIV) and the transmembrane protonmotive force, pmf. Coenzyme Q (Q) and the 414 415 cytochromes b, c, and aa_3 are redox systems of the mitochondrial inner membrane, mtIM.

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Glycerol-3-phosphate, Gp.

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(B) Respiration in mitochondrial preparations: The mitochondrial electron transfer system 417 418 (ETS) is (1) fuelled by diffusion and transport of substrates across the mtOM and mtIM, 419 and in addition consists of the (2) matrix-ETS, and (3) membrane-ETS. Electron transfer 420 converges at the N-junction, and from CI, CII and electron transferring flavoprotein 421 complex (CETF) at the Q-junction. Unspecified arrows converging at the Q-junction 422 indicate additional ETS-sections with electron entry into Q through glycerophosphate 423 dehydrogenase, dihydro-orotate dehydrogenase, proline dehydrogenase, choline 424 dehydrogenase, and sulfide-ubiquinone oxidoreductase. The dotted arrow indicates the 425 branched pathway of oxygen consumption by alternative quinol oxidase (AOX). ET-426 pathways are coupled to the phosphorylation-pathway. The H⁺_{pos}/O₂ ratio is the outward 427 proton flux from the matrix space to the positively (pos) charged vesicular compartment, 428 divided by catabolic O₂ flux in the NADH-pathway. The H⁺_{neg}/P» ratio is the inward proton 429 flux from the inter-membrane space to the negatively (neg) charged matrix space, divided 430 by the flux of phosphorylation of ADP to ATP. These stoichiometries are not fixed due to 431 ion leaks and proton slip. Modified from Lemieux et al. (2017) and Rich (2013). 432 (C) OXPHOS coupling: O₂ flux through the catabolic ET-pathway, J_{kO_2} , is coupled by the H⁺ circuit to flux through the phosphorylation-pathway of ADP to ATP, J_{P*} . 433 434 (**D**) Chemiosmotic phosphorylation-pathway catalyzed by the proton pump F_1F_0 -ATPase 435 (F-ATPase, ATP synthase), adenine nucleotide translocase (ANT), and inorganic phosphate carrier (PiC). The H⁺_{neg}/P» stoichiometry is the sum of the coupling 436

- 438 2.7 H_{neg}^+ to the matrix, *i.e.*, the negative compartment) and the proton balance in the translocation of ADP³⁻, ATP⁴⁻ and P_i²⁻. Modified from Gnaiger (2014).
- 440

437

441 The plasma membrane separates the intracellular compartment including the cytosol, nucleus, and 442 organelles from the extracellular environment. The plasma membrane consists of a lipid bilayer with 443 embedded proteins and attached organic molecules that collectively control the selective permeability 444 of ions, organic molecules, and particles across the cell boundary. The intact plasma membrane prevents 445 the passage of many water-soluble mitochondrial substrates and inorganic ions—such as succinate, 446 adenosine diphosphate (ADP) and inorganic phosphate (P_i) , that must be precisely controlled at 447 kinetically-saturating concentrations for the analysis of mitochondrial respiratory capacities. 448 Respiratory capacities delineate, comparable to channel capacity in information theory (Schneider 449 2006), the upper bound of the rate of O₂ consumption measured in defined respiratory states. Despite 450 the activity of solute carriers, e.g., SLC13A3 and SLC20A2, which transport specific metabolites across 451 the plasma membrane of various cell types, the intact plasma membrane limits the scope of 452 investigations into mitochondrial respiratory function in intact cells.

stoichiometry in the F-ATPase reaction (-2.7 H⁺_{pos} from the positive intermembrane space,

453 2.1.2. Specification of biochemical dose: Substrates, uncouplers, inhibitors, and other chemical 454 reagents are titrated to analyse cellular and mitochondrial function. Nominal concentrations of these 455 substances are usually reported as initial amount of substance concentration [mol· L^{-1}] in the incubation 456 medium. When aiming at the measurement of kinetically saturated processes—such as OXPHOS-457 capacities, the concentrations for substrates can be chosen according to the apparent equilibrium 458 constant, $K_{\rm m}$ '. In the case of hyperbolic kinetics, only 80% of maximum respiratory capacity is obtained 459 at a substrate concentration of four times the $K_{\rm m}$ ', whereas substrate concentrations of 5, 9, 19 and 49 460 times the $K_{\rm m}$ ' are theoretically required for reaching 83%, 90%, 95% or 98% of the maximal rate 461 (Gnaiger 2001). Other reagents are chosen to inhibit or alter a particular process. The amount of these 462 chemicals in an experimental incubation is selected to maximize effect, avoiding unacceptable off-target 463 consequences that would adversely affect the data being sought. Specifying the amount of substance in 464 an incubation as nominal concentration in the aqueous incubation medium can be ambiguous (Doskey et al. 2015), particularly for cations (TPP⁺; fluorescent dyes such as safranin, TMRM; Chowdhury et al. 465 466 2015) and lipophilic substances (oligomycin, uncouplers, permeabilization agents; Doerrier et al. 2018), 467 which accumulate in the mitochondrial matrix or in biological membranes, respectively. Generally, dose/exposure can be specified per unit of biological sample, i.e., (nominal moles of 468 469 xenobiotic)/(number of cells) [mol·cell⁻¹] or, as appropriate, per mass of biological sample [mol·kg⁻¹]. 470 This approach to specification of dose/exposure provides a scalable parameter that can be used to design 471 experiments, help interpret a wide variety of experimental results, and provide absolute information that 472 allows researchers worldwide to make the most use of published data (Doskey et al. 2015).

473 2.2. Mitochondrial preparations

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475 Mitochondrial preparations are defined as either isolated mitochondria, or tissue and cellular 476 preparations in which the barrier function of the plasma membrane is disrupted. Since this entails the 477 loss of cell viability, mitochondrial preparations are not studied in vivo. In contrast to isolated mitochondria and tissue homogenate preparations, mitochondria in permeabilized tissues and cells are 478 479 in situ relative to the plasma membrane. When studying mitochondrial preparations, substrate-480 uncoupler-inhibitor-titration (SUIT) protocols are used to establish respiratory coupling control states (CCS) and pathway control states (PCS) that provide reference values for various output variables 481 482 (Table 1). Physiological conditions in vivo deviate from these experimentally obtained states; this is 483 because kinetically-saturating concentrations, e.g., of ADP, oxygen (O₂; dioxygen) or fuel substrates, 484 may not apply to physiological intracellular conditions. Further information is obtained in studies of 485 kinetic responses to variations in fuel substrate concentrations, [ADP], or [O₂] in the range between 486 kinetically-saturating concentrations and anoxia (Gnaiger 2001).

487 The cholesterol content of the plasma membrane is high compared to mitochondrial membranes 488 (Korn 1969). Therefore, mild detergents—such as digitonin and saponin—can be applied to selectively 489 permeabilize the plasma membrane via interaction with cholesterol; this allows free exchange of organic 490 molecules and inorganic ions between the cytosol and the immediate cell environment, while 491 maintaining the integrity and localization of organelles, cytoskeleton, and the nucleus. Application of 492 permeabilization agents (mild detergents or toxins) leads to washout of cytosolic marker enzymes— 493 such as lactate dehydrogenase—and results in the complete loss of cell viability (tested by nuclear 494 staining using plasma membrane-impermeable dyes), while mitochondrial function remains intact 495 (tested by cytochrome c stimulation of respiration). Digitonin concentrations have to be optimizeded 496 according to cell type, particularly since mitochondria from cancer cells contain significantly higher 497 contents of cholesterol in both membranes (Baggetto and Testa-Perussini, 1990). For example, a dose of digitonin of 8 fmol·cell⁻¹ (10 pg·cell⁻¹; 10 μ g·10⁻⁶ cells) is optimal for permeabilization of endothelial 498 499 cells, and the concentration in the incubation medium has to be adjusted according to the cell density 500 applied (Doerrier et al. 2018). Respiration of isolated mitochondria remains unaltered after the addition 501 of low concentrations of digitonin or saponin. In addition to mechanical cell disruption during 502 homogenization of tissue, permeabilization agents may be applied to ensure permeabilization of all cells 503 in tissue homogenates.

504 Suspensions of cells permeabilized in the respiration chamber and crude tissue homogenates 505 contain all components of the cell at highly dilute concentrations. All mitochondria are retained in 506 chemically-permeabilized mitochondrial preparations and crude tissue homogenates. In the preparation 507 of isolated mitochondria, however, the mitochondria are separated from other cell fractions and purified 508 by differential centrifugation, entailing the loss of mitochondria at typical recoveries ranging from 30% 509 to 80% of total mitochondrial content (Lai et al. 2018). Using Percoll or sucrose density gradients to 510 maximize the purity of isolated mitochondria may compromise the mitochondrial yield or structural and 511 functional integrity. Therefore, mitochondrial isolation protocols need to be optimized according to each 512 study. The term mitochondrial preparation does neither include intact cells, nor submitochondrial 513 particles and further fractionation of mitochondrial components.

- 515 2.3. Electron transfer pathways
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517 Mitochondrial electron transfer (ET) pathways are fuelled by diffusion and transport of substrates 518 across the mtOM and mtIM. In addition, the mitochondrial electron transfer system (ETS) consists of 519 the matrix-ETS, and membrane-ETS (Figure 2B). Upstream sections of ET-pathways converge at the 520 NADH-junction (N-junction). NADH is mainly generated in the tricarboxylic acid (TCA) cycle and is 521 oxidized by Complex I (CI), with further electron entry into the coenzyme Q-junction (Q-junction). 522 Similarly, succinate is formed in the TCA cycle and oxidized by CII to fumarate. CII is part of both the 523 TCA cycle and the ETS, and reduces FAD to FADH₂ with further reduction of ubiquinone to ubiquinol 524 downstream of the TCA cycle in the Q-junction. Thus FADH₂ is not a substrate but is the product of 525 CII, in contrast to erroneous metabolic maps shown in many publications. β-oxidation of fatty acids 526 (FA) generates FADH₂ as the substrate of electron transferring flavoprotein complex (CETF).

527 Selected mitochondrial catabolic pathways, k, of electron transfer from the oxidation of fuel 528 substrates to the reduction of O_2 are activated by depletion of endogenous substrates and addition of fuel substrates to the mitochondrial respiration medium (Figure 2B). Substrate combinations and specific
 inhibitors of ET-pathway enzymes are used to obtain defined pathway control states in mitochondrial
 preparations (Gnaiger 2014).

533 2.4. Respiratory coupling control

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535 2.4.1. Coupling: In mitochondrial electron transfer, vectorial transmembrane proton flux is 536 coupled through the redox proton pumps CI, CIII and CIV to the catabolic flux of scalar reactions, 537 collectively measured as O_2 flux, J_{kO_2} (Figure 2). Thus mitochondria are elementary components of 538 energy transformation. Energy is a conserved quantity and cannot be lost or produced in any internal 539 process (First Law of thermodynamics). Open and closed systems can gain or lose energy only by 540 external fluxes—by exchange with the environment. Therefore, energy can neither be produced by 541 mitochondria, nor is there any internal process without energy conservation. Exergy or Gibbs energy 542 ('free energy') is the part of energy that can potentially be transformed into work under conditions of 543 constant temperature and pressure. *Coupling* is the interaction of an exergonic process (spontaneous, 544 negative exergy change) with an endergonic process (positive exergy change) in energy transformations 545 which conserve part of the exergy that would be irreversibly lost or dissipated in an uncoupled process.

Pathway control states (PCS) and coupling control states (CCS) are complementary, since
mitochondrial preparations depend on (1) an exogenous supply of pathway-specific fuel substrates and
oxygen, and (2) exogenous control of phosphorylation (Figure 2).

549 **2.4.2.** Phosphorylation, P_{*} , and P_{*}/O_{2} ratio: Phosphorylation in the context of OXPHOS is 550 defined as phosphorylation of ADP by P_i to form ATP. On the other hand, the term phosphorylation is 551 used generally in many contexts, *e.g.*, protein phosphorylation. This justifies consideration of a symbol 552 more discriminating and specific than P as used in the P/O ratio (phosphate to atomic oxygen ratio), 553 where P indicates phosphorylation of ADP to ATP or GDP to GTP (Figure 2). We propose the symbol 554 P» for the endergonic (uphill) direction of phosphorylation ADP-ATP, and likewise the symbol P« for the corresponding exergonic (downhill) hydrolysis ATP \rightarrow ADP. P» refers mainly to electrontransfer 555 phosphorylation but may also involve substrate-level phosphorylation as part of the TCA cycle 556 557 (succinyl-CoA ligase; phosphoglycerate kinase) and phosphorylation of ADP catalyzed by pyruvate 558 kinase, and of GDP phosphorylated by phosphoenolpyruvate carboxykinase. Transphosphorylation is 559 performed by adenylate kinase, creatine kinase (mtCK), hexokinase and nucleoside diphosphate kinase. 560 In isolated mammalian mitochondria, ATP production catalyzed by adenylate kinase (2 ADP \leftrightarrow ATP + AMP) proceeds without fuel substrates in the presence of ADP (Komlódi and Tretter 2017). Kinase 561 562 cycles are involved in intracellular energy transfer and signal transduction for regulation of energy flux.

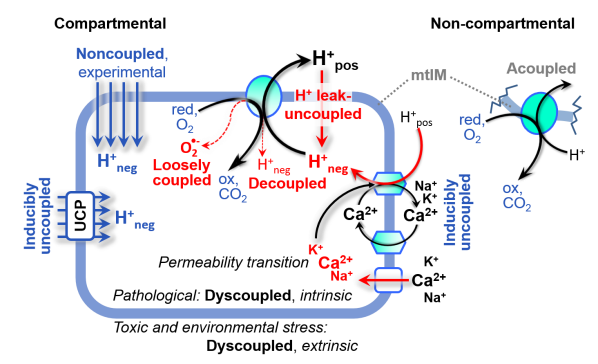
The P»/O₂ ratio (P»/4 e⁻) is two times the 'P/O' ratio (P»/2 e⁻) of classical bioenergetics. P»/O₂ is a generalized symbol, not specific for determination of P_i consumption (P_i/O₂ flux ratio), ADP depletion (ADP/O₂ flux ratio), or ATP production (ATP/O₂ flux ratio). The mechanistic P»/O₂ ratio—or P»/O₂ stoichiometry—is calculated from the proton–to–O₂ and proton–to–phosphorylation coupling stoichiometries (**Figure 2B**):

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$$P \gg /O_2 = \frac{H_{pos}^+/O_2}{H_{neg}^+/P \gg}$$
 (1)

The H^+_{pos}/O_2 coupling stoichiometry (referring to the full 4 electron reduction of O_2) depends on the relative involvement of the three coupling sites (respiratory Complexes CI, CIII and CIV) in the catabolic ET-pathway from reduced fuel substrates (electron donors) to the reduction of O_2 (electron acceptor). This varies with: (1) a bypass of CI by single or multiple electron input into the Q-junction; and (2) a bypass of CIV by involvement of alternative oxidases, AOX, which are not expressed in mammalian mitochondria.

The H^+_{pos}/O_2 coupling stoichiometry equals 12 in the ET-pathways involving CIII and CIV as 577 proton pumps, increasing to 20 for the NADH-pathway through CI (Figure 2B), but a general consensus 578 579 on H⁺_{pos}/O₂ stoichiometries remains to be reached (Hinkle 2005; Wikström and Hummer 2012; Sazanov 2015). The H_{neg}^+/P coupling stoichiometry (3.7; Figure 2B) is the sum of 2.7 H_{neg}^+ required by the F-580 581 ATPase of vertebrate and most invertebrate species (Watt et al. 2010) and the proton balance in the 582 translocation of ADP, ATP and P_i (Figure 2C). Taken together, the mechanistic $P \gg O_2$ ratio is calculated 583 at 5.4 and 3.3 for NADH- and succinate-linked respiration, respectively (Eq. 1). The corresponding 584 classical P»/O ratios (referring to the 2 electron reduction of $0.5 O_2$) are 2.7 and 1.6 (Watt *et al.* 2010), 585 in agreement with the measured P»/O ratio for succinate of 1.58 ± 0.02 (Gnaiger *et al.* 2000).



587 588 Figure 3. Mechanisms of respiratory uncoupling

589 An intact mitochondrial inner membrane, mtIM, is required for vectorial, compartmental coupling. 'Acoupled' respiration is the consequence of structural disruption with catalytic activity of non-590 compartmental mitochondrial fragments. Inducible uncoupling (e.g., by activation of UCP1) increases 591 592 LEAK respiration; experimentally noncoupled respiration provides an estimate of ET-capacity obtained 593 by titration of protonophores stimulating respiration to maximum O_2 flux. H⁺ leak-uncoupled, 594 decoupled, and loosely coupled respiration are components of intrinsic uncoupling (Table 2). 595 Pathological dysfunction may affect all types of uncoupling, including permeability transition, causing 596 intrinsically dyscoupled respiration. Similarly, toxicological and environmental stress factors can cause 597 extrinsically dyscoupled respiration. Reduced fuel substrates, red; oxidized products, ox.

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600 **2.4.3. Uncoupling:** The effective $P \gg O_2$ flux ratio $(Y_{P \gg O_2} = J_{P \gg}/J_{kO_2})$ is diminished relative to the mechanistic P»/O₂ ratio by intrinsic and extrinsic uncoupling or dyscoupling (Figure 3). Such 601 602 generalized uncoupling is different from switching to mitochondrial pathways that involve fewer than 603 three proton pumps ('coupling sites': Complexes CI, CIII and CIV), bypassing CI through multiple 604 electron entries into the O-junction, or CIII and CIV through AOX (Figure 2B). Reprogramming of 605 mitochondrial pathways leading to different types of substrates being oxidized may be considered as a 606 switch of gears (changing the stoichiometry by altering the substrate that is oxidized) rather than 607 uncoupling (loosening the tightness of coupling relative to a fixed stoichiometry). In addition, $Y_{P \gg /Q_2}$ depends on several experimental conditions of flux control, increasing as a hyperbolic function of [ADP] 608 609 to a maximum value (Gnaiger 2001).

Uncoupling of mitochondrial respiration is a general term comprising diverse mechanisms:

- 611 1. Proton leak across the mtIM from the pos- to the neg-compartment (H⁺ leak-uncoupled; Figure 3).
 - 2. Cycling of other cations, strongly stimulated by permeability transition; comparable to the use of protonophores, cation cycling is experimentally induced by valinomycin in the presence of K⁺;
- 615
 3. Decoupling by proton slip in the redox proton pumps when protons are effectively not pumped
 616 (CI, CIII and CIV) or are not driving phosphorylation (F-ATPase);
- 617 4. Loss of vesicular (compartmental) integrity when electron transfer is acoupled;
- 5. Electron leak in the loosely coupled univalent reduction of O₂ to superoxide (O₂⁻; superoxide anion radical).
- Differences of terms—uncoupled *vs.* noncoupled—are easily overlooked, although they relate to
 different meanings of uncoupling (Figure 3 and Table 2).

622 2.5. Coupling states and respiratory rates

To extend the classical nomenclature on mitochondrial coupling states (Section 2.6) by a conceptdriven terminology that explicitly incorporates information on the meaning of respiratory states, the terminology must be general and not restricted to any particular experimental protocol or mitochondrial preparation (Gnaiger 2009). Concept-driven nomenclature aims at mapping the meaning and concept behind the words and acronyms onto the forms of words and acronyms (Miller 1991). The focus of concept-driven nomenclature is primarily the conceptual *why*, along with clarification of the experimental *how*.

- 632 Table 1. Coupling states and residual oxygen consumption in mitochondrial 633 preparations in relation to respiration- and phosphorylation-flux, J_{kO_2} and J_{P_N} , and
- 634 **protonmotive force, pmf.** Coupling states are established at kinetically-saturating 635 concentrations of fuel substrates and O₂.

State	$J_{ m kO2}$	$J_{\mathrm{P}*}$	pmf	Inducing factors	Limiting factors
LEAK	<i>L</i> ; low, cation leak- dependent respiration	0	max.	back-flux of cations including proton leak, proton slip	J_{P} = 0: (1) without ADP, L_N ; (2) max. ATP/ADP ratio, L_T ; or (3) inhibition of the phosphorylation- pathway, L_{Omy}
OXPHOS	<i>P</i> ; high, ADP- stimulated respiration, OXPHOS-capacity	max.	high	kinetically- saturating [ADP] and [P _i]	J_{P} by phosphorylation- pathway; or J_{kO2} by ET- capacity
ET	<i>E</i> ; max., noncoupled respiration, ET-capacity	0	low	optimal external uncoupler concentration for max. $J_{O2,E}$	$J_{\rm kO_2}$ by ET-capacity
ROX	<i>Rox</i> ; min., residual O_2 consumption	0	0	<i>J</i> _{O2,<i>Rox</i>} in non-ET- pathway oxidation reactions	inhibition of all ET- pathways; or absence of fuel substrates

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638 To provide a diagnostic reference for respiratory capacities of core energy metabolism, the 639 capacity of oxidative phosphorylation, OXPHOS, is measured at kinetically-saturating concentrations 640 of ADP and P_i. The oxidative ET-capacity reveals the limitation of OXPHOS-capacity mediated by the phosphorylation-pathway. The ET- and phosphorylation-pathways comprise coupled segments of the 641 642 OXPHOS-system. ET-capacity is measured as noncoupled respiration by application of external uncouplers. The contribution of intrinsically uncoupled O₂ consumption is studied by preventing the 643 644 stimulation of phosphorylation either in the absence of ADP or by inhibition of the phosphorylationpathway. The corresponding states are collectively classified as LEAK-states, when O₂ consumption 645 646 compensates mainly for ion leaks, including the proton leak. Defined coupling states are induced by: (1) adding cation chelators such as EGTA, binding free Ca^{2+} and thus limiting cation cycling; (2) adding 647 648 ADP and P_i ; (3) inhibiting the phosphorylation-pathway; and (4) uncoupler titrations, while maintaining 649 a defined ET-pathway state with constant fuel substrates and inhibitors of specific branches of the ET-650 pathway.

The three coupling states, ET, LEAK and OXPHOS, are shown schematically with the corresponding respiratory rates, abbreviated as *E*, *L* and *P*, respectively (**Figure 4**). We distinguish metabolic *pathways* from metabolic *states* and the corresponding metabolic *rates*; for example: ETpathways, ET-states, and ET-capacities, *E*, respectively (**Table 1**). The protonmotive force is *high* in the OXPHOS-state when it drives phosphorylation, *maximum* in the LEAK-state of coupled mitochondria, driven by LEAK-respiration at a minimum back-flux of cations to the matrix side, and *very low* in the ET-state when uncouplers short-circuit the proton cycle (**Table 1**). 661 Respiratory states (ET, OXPHOS, LEAK; Table 1) and corresponding 662 663 rates (E, P, L) are connected by the 664 protonmotive force, pmf. (1) ETcapacity, E, is partitioned into (2) 665 dissipative LEAK-respiration, L, 666 667 when the Gibbs energy change of catabolic O₂ flux is irreversibly lost, 668

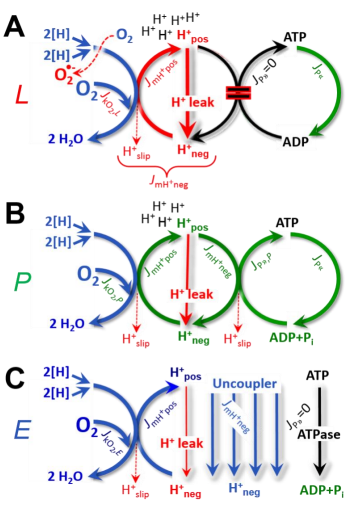
- 669 (3) net OXPHOS-capacity, P-L, with
- 670 partial conservation of the capacity to perform work, and (4) the excess capacity, *E-P*. Modified from
- 671 Gnaiger (2014).

Figure 5. Respiratory coupling states

(A) **LEAK-state and rate**, *L*: Oxidation only, since phosphorylation is arrested, $J_{P} = 0$, and catabolic O₂ flux, $J_{kO_2,L}$, is controlled mainly by the proton leak and slip, J_{mH^+neg} , at maximum protonmotive force (**Figure 4**). Extramitochondrial ATP may be hydrolyzed by extramitochondrial ATPases, J_{P*} ; then phosphorylation must be blocked.

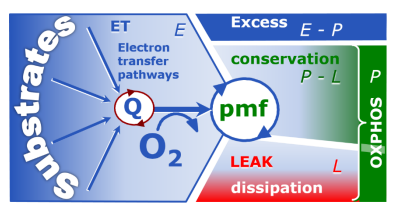
(B) OXPHOS-state and rate, P: Oxidation coupled to phosphorylation, $J_{\rm P*}$. which is stimulated by kinetically-saturating [ADP] and [P_i], supported by a high protonmotive force. O₂ flux, $J_{kO_2,P}$, is well-coupled at a P»/O2 ratio of $J_{\mathrm{P}\gg,P}/J_{\mathrm{O}_2,P}$. Extramitochondrial ATPases may recycle ATP, $J_{P^{*}}$. (C) ET-state and rate, E: Oxidation only, since phosphorylation is zero, $J_{\rm P*} = 0$, at optimum exogenous uncoupler concentration when

uncoupler concentration when noncoupled respiration, $J_{kO2,E}$, is maximum. The F-ATPase may hydrolyze extramitochondrial ATP.





673 **2.5.1.** LEAK-state (Figure 5A): The LEAK-state is defined as a state of mitochondrial respiration when O₂ flux mainly compensates for ion leaks in the absence of ATP synthesis, at 674 675 kinetically-saturating concentrations of O₂, respiratory fuel substrates and P_i. LEAK-respiration is 676 measured to obtain an estimate of intrinsic uncoupling without addition of an experimental uncoupler: (1) in the absence of adenylates, *i.e.*, AMP, ADP and ATP; (2) after depletion of ADP at a maximum 677 ATP/ADP ratio; or (3) after inhibition of the phosphorylation-pathway by inhibitors of F-ATPase—such 678 679 as oligomycin, or of adenine nucleotide translocase—such as carboxyatractyloside. Adjustment of the nominal concentration of these inhibitors to the density of biological sample applied can minimize or 680 681 avoid inhibitory side-effects exerted on ET-capacity or even some dyscoupling.



Term	Term		P»/O ₂	Notes
acoupl	ed		0	electron transfer in mitochondrial fragments without vectorial proton translocation (Figure 3)
	uncoupled	L	0	non-phosphorylating LEAK-respiration (Figure 5A)
e addeo	proton leak- uncoupled		0	component of <i>L</i> , H ⁺ diffusion across the mtIM (Figure 3)
hore	decoupled		0	component of <i>L</i> , proton slip (Figure 3)
intrinsic, no protonophore added λ	loosely coupled		0	component of <i>L</i> , lower coupling due to superoxide formation and bypass of proton pumps by electron leak (Figure 3)
sic, no	dyscoupled		0	pathologically, toxicologically, environmentally increased uncoupling, mitochondrial dysfunction
intrins	inducibly uncoupled		0	by UCP1 or cation (<i>e.g.</i> , Ca ²⁺) cycling (Figure 3)
noncou	noncoupled		0	ET-capacity, non-phosphorylating respiration stimulated to maximum flux at optimum exogenous protonophore concentration (Figure 5C)
well-co	well-coupled		high	OXPHOS-capacity, phosphorylating respiration with an intrinsic LEAK component (Figure 5B)
fully c	oupled	P-L	max.	OXPHOS-capacity corrected for LEAK-respiration (Figure 4)

Table 2. Terms on respiratory coupling and uncoupling.

• **Proton leak and uncoupled respiration:** The intrinsic proton leak is the *uncoupled* leak current of protons in which protons diffuse across the mtIM in the dissipative direction of the downhill protonmotive force without coupling to phosphorylation (**Figure 5A**). The proton leak flux depends non-linearly on the protonmotive force (Garlid *et al.* 1989; Divakaruni and Brand 2011), which is a temperature-dependent property of the mtIM and may be enhanced due to possible contaminations by free fatty acids. Inducible uncoupling mediated by uncoupling protein 1 (UCP1) is physiologically controlled, *e.g.*, in brown adipose tissue. UCP1 is a member of the mitochondrial carrier family that is involved in the translocation of protons across the mtIM (Klingenberg 2017). Consequently, this short-circuit lowers the protonmotive force and stimulates electron transfer, respiration, and heat dissipation in the absence of phosphorylation of ADP.

- Cation cycling: There can be other cation contributors to leak current including calcium and probably magnesium. Calcium influx is balanced by mitochondrial Na⁺/Ca²⁺ or H⁺/Ca²⁺ exchange, which is balanced by Na⁺/H⁺ or K⁺/H⁺ exchanges. This is another effective uncoupling mechanism different from proton leak (Table 2).
- Proton slip and decoupled respiration: Proton slip is the *decoupled* process in which protons • are only partially translocated by a redox proton pump of the ET-pathways and slip back to the original vesicular compartment. The proton leak is the dominant contributor to the overall leak current in mammalian mitochondria incubated under physiological conditions at 37 °C, whereas proton slip increases at lower experimental temperature (Canton et al. 1995). Proton slip can also happen in association with the F-ATPase, in which the proton slips downhill across the pump to the matrix without contributing to ATP synthesis. In each case, proton slip is a property of the proton pump and increases with the pump turnover rate.
- Electron leak and loosely coupled respiration: Superoxide production by the ETS leads to a bypass of redox proton pumps and correspondingly lower P»/O₂ ratio. This depends on the actual site of electron leak and the scavenging of hydrogen peroxide by cytochrome *c*, whereby electrons may re-enter the ETS with proton translocation by CIV.

- Loss of compartmental integrity and acoupled respiration: Electron transfer and catabolic O₂
 flux proceed without compartmental proton translocation in disrupted mitochondrial fragments.
 Such fragments are an artefact of mitochondrial isolation, and may not fully fuse to re-establish
 structurally intact mitochondria. Loss of mtIM integrity, therefore, is the cause of acoupled
 respiration, which is a nonvectorial dissipative process without control by the protonmotive force.
- Dyscoupled respiration: Mitochondrial injuries may lead to *dyscoupling* as a pathological or toxicological cause of *uncoupled* respiration. Dyscoupling may involve any type of uncoupling mechanism, *e.g.*, opening the permeability transition pore. Dyscoupled respiration is distinguished from the experimentally induced *noncoupled* respiration in the ET-state (Table 2).

2.5.2. OXPHOS-state (Figure 5B): The OXPHOS-state is defined as the respiratory state with
 kinetically-saturating concentrations of O₂, respiratory and phosphorylation substrates, and absence of
 exogenous uncoupler, which provides an estimate of the maximal respiratory capacity in the OXPHOS state for any given ET-pathway state. Respiratory capacities at kinetically-saturating substrate
 concentrations provide reference values or upper limits of performance, aiming at the generation of data
 sets for comparative purposes. Physiological activities and effects of substrate kinetics can be evaluated
 relative to the OXPHOS-capacity.

728 As discussed previously, 0.2 mM ADP does not fully saturate flux in isolated mitochondria 729 (Gnaiger 2001; Puchowicz et al. 2004); greater [ADP] is required, particularly in permeabilized muscle 730 fibres and cardiomyocytes, to overcome limitations by intracellular diffusion and by the reduced conductance of the mtOM (Jepihhina et al. 2011, Illaste et al. 2012, Simson et al. 2016), either through 731 732 interaction with tubulin (Rostovtseva et al. 2008) or other intracellular structures (Birkedal et al. 2014). 733 In addition, saturating ADP concentrations need to be evaluated under different experimental conditions 734 such as temperature (Lemieux et al. 2017) and with different animal models (Blier and Guderley, 1993). 735 In permeabilized muscle fibre bundles of high respiratory capacity, the apparent K_m for ADP increases up to 0.5 mM (Saks et al. 1998), consistent with experimental evidence that >90% saturation is reached 736 only at >5 mM ADP (Pesta and Gnaiger 2012). Similar ADP concentrations are also required for 737 738 accurate determination of OXPHOS-capacity in human clinical cancer samples and permeabilized cells 739 (Klepinin et al. 2016; Koit et al. 2017). Whereas 2.5 to 5 mM ADP is sufficient to obtain the actual 740 OXPHOS-capacity in many types of permeabilized tissue and cell preparations, experimental validation 741 is required in each specific case.

742 **2.5.3. Electron transfer-state (Figure 5C)**: O_2 flux determined in the ET-state yields an estimate 743 of ET-capacity. The ET-state is defined as the noncoupled state with kinetically-saturating 744 concentrations of O₂, respiratory substrate and optimum exogenous uncoupler concentration for 745 maximum O_2 flux. Uncouplers are weak lipid-soluble acids which function as protonophores. These 746 disrupt the barrier function of the mtIM and thus short circuit the protonmotive system, functioning like 747 a clutch in a mechanical system. As a consequence of the nearly collapsed protonmotive force, the driving force is insufficient for phosphorylation, and $J_{P*} = 0$. The most frequently used uncouplers are 748 749 cvanide m-chloro phenyl hydrazone carbonyl carbonyl (CCCP), cyanide p-750 trifluoromethoxyphenylhydrazone (FCCP), or dinitrophenole (DNP). Stepwise titration of uncouplers 751 stimulates respiration up to or above the level of O₂ consumption rates in the OXPHOS-state; respiration 752 is inhibited, however, above optimum uncoupler concentrations (Mitchell 2011). Data obtained with a 753 single dose of uncoupler must be evaluated with caution, particularly when a fixed uncoupler 754 concentration is used in studies exploring a treatment or disease that may alter the mitochondrial content 755 or mitochondrial sensitivity to inhibition by uncouplers. The effect on ET-capacity of the reversed 756 function of F-ATPase ($J_{P_{\alpha}}$; Figure 5C) can be evaluated in the presence and absence of 757 extramitochondrial ATP.

758 2.5.4. ROX state and Rox: Besides the three fundamental coupling states of mitochondrial 759 preparations, the state of residual O_2 consumption, ROX, is relevant to assess respiratory function 760 (Figure 1). ROX is not a coupling state. The rate of residual oxygen consumption, Rox, is defined as O_2 761 consumption due to oxidative reactions measured after inhibition of ET-with rotenone, malonic acid 762 and antimycin A. Cyanide and azide inhibit not only CIV but catalase and several peroxidases involved in Rox. High concentrations of antimycin A, but not rotenone or cyanide, inhibit peroxisomal acvl-CoA 763 764 oxidase and D-amino acid oxidase (Vamecq et al. 1987). Rox represents a baseline that is used to correct 765 respiration measured in defined coupling states. Rox-corrected L, P and E not only lower the values of 766 total fluxes, but also change the flux control ratios L/P and L/E. Rox is not necessarily equivalent to non767 mitochondrial reduction of O₂, considering O₂-consuming reactions in mitochondria that are not related 768 to ET—such as O₂ consumption in reactions catalyzed by monoamine oxidases (type A and B), monooxygenases (cytochrome P450 monooxygenases), dioxygenase (sulfur dioxygenase and 769 770 trimethyllysine dioxygenase), and several hydoxylases. Even isolated mitochondrial fractions, 771 especially those obtained from liver, may be contaminated by peroxisomes. This fact makes the exact 772 determination of mitochondrial O₂ consumption and mitochondria-associated generation of reactive oxygen species complicated (Schönfeld et al. 2009; Speijer 2016; Figure 2). The dependence of ROX-773 774 linked O_2 consumption needs to be studied in detail together with non-ET enzyme activities, availability 775 of specific substrates, O₂ concentration, and electron leakage leading to the formation of reactive oxygen 776 species.

777 **2.5.5. Quantitative relations:** E may exceed or be equal to P. E > P is observed in many types 778 of mitochondria, varying between species, tissues and cell types (Gnaiger 2009). E-P is the excess ET-779 capacity pushing the phosphorylation-flux (Figure 2C) to the limit of its capacity of utilizing the 780 protonmotive force. In addition, the magnitude of E-P depends on the tightness of respiratory coupling 781 or degree of uncoupling, since an increase of L causes P to increase towards the limit of E. The excess 782 E-P capacity, E-P, therefore, provides a sensitive diagnostic indicator of specific injuries of the 783 phosphorylation-pathway, under conditions when E remains constant but P declines relative to controls 784 (Figure 4). Substrate cocktails supporting simultaneous convergent electron transfer to the Q-junction 785 for reconstitution of TCA cycle function establish pathway control states with high ET-capacity, and 786 consequently increase the sensitivity of the *E*-*P* assay.

E cannot theoretically be lower than *P*. E < P must be discounted as an artefact, which may be caused experimentally by: (1) loss of oxidative capacity during the time course of the respirometric assay, since *E* is measured subsequently to *P*; (2) using insufficient uncoupler concentrations; (3) using high uncoupler concentrations which inhibit ET (Gnaiger 2008); (4) high oligomycin concentrations applied for measurement of *L* before titrations of uncoupler, when oligomycin exerts an inhibitory effect on *E*. On the other hand, the excess ET-capacity is overestimated if non-saturating [ADP] or [P_i] are used. See State 3 in the next section.

794 The net OXPHOS-capacity is calculated by subtracting L from P (Figure 4). The net $P \gg O_2$ equals 795 P»/(P-L), wherein the dissipative LEAK component in the OXPHOS-state may be overestimated. This 796 can be avoided by measuring LEAK-respiration in a state when the protonmotive force is adjusted to its 797 slightly lower value in the OXPHOS-state—by titration of an ET inhibitor (Divakaruni and Brand 2011). 798 Any turnover-dependent components of proton leak and slip, however, are underestimated under these 799 conditions (Garlid et al. 1993). In general, it is inappropriate to use the term ATP production or ATP 800 turnover for the difference of O₂ flux measured in the OXPHOS and LEAK states. P-L is the upper limit 801 of OXPHOS-capacity that is freely available for ATP production (corrected for LEAK-respiration) and 802 is fully coupled to phosphorylation with a maximum mechanistic stoichiometry (Figure 4).

803 The rates of LEAK respiration and OXPHOS capacity depend on (1) the tightness of coupling 804 under the influence of the respiratory uncoupling mechanisms (Figure 3), and (2) the coupling 805 stoichiometry, which varies as a function of the substrate type undergoing oxidation in ET-pathways 806 with either two or three coupling sites (Figure 2B). When cocktails with NADH-linked substrates and succinate are used, the relative contribution of ET-pathways with three or two coupling sites cannot be 807 808 controlled experimentally, is difficult to determine, and may shift in transitions between LEAK-, 809 OXPHOS- and ET-states (Gnaiger 2014). Under these experimental conditions, we cannot separate the 810 tightness of coupling versus coupling stoichiometry as the mechanisms of respiratory control in the shift 811 of L/P ratios. The tightness of coupling and fully coupled O₂ flux, P-L (Table 2), therefore, are obtained 812 from measurements of coupling control of LEAK respiration, OXPHOS- and ET-capacities in well 813 defined pathway states, using either pyruvate and malate as substrates or the classical succinate and 814 rotenone substrate-inhibitor combination (Figure 2B).

815 2.5.6. The steady-state: Mitochondria represent a thermodynamically open system in non-816 equilibrium states of biochemical energy transformation. State variables (protonmotive force; redox 817 states) and metabolic *rates* (fluxes) are measured in defined mitochondrial respiratory *states*. Steady-818 states can be obtained only in open systems, in which changes by internal transformations, e.g., O₂ 819 consumption, are instantaneously compensated for by external fluxes, e.g., O_2 supply, preventing a 820 change of O₂ concentration in the system (Gnaiger 1993b). Mitochondrial respiratory states monitored 821 in closed systems satisfy the criteria of pseudo-steady states for limited periods of time, when changes 822 in the system (concentrations of O₂, fuel substrates, ADP, P_i, H⁺) do not exert significant effects on metabolic fluxes (respiration, phosphorylation). Such pseudo-steady states require respiratory media
 with sufficient buffering capacity and substrates maintained at kinetically-saturating concentrations, and
 thus depend on the kinetics of the processes under investigation.

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827 2.6. Classical terminology for isolated mitochondria

When a code is familiar enough, it ceases appearing like a code; one forgets that there is a
decoding mechanism. The message is identical with its meaning' (Hofstadter 1979).

Chance and Williams (1955; 1956) introduced five classical states of mitochondrial respiration
and cytochrome redox states. Table 3 shows a protocol with isolated mitochondria in a closed
respirometric chamber, defining a sequence of respiratory states. States and rates are not specifically
distinguished in this nomenclature.

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- 836
- 837
- 838

Table 3. Metabolic states of mitochondria (Chance and Williams, 1956; Table V).

State	[O ₂]	ADP level	Substrate level	Respiration rate	Rate-limiting substance
1	>0	low	low	slow	ADP
2	>0	high	~0	slow	substrate
3	>0	high	high	fast	respiratory chain
4	>0	low	high	slow	ADP
5	0	high	high	0	oxygen

839 840

841 2.6.1. State 1 is obtained after addition of isolated mitochondria to air-saturated
842 isoosmotic/isotonic respiration medium containing P_i, but no fuel substrates and no adenylates, *i.e.*,
843 AMP, ADP, ATP.

2.6.2. State 2 is induced by addition of a 'high' concentration of ADP (typically 100 to $300 \,\mu$ M), 844 845 which stimulates respiration transiently on the basis of endogenous fuel substrates and phosphorylates 846 only a small portion of the added ADP. State 2 is then obtained at a low respiratory activity limited by 847 exhausted endogenous fuel substrate availability (Table 3). If addition of specific inhibitors of respiratory complexes-such as rotenone-does not cause a further decline of O2 flux, State 2 is 848 equivalent to the ROX state (See below.). If inhibition is observed, undefined endogenous fuel substrates 849 850 are a confounding factor of pathway control, contributing to the effect of subsequently externally added 851 substrates and inhibitors. In contrast to the original protocol, an alternative sequence of titration steps is frequently applied, in which the alternative 'State 2' has an entirely different meaning, when this second 852 state is induced by addition of fuel substrate without ADP or ATP (LEAK-state; in contrast to State 2 853 854 defined in Table 1 as a ROX state). Some researchers have called this condition as "pseudostate 4" 855 because it has no significant concentrations of adenine nucleotides and hence it is not a near-856 physiological condition, although it should be used for calculating the net OXPHOS-capacity, P-L.

857 2.6.3. State 3 is the state stimulated by addition of fuel substrates while the ADP concentration 858 is still high (Table 3) and supports coupled energy transformation through oxidative phosphorylation. 'High ADP' is a concentration of ADP specifically selected to allow the measurement of State 3 to State 859 860 4 transitions of isolated mitochondria in a closed respirometric chamber. Repeated ADP titration reestablishes State 3 at 'high ADP'. Starting at O_2 concentrations near air-saturation (193 or 238 μ M O_2 861 862 at 37 °C or 25 °C and sea level at 1 atm or 101.32 kPa, and an oxygen solubility of respiration medium 863 at 0.92 times that of pure water; Forstner and Gnaiger 1983), the total ADP concentration added must 864 be low enough (typically 100 to 300 μ M) to allow phosphorylation to ATP at a coupled O₂ flux that does not lead to O_2 depletion during the transition to State 4. In contrast, kinetically-saturating ADP 865 866 concentrations usually are 10-fold higher than 'high ADP', e.g., 2.5 mM in isolated mitochondria. The abbreviation State 3u is occasionally used in bioenergetics, to indicate the state of respiration after 867 868 titration of an uncoupler, without sufficient emphasis on the fundamental difference between OXPHOScapacity (well-coupled with an endogenous uncoupled component) and ET-capacity (noncoupled). 869

870 **2.6.4.** State 4 is a LEAK-state that is obtained only if the mitochondrial preparation is intact and 871 well-coupled. Depletion of ADP by phosphorylation to ATP causes a decline of O_2 flux in the transition from State 3 to State 4. Under the conditions of State 4, a maximum protonmotive force and high 872 873 ATP/ADP ratio are maintained. The gradual decline of $Y_{P \times /O2}$ towards diminishing [ADP] at State 4 must 874 be taken into account for calculation of P»/O₂ ratios (Gnaiger 2001). State 4 respiration, L_T (Table 1), reflects intrinsic proton leak and ATP hydrolysis activity. O₂ flux in State 4 is an overestimation of 875 LEAK-respiration if the contaminating ATP hydrolysis activity recycles some ATP to ADP, J_{P*} , which 876 877 stimulates respiration coupled to phosphorylation, $J_{P_{0}} > 0$. Some degree of mechanical disruption and 878 loss of mitochondrial integrity allows the exposed mitochondrial F-ATPases to hydrolyze the ATP 879 synthesized by the fraction of coupled mitochondria. This can be tested by inhibition of the 880 phosphorylation-pathway using oligomycin, ensuring that $J_{P} = 0$ (State 40). On the other hand, the State 881 4 respiration reached after exhaustion of added ADP is a more physiological condition (*i.e.*, presence of 882 ATP, ADP and even AMP). Sequential ADP titrations re-establish State 3, followed by State 3 to State 883 4 transitions while sufficient O_2 is available. Anoxia may be reached, however, before exhaustion of 884 ADP (State 5).

885 **2.6.5. State 5** is the state after exhaustion of O_2 in a closed respirometric chamber. Diffusion of 886 O_2 from the surroundings into the aqueous solution may be a confounding factor preventing complete 887 anoxia (Gnaiger 2001). Chance and Williams (1955) provide an alternative definition of State 5, which 888 gives it the different meaning of ROX versus anoxia: 'State 5 may be obtained by antimycin A treatment 889 or by anaerobiosis'.

In Table 3, only States 3 and 4 are coupling control states, with the restriction that rates in State
3 may be limited kinetically by non-saturating ADP concentrations.

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893 2.7. Control and regulation894

895 The terms metabolic control and regulation are frequently used synonymously, but are 896 distinguished in metabolic control analysis: 'We could understand the regulation as the mechanism that 897 occurs when a system maintains some variable constant over time, in spite of fluctuations in external 898 conditions (homeostasis of the internal state). On the other hand, metabolic control is the power to 899 change the state of the metabolism in response to an external signal' (Fell 1997). Respiratory control 900 may be induced by experimental control signals that exert an influence on: (1) ATP demand and ADP 901 phosphorylation-rate; (2) fuel substrate composition, pathway competition; (3) available amounts of 902 substrates and O₂, e.g., starvation and hypoxia; (4) the protonmotive force, redox states, flux-force relationships, coupling and efficiency; (5) Ca^{2+} and other ions including H⁺; (6) inhibitors, *e.g.*, nitric 903 904 oxide or intermediary metabolites such as oxaloacetate; (7) signalling pathways and regulatory proteins, 905 *e.g.*, insulin resistance, transcription factor hypoxia inducible factor 1.

906 Mechanisms of respiratory control and regulation include adjustments of: (1) enzyme activities 907 by allosteric mechanisms and phosphorylation; (2) enzyme content, concentrations of cofactors and 908 conserved moieties—such as adenylates, nicotinamide adenine dinucleotide [NAD⁺/NADH], coenzyme 909 Q, cytochrome c; (3) metabolic channeling by supercomplexes; and (4) mitochondrial density (enzyme 910 concentrations and membrane area) and morphology (cristae folding, fission and fusion). Mitochondria 911 are targeted directly by hormones, e.g., progesterone and glucacorticoids, which affect their energy 912 metabolism (Lee et al. 2013; Gerö and Szabo 2016; Price and Dai 2016; Moreno et al. 2017). 913 Evolutionary or acquired differences in the genetic and epigenetic basis of mitochondrial function (or 914 dysfunction) between individuals; age; biological sex, and hormone concentrations; life style including 915 exercise and nutrition; and environmental issues including thermal, atmospheric, toxic and 916 pharmacological factors, exert an influence on all control mechanisms listed above. For reviews, see 917 Brown 1992; Gnaiger 1993a, 2009; 2014; Paradies et al. 2014; Morrow et al. 2017.

918 Lack of control by a metabolic pathway, *e.g.*, phosphorylation-pathway, means that there will 919 be no response to a variable activating it, *e.g.*, [ADP]. The reverse, however, is not true as the absence 920 of a response to [ADP] does not exclude the phosphorylation-pathway from having some degree of 921 control. The degree of control of a component of the OXPHOS-pathway on an output variable—such 922 as O_2 flux, will in general be different from the degree of control on other outputs—such as 923 phosphorylation-flux or proton leak flux. Therefore, it is necessary to be specific as to which input and 924 output are under consideration (Fell 1997). Respiratory control refers to the ability of mitochondria to adjust O₂ flux in response to external

926 control signals by engaging various mechanisms of control and regulation. Respiratory control is monitored in a mitochondrial preparation under conditions defined as respiratory states, preferentially 927 928 under near-physiological conditions of temperature, pH and medium ionic composition, to generate data 929 of higher biological relevance. When phosphorylation of ADP to ATP is stimulated or depressed, an 930 increase or decrease is observed in electron transfer measured as O_2 flux in respiratory coupling states 931 of intact mitochondria ('controlled states' in the classical terminology of bioenergetics). Alternatively, 932 coupling of electron transfer with phosphorylation is diminished by uncouplers. The corresponding 933 coupling control state is characterized by a high respiratory rate without control by P» (noncoupled or 934 'uncontrolled state').

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937 3. What is a rate?938

939 The term *rate* is not adequately defined to be useful for reporting data. Normalization of 'rates' 940 leads to a diversity of formats. Application of common and defined units is required for direct transfer 941 of reported results into a database. The second [s] is the SI unit for the base quantity *time*. It is also the 942 standard time-unit used in solution chemical kinetics.

The inconsistency of the meanings of rate becomes apparent when considering Galileo Galilei's famous principle, that 'bodies of different weight all fall at the same rate (have a constant acceleration)' (Coopersmith 2010). A rate may be an extensive quantity, which is a *flow*, *I*, when expressed per object (per number of cells or organisms) or per chamber (per system). 'System' is defined as the open or closed chamber of the measuring device. A rate is a *flux*, *J*, when expressed as a size-specific quantity (**Figure 6A; Box 2**).

- **Extensive quantities:** An extensive quantity increases proportionally with system size. For example, mass and volume are extensive quantities. Flow is an extensive quantity. The magnitude of an extensive quantity is completely additive for non-interacting subsystems. The magnitude of these quantities depends on the extent or size of the system (Cohen *et al.* 2008).
- 954 Size-specific quantities: 'The adjective *specific* before the name of an extensive quantity is • 955 often used to mean divided by mass' (Cohen et al. 2008). In this system-paradigm, mass-956 specific flux is flow divided by mass of the system (the total mass of everything within the 957 measuring chamber or reactor). Rates are frequently expressed as volume-specific flux. A mass-specific or volume-specific quantity is independent of the extent of non-interacting 958 959 homogenous subsystems. Tissue-specific quantities (related to the *sample* in contrast to the 960 system) are of fundamental interest in the field of comparative mitochondrial physiology, 961 where *specific* refers to the *type of the sample* rather than *mass of the system*. The term specific, therefore, must be clarified; sample-specific, e.g., muscle mass-specific 962 963 normalization, is distinguished from *system*-specific quantities (mass or volume; **Figure 6**).
- Intensive quantities: In contrast to size-specific properties, forces are intensive quantities defined as the change of an extensive quantity per advancement of an energy transformation (Gnaiger 1993b).
- *N_X* and *m_X* indicate the number format and mass format, respectively, for expressing the quantity of a sample *X*. When different formats are indicated in symbols of derived quantities, the format (*N*, *m*) is shown as a subscript (*underlined italic*), as in *I*_{O2/NX} and *J*_{O2/mX}. Oxygen flow and flux are expressed in the molar format, *n*_{O2} [mol], but in the volume format, *V*_{O2} [m³] in ergometry. For mass-specific flux these formats can be distinguished as *J_{nO2/mX}* and *J_{VO2/mX}*, respectively. Further examples are given in Figure 6 and Table 4.

974 Figure 6. Flow and flux, and
975 normalization in structure976 function analysis
977 (A) When expressing metabolic
978 'rate' measured in a chember a

978 'rate' measured in a chamber, a 979 fundamental distinction is made 980 between relating the rate to the 981 experimental sample (left) or 982 chamber (right). The different 983 meanings of rate need to be 984 specified by the chosen 985 normalization. Left: Results are 986 expressed as mass-specific *flux*, J_{mX} , 987 per mg protein, dry or wet mass. 988 Cell volume, V_{ce} , may be used for 989 normalization (volume-specific 990 flux, J_{Vce}). Right: Flow per chamber, 991 I, or flux per chamber volume, J_V , 992 are merely reported for 993 methodological reasons. 994 (B) O_2 flow per cell, $I_{O_2/N_{ce}}$, is the 995 product of mitochondria-specific 996 flux, mt-density and mass per cell. 997 Unstructured analysis: performance 998 is the product of mass-specific flux, 999 $J_{O_2/MX}$ [mol·s⁻¹·kg⁻¹], and size (mass 1000 per cell). Structured analysis: 1001 performance is the product of mitochondrial function (mt-specific 1002 1003 flux) and structure (mt-content). 1004 Modified from Gnaiger (2014). For further details see Table 4. 1005

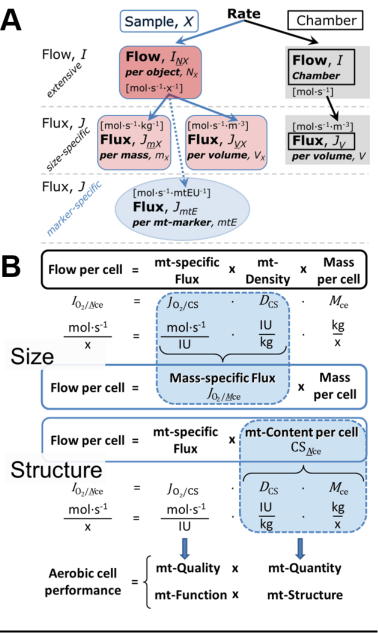
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1011 Box 2: Metabolic flows and fluxes: vectoral, vectorial, and scalar

1013 In a generalization of electrical terms, flow as an extensive quantity (*I*; per system) is distinguished from flux as a size-specific quantity (J; per system size). Flows, I_{tr} , are defined for all 1014 1015 transformations as extensive quantities. Electric charge per unit time is electric flow or current, I_{el} = $dQ_{el} dt^{-1}$ [A = C·s⁻¹]. When dividing I_{el} by size of the system (cross-sectional area of a 'wire'), we obtain 1016 flux as a size-specific quantity, which is the current density (surface-density of flow) perpendicular to 1017 the direction of flux, $J_{el} = I_{el} \cdot A^{-1} [A \cdot m^{-2}]$ (Cohen et al. 2008). Fluxes with *spatial* geometric direction and 1018 magnitude are vectors. Vector and scalar *fluxes* are related to flows as $J_{tr} = I_{tr} \cdot A^{-1} \text{ [mol} \cdot \text{s}^{-1} \cdot \text{m}^{-2} \text{]}$ and $J_{tr} =$ 1019 $I_{tt} \cdot V^{-1}$ [mol·s⁻¹·m⁻³], expressing flux as an area-specific vector or volume-specific vectorial or scalar 1020 1021 quantity, respectively (Gnaiger 1993b). We use the metre-kilogram-second-ampere (MKSA) international system of units (SI) for general cases ([m], [kg], [s] and [A]), with decimal SI prefixes for 1022 1023 specific applications (Table 4).

1024 We suggest to define: (1) vectoral fluxes, which are translocations as functions of gradients with 1025 direction in geometric space in continuous systems; (2) vectorial fluxes, which describe translocations 1026 in discontinuous systems and are restricted to information on *compartmental differences* 1027 (transmembrane proton flux); and (3) scalar fluxes, which are transformations in a homogenous system 1028 (catabolic O_2 flux, J_{kO_2}). 1029 1030

4. Normalization of rate per sample

1031 The challenges of measuring mitochondrial respiratory flux are matched by those of 1032 normalization. Normalization (**Table 4**) is guided by physicochemical principles, methodological 1033 considerations, and conceptual strategies (**Figure 6**).

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Table 4. Sample concentrations and normalization of flux.

Expression	Symbol	Definition	Unit	Notes
Sample				
identity of sample	X	object: cell, tissue, animal, patient		
number of sample entities X	N_X	number of objects	х	1
mass of sample X	m_X		kg	2
mass of object X	M_X	$M_X = m_X \cdot N_X^{-1}$	kg·x ⁻¹	2
Mitochondria				
mitochondria	mt	X = mt		
amount of				
mt-elementary components	mtE	quantity of mt-marker	mtEU	
Concentrations				
object number concentration	$C_{\underline{N}X}$	$C_{\underline{N}\underline{X}} = N_{\underline{X}} \cdot V^{-1}$	x ⋅ m ⁻³	3
sample mass concentration	$C_{\underline{m}X}$	$C_{\underline{m}X} = m_X \cdot V^{-1}$	kg⋅m ⁻³	
mitochondrial concentration	C_{mtE}	$C_{mtE} = mtE \cdot V^{-1}$	mtEU·m ⁻³	4
specific mitochondrial density	D_{mtE}	$D_{mtE} = mtE \cdot m_X^{-1}$	mtEU·kg ⁻¹	5
mitochondrial content,				
<i>mtE</i> per object X	mtE_{NX}	$mtE_{\underline{N}X} = mtE \cdot N_X^{-1}$	mtEU·x ⁻¹	6
O ₂ flow and flux				7
flow, system	I_{O2}	internal flow	mol·s ⁻¹	8
volume-specific flux	$J_{V,\mathrm{O2}}$	$J_{V,\mathrm{O2}} = I_{\mathrm{O2}} \cdot V^{-1}$	mol·s ⁻¹ ·m ⁻³	9
flow per object X	<i>I</i> _{O2/<u>N</u>X}	$I_{O_2/\underline{N}X} = J_{V,O_2} \cdot C_{\underline{N}X}^{-1}$	mol·s ⁻¹ ·x ⁻¹	10
mass-specific flux	$J_{{ m O2}/\underline{m}X}$	$J_{\mathrm{O}_{2}/\underline{m}_{X}} = J_{V,\mathrm{O}_{2}} \cdot C_{\underline{m}_{X}}^{-1}$	mol·s ⁻¹ ·kg ⁻¹	
mt-marker-specific flux	$J_{O_2/mtE}$	$J_{\text{O2/mtE}} = J_{V,\text{O2}} \cdot C_{mtE}^{-1}$	mol·s ⁻¹ ·mtEU ⁻¹	11

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1 The unit x for a number is not used by IUPAC. To avoid confusion, the units [kg·x⁻¹] and [kg] distinguish the mass per object from the mass of a sample that may contain any number of objects. Similarly, the units for flow per system *versus* flow per object are [mol·s⁻¹] (Note 8) and [mol·s⁻¹·x⁻¹] (Note 10).

1041 2 Units are given in the MKSA system (Box 2). The S/ prefix k is used for the S/ base unit of mass (kg = 1,000 g). In praxis, various S/ prefixes are used for convenience, to make numbers easily readable, e.g., 1 mg tissue, cell or mitochondrial mass instead of 0.000001 kg.

1044 3 In case of cells (sample X = cells), the object number concentration is $C_{\underline{Nce}} = N_{ce} \cdot V^1$, and volume 1045 may be expressed in [dm³ = L] or [cm³ = mL]. See **Table 5** for different object types.

1046 4 mt-concentration is an experimental variable, dependent on sample concentration: (1) $C_{mtE} = mtE \cdot V^{-1}$; 1047 (2) $C_{mtE} = mtE_{X} \cdot C_{\underline{N}X}$; (3) $C_{mtE} = C_{\underline{m}X} \cdot D_{mtE}$.

1048 5 If the amount of mitochondria, mtE, is expressed as mitochondrial mass, then D_{mtE} is the mass 1049 fraction of mitochondria in the sample. If mtE is expressed as mitochondrial volume, V_{mt} , and the 1050 mass of sample, m_X , is replaced by volume of sample, V_X , then D_{mtE} is the volume fraction of 1051 mitochondria in the sample.

1052 6 $mtE_{\underline{N}X} = mtE \cdot N_{X}^{-1} = C_{mtE} \cdot C_{\underline{N}X}^{-1}$.

1053 7 O₂ can be replaced by other chemicals to study different reactions, *e.g.*, ATP, H₂O₂, or vesicular compartmental translocations, *e.g.*, Ca²⁺.

- 1055 8 IO2 and V are defined per instrument chamber as a system of constant volume (and constant 1056 temperature), which may be closed or open. I_{O2} is abbreviated for I_{rO2} , *i.e.*, the metabolic or internal 1057 O_2 flow of the chemical reaction r in which O_2 is consumed, hence the negative stoichiometric number, $v_{O2} = -1$. $I_{rO2} = d_r n_{O2}/dt \cdot v_{O2}^{-1}$. If r includes all chemical reactions in which O₂ participates, then 1058 1059 $d_{r}n_{O2} = dn_{O2} - d_{e}n_{O2}$, where dn_{O2} is the change in the amount of O₂ in the instrument chamber and $d_{e}n_{O2}$ 1060 is the amount of O₂ added externally to the system. At steady state, by definition $d_{O_2} = 0$, hence d_{rO_2} 1061 $= -d_e n_{O_2}$. Note that in this context 'external', e, refers to the system, whereas in Figure 1 'external', 1062 ext, refers to the organism.
- 1063 9 J_{V,O_2} is an experimental variable, expressed per volume of the instrument chamber.
- 1064 10 $I_{O2/\underline{N}X}$ is a physiological variable, depending on the size of entity X.
- 1065 11 There are many ways to normalize for a mitochondrial marker, that are used in different experimental approaches: (1) $J_{02/mtE} = J_{V,02} \cdot C_{mtE}^{-1}$; (2) $J_{02/mtE} = J_{V,02} \cdot C_{\underline{m}X}^{-1} \cdot D_{mtE}^{-1} = J_{02/\underline{m}X} \cdot D_{mtE}^{-1}$; (3) $J_{02/mtE} = J_{V,02} \cdot C_{\underline{m}X}^{-1} \cdot mtE_{\underline{N}X}^{-1} = I_{02/\underline{N}X} \cdot mtE_{\underline{N}X}^{-1}$; (4) $J_{02/mtE} = I_{02} \cdot mtE^{-1}$. The mt-elementary unit [mtEU] varies depending on the mt-marker.
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Identity of sample mitochondrial preparation	X	N_X [x]	Mass^a [kg]	Volume [m ³]	mt-Marker [mtEU]
isolated mitochondria	imt		m _{mt}	$V_{ m mt}$	mtE
tissue homogenate	thom		$m_{\rm thom}$		$mtE_{\rm thom}$
permeabilized tissue	pti		m _{pti}		mtE_{pti}
permeabilized fibre	pfi		$m_{\rm pfi}$		mtE_{pfi}
permeabilized cell	pce	$N_{\rm pce}$	$M_{\rm pce}$	$V_{ m pce}$	mtE_{pce}
cells ^b	ce	$N_{\rm ce}$	$M_{ m ce}$	$V_{\rm ce}$	mtE_{ce}
intact cell, viable cell	vce	$N_{ m vce}$	$M_{ m vce}$	$V_{ m vce}$	
dead cell	dce	$N_{ m dce}$	$M_{ m dce}$	$V_{ m dce}$	
organism	org	$N_{ m org}$	$M_{ m org}$	$V_{ m org}$	

1071 Table 5. Sample types, *X*, abbreviations, and quantification.

^{*a*} Instead of mass, the wet weight or dry weight is frequently stated, W_w or W_d . m_X is mass of the sample [kg], M_X is mass of the object [kg·x⁻¹] (**Table 4**).

^b Total cell count, $N_{ce} = N_{vce} + N_{dce}$

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1076 *4.1. Flow: per object*

4.1.1. Number concentration, $C_{\underline{N}X}$: Normalization per sample concentration is routinely required to report respiratory data. $C_{\underline{N}X}$ is the experimental number concentration of sample *X*. In the case of animals, *e.g.*, nematodes, $C_{\underline{N}X} = N_X/V$ [x·L⁻¹], where N_X is the number of organisms in the chamber. Similarly, the number of cells per chamber volume is the number concentration of permeabilized or intact cells $C_{\underline{N}ce} = N_{ce}/V$ [x·L⁻¹], where N_{ce} is the number of cells in the chamber (**Table 4**).

4.1.2. Flow per object, I_{O_2/N_X} : O₂ flow per cell is calculated from volume-specific O₂ flux, J_{V,O_2} 1083 $[nmol \cdot s^{-1} \cdot L^{-1}]$ (per V of the measurement chamber [L]), divided by the number concentration of cells. 1084 1085 The total cell count is the sum of viable and dead cells, $N_{ce} = N_{vce} + N_{dce}$ (Table 5). The cell viability 1086 index, $VI = N_{vce}/N_{ce}$, is the ratio of viable cells (N_{vce} ; before experimental permeabilization) per total cell 1087 count. After experimental permeabilization, all cells are permeabilized, $N_{\rm pce} = N_{\rm ce}$. The cell viability 1088 index can be used to normalize respiration for the number of cells that have been viable before experimental permeabilization, $I_{O2/Nvce} = I_{O2/Nce}/VI$, considering that mitochondrial respiratory 1089 1090 dysfunction in dead cells should be eliminated as a confounding factor.

1091 The complexity changes when the object is a whole organism studied as an experimental model. 1092 The scaling law in respiratory physiology reveals a strong interaction between O_2 flow and individual 1093 body mass: *basal* metabolic rate (flow) does not increase linearly with body mass, whereas *maximum* 1094 mass-specific O_2 flux, \dot{V}_{O2max} or \dot{V}_{O2peak} , is approximately constant across a large range of individual 1095 body mass (Weibel and Hoppeler 2005). Individuals, breeds and species, however, deviate substantially 1096 from this relationship. \dot{V}_{O2peak} of human endurance athletes is 60 to 80 mL $O_2 \cdot \min^{-1} \cdot kg^{-1}$ body mass, 1097 converted to $J_{O2peak/Morg}$ of 45 to 60 nmol·s⁻¹·g⁻¹ (Gnaiger 2014; **Table 6**).

1098 4.2. Size-specific flux: per sample size

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4.2.1. Sample concentration, $C_{\underline{m}X}$: Considering permeabilized tissue, homogenate or cells as the sample, *X*, the sample mass is m_X [mg], which is frequently measured as wet or dry weight, W_w or W_d [mg], respectively, or as amount of protein, m_{Protein} . The sample concentration is the mass of the subsample per volume of the measurement chamber, $C_{\underline{m}X} = m_X/V$ [g·L⁻¹ = mg·mL⁻¹]. *X* is the type of sample—isolated mitochondria, tissue homogenate, permeabilized fibres or cells (**Table 5**).

1105**4.2.2. Size-specific flux:** Cellular O_2 flow can be compared between cells of identical size. To1106take into account changes and differences in cell size, normalization is required to obtain cell size-1107specific or mitochondrial marker-specific O_2 flux (Renner *et al.* 2003).

- **Mass-specific flux**, $J_{O2/mX}$ [mol·s⁻¹·kg⁻¹]: Mass-specific flux is obtained by expressing respiration per mass of sample, m_X [mg]. Flow per cell is divided by mass per cell, $J_{O2/mce} = I_{O2/Nce}/M_{Nce}$. Or chamber volume-specific flux, $J_{V,O2}$, is divided by mass concentration of X in the chamber, $J_{O2/mX} = J_{V,O2}/C_{mX}$.
 - **Cell volume-specific flux**, $J_{O_2/\underline{VX}}$ [mol·s⁻¹·m⁻³]: Sample volume-specific flux is obtained by expressing respiration per volume of sample. For example, in the case of using cells as sample will be the volume of cells added to the chamber (**Figure 6**).

1115 If size-specific O_2 flux is constant and independent of sample size, then there is no interaction 1116 between the subsystems. For example, a 1.5 mg and a 3.0 mg muscle sample respires at identical mass-1117 specific flux. Mass-specific O_2 flux, however, may change with the mass of a tissue sample, cells or 1118 isolated mitochondria in the measuring chamber, in which the nature of the interaction becomes an issue. 1119 Therefore, cell density must be optimized, particularly in experiments carried out in wells, considering 1120 the confluency of the cell monolayer or clumps of cells (Salabei *et al.* 2014).

1122 4.3. Marker-specific flux: per mitochondrial content

1124 Tissues can contain multiple cell populations that may have distinct mitochondrial subtypes. 1125 Mitochondria undergo dynamic fission and fusion cycles, and can exist in multiple stages and sizes that 1126 may be altered by a range of factors. The isolation of mitochondria (often achieved through differential 1127 centrifugation) can therefore yield a subsample of the mitochondrial types present in a tissue, depending 1128 on the isolation protocols utilized (e.g., centrifugation speed). This possible bias should be taken into 1129 account when planning experiments using isolated mitochondria. Different sizes of mitochondria are 1130 enriched at specific centrifugation speeds, which can be used strategically for isolation of mitochondrial 1131 subpopulations.

1132 Part of the mitochondrial content of a tissue is lost during preparation of isolated mitochondria. 1133 The fraction of isolated mitochondria obtained from a tissue sample is expressed as mitochondrial 1134 recovery. At a high mitochondrial recovery the fraction of isolated mitochondria is more representative 1135 of the total mitochondrial population than in preparations characterized by low recovery. Determination 1136 of the mitochondrial recovery and yield is based on measurement of the concentration of a mitochondrial marker in the stock of isolated mitochondria, $C_{mtE,stock}$, and crude tissue homogenate, $C_{mtE,thom}$, which 1137 1138 simultaneously provides information on the specific mitochondrial density in the sample, D_{mtE} (Table 1139 **4**).

1140 When discussing concepts on normalization, it is essential to consider the question of the study. 1141 If the study aims at comparing tissue performance—such as the effects of a treatment on a specific 1142 tissue, then normalization for tissue mass or protein content is appropriate. However, if the aim is to 1143 find differences on mitochondrial function independent of mitochondrial density (Table 4), then 1144 normalization to a mitochondrial marker is imperative (Figure 6). One cannot assume that quantitative 1145 changes in various markers—such as mitochondrial proteins—necessarily occur in parallel with one 1146 another. It should be established that the marker chosen is not selectively altered by the performed 1147 treatment. In conclusion, the normalization must reflect the question under investigation to reach a 1148 satisfying answer. On the other hand, the goal of comparing results across projects and institutions 1149 requires standardization on normalization for entry into a databank.

1150 **4.3.1. Mitochondrial concentration**, C_{mtE} , and mitochondrial markers: Mitochondrial 1151 organelles comprise a dynamic cellular reticulum in various states of fusion and fission. Hence, the 1152 definition of an "amount" of mitochondria is often misconceived: mitochondria cannot be counted 1153 reliably as a number of occurring elementary components. Therefore, quantification of the "amount" of 1154 mitochondria depends on the measurement of chosen mitochondrial markers. 'Mitochondria are the 1155 structural and functional elementary units of cell respiration' (Gnaiger 2014). The quantity of a 1156 mitochondrial marker can reflect the amount of *mitochondrial elementary components*, *mtE*, expressed 1157 in various mitochondrial elementary units [mtEU] specific for each measured mt-marker (Table 4). 1158 However, since mitochondrial quality may change in response to stimuli-particularly in mitochondrial 1159 dysfunction (Campos et al. 2017) and after exercise training (Pesta et al. 2011) and during aging (Daum 1160 et al. 2013)—some markers can vary while others are unchanged: (1) Mitochondrial volume and 1161 membrane area are structural markers, whereas mitochondrial protein mass is commonly used as a 1162 marker for isolated mitochondria. (2) Molecular and enzymatic mitochondrial markers (amounts or 1163 activities) can be selected as matrix markers, e.g., citrate synthase activity, mtDNA; mtIM-markers, e.g., cytochrome c oxidase activity, aa_3 content, cardiolipin, or mtOM-markers, e.g., the voltage-dependent 1164 1165 anion channel (VDAC), TOM20. (3) Extending the measurement of mitochondrial marker enzyme 1166 activity to mitochondrial pathway capacity, ET- or OXPHOS-capacity can be considered as an integrative functional mitochondrial marker. 1167

1168 Depending on the type of mitochondrial marker, the mitochondrial elementary component, *mtE*, 1169 is expressed in marker-specific units. Mitochondrial concentration in the measurement chamber and the 1170 tissue of origin are quantified as (1) a quantity for normalization in functional analyses, C_{mtE} , and (2) a 1171 physiological output that is the result of mitochondrial biogenesis and degradation, D_{mtE} , respectively (Table 4). It is recommended, therefore, to distinguish experimental mitochondrial concentration, C_{mtE} 1172 = mtE/V and physiological mitochondrial density, $D_{mtE} = mtE/m_X$. Then mitochondrial density is the 1173 1174 amount of mitochondrial elementary components per mass of tissue, which is a biological variable 1175 (Figure 6). The experimental variable is mitochondrial density multiplied by sample mass concentration 1176 in the measuring chamber, $C_{mtE} = D_{mtE} \cdot C_{mX}$, or mitochondrial content multiplied by sample number 1177 concentration, $C_{mtE} = mtE_X \cdot C_{NX}$ (Table 4).

4.3.2. mt-Marker-specific flux, $J_{O2/mtE}$: Volume-specific metabolic O₂ flux depends on: (1) the sample concentration in the volume of the instrument chamber, $C_{\underline{m}X}$, or $C_{\underline{N}X}$; (2) the mitochondrial density in the sample, $D_{mtE} = mtE/m_X$ or $mtE_X = mtE/N_X$; and (3) the specific mitochondrial activity or performance per elementary mitochondrial unit, $J_{O2/mtE} = J_{V,O2}/C_{mtE}$ [mol·s⁻¹·mtEU⁻¹] (**Table 4**). Obviously, the numerical results for $J_{O2/mtE}$ vary with the type of mitochondrial marker chosen for measurement of mtE and $C_{mtE} = mtE/V$ [mtEU·m⁻³].

Different methods are implicated in the quantification of mitochondrial markers and have 1184 different strengths. Some problems are common for all mitochondrial markers, mtE: (1) Accuracy of 1185 1186 measurement is crucial, since even a highly accurate and reproducible measurement of O_2 flux results 1187 in an inaccurate and noisy expression if normalized by a biased and noisy measurement of a mitochondrial marker. This problem is acute in mitochondrial respiration because the denominators used 1188 1189 (the mitochondrial markers) are often small moieties of which accurate and precise determination is 1190 difficult. This problem can be avoided when O₂ fluxes measured in substrate-uncoupler-inhibitor 1191 titration protocols are normalized for flux in a defined respiratory reference state, which is used as an 1192 internal marker and yields flux control ratios, FCRs. FCRs are independent of externally measured 1193 markers and, therefore, are statistically robust, considering the limitations of ratios in general (Jasienski and Bazzaz 1999). FCRs indicate qualitative changes of mitochondrial respiratory control, with highest 1194 1195 quantitative resolution, separating the effect of mitochondrial density or concentration on J_{O_2/m_X} and $I_{O2/NX}$ from that of function per elementary mitochondrial marker, $J_{O2/mtE}$ (Pesta et al. 2011; Gnaiger 1196 1197 2014). (2) If mitochondrial quality does not change and only the amount of mitochondria varies as a 1198 determinant of mass-specific flux, any marker is equally qualified in principle; then in practice selection 1199 of the optimum marker depends only on the accuracy and precision of measurement of the mitochondrial 1200 marker. (3) If mitochondrial flux control ratios change, then there may not be any best mitochondrial 1201 marker. In general, measurement of multiple mitochondrial markers enables a comparison and 1202 evaluation of normalization for a variety of mitochondrial markers. Particularly during postnatal 1203 development, the activity of marker enzymes—such as cytochrome c oxidase and citrate synthase— 1204 follows different time courses (Drahota et al. 2004). Evaluation of mitochondrial markers in healthy 1205 controls is insufficient for providing guidelines for application in the diagnosis of pathological states 1206 and specific treatments.

1207 In line with the concept of the respiratory control ratio (Chance and Williams 1955a), the most 1208 readily used normalization is that of flux control ratios and flux control factors (Gnaiger 2014). Selection 1209 of the state of maximum flux in a protocol as the reference state has the advantages of: (1) internal normalization; (2) statistically validated linearization of the response in the range of 0 to 1; and (3) consideration of maximum flux for integrating a large number of elementary steps in the OXPHOS- or ET-pathways. This reduces the risk of selecting a functional marker that is specifically altered by the treatment or pathology, yet increases the chance that the highly integrative pathway is disproportionately affected, *e.g.*, the OXPHOS- rather than ET-pathway in case of an enzymatic defect in the phosphorylation-pathway. In this case, additional information can be obtained by reporting flux control ratios based on a reference state which indicates stable tissue-mass specific flux.

Stereological determination of mitochondrial content via two-dimensional transmission electron
microscopy can have limitations due to the dynamics of mitochondrial size (Meinild Lundby *et al.*2017). Accurate determination of three-dimensional volume by two-dimensional microscopy can be
both time consuming and statistically challenging (Larsen *et al.* 2012).

1221 The validity of using mitochondrial marker enzymes (citrate synthase activity, CI to CIV amount 1222 or activity) for normalization of flux is limited in part by the same factors that apply to flux control 1223 ratios. Strong correlations between various mitochondrial markers and citrate synthase activity 1224 (Reichmann et al. 1985; Boushel et al. 2007; Mogensen et al. 2007) are expected in a specific tissue of 1225 healthy persons and in disease states not specifically targeting citrate synthase. Citrate synthase activity is acutely modifiable by exercise (Tonkonogi et al. 1997; Leek et al. 2001). Evaluation of mitochondrial 1226 1227 markers related to a selected age and sex cohort cannot be extrapolated to provide recommendations for normalization in respirometric diagnosis of disease, in different states of development and ageing, 1228 1229 different cell types, tissues, and species. mtDNA normalized to nDNA via qPCR is correlated to 1230 functional mitochondrial markers including OXPHOS- and ET-capacity in some cases (Puntschart et al. 1231 1995; Wang et al. 1999; Menshikova et al. 2006; Boushel et al. 2007; Ehinger et al. 2015), but lack of 1232 such correlations have been reported (Menshikova et al. 2005; Schultz and Wiesner 2000; Pesta et al. 1233 2011). Several studies indicate a strong correlation between cardiolipin content and increase in 1234 mitochondrial function with exercise (Menshikova et al. 2005; Menshikova et al. 2007; Larsen et al. 1235 2012; Faber et al. 2014), but it has not been evaluated as a general mitochondrial biomarker in disease. 1236 With no single best mitochondrial marker, a good strategy is to quantify several different biomarkers to minimize the decorrelating effects caused by diseases, treatments, or other factors. Determination of 1237 1238 multiple markers, particularly a matrix marker and a marker from the mtIM, allows tracking changes in 1239 mitochondrial quality defined by their ratio.

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1242 **5. Normalization of rate per system**

1244 5.1. Flow: per chamber

1246 The experimental system (experimental chamber) is part of the measurement instrument, 1247 separated from the environment as an isolated, closed, open, isothermal or non-isothermal system 1248 (**Table 4**). Reporting O₂ flows per respiratory chamber, I_{O2} [nmol·s⁻¹], restricts the analysis to intra-1249 experimental comparison of relative differences.

1251 *5.2. Flux: per chamber volume*

1253 **5.2.1.** System-specific flux, $J_{V,02}$: We distinguish between (1) the system with volume V and mass m defined by the system boundaries, and (2) the sample or objects with volume V_X and mass m_X that are 1254 1255 enclosed in the experimental chamber (Figure 6). Metabolic O_2 flow per object, I_{O_2/N_X} , is the total O_2 flow in the system divided by the number of objects, N_X , in the system. $I_{O2/NX}$ increases as the mass of 1256 the object is increased. Sample mass-specific O₂ flux, $J_{O_2/\underline{m}X}$ should be independent of the mass of the 1257 sample studied in the instrument chamber, but system volume-specific O_2 flux, J_{V,O_2} (per volume of the 1258 1259 instrument chamber), increases in proportion to the mass of the sample in the chamber. Whereas J_{VQ_2} 1260 depends on mass-concentration of the sample in the chamber, it should be independent of the chamber (system) volume at constant sample mass. There are practical limitations to increase the mass-1261 1262 concentration of the sample in the chamber, when one is concerned about crowding effects and 1263 instrumental time resolution.

1264 5.2.2. Advancement per volume: When the reactor volume does not change during the reaction,
1265 which is typical for liquid phase reactions, the volume-specific *flux of a chemical reaction* r is the time

1266 derivative of the advancement of the reaction per unit volume, $J_{V,rB} = d_r \xi_B / dt \cdot V^{-1} [(mol \cdot s^{-1}) \cdot L^{-1}]$. The rate of concentration change is dc_B/dt [(mol·L⁻¹)·s⁻¹], where concentration is $c_B = n_B/V$. There is a difference 1267 between (1) $J_{V,rO2}$ [mol·s⁻¹·L⁻¹] and (2) rate of concentration change [mol·L⁻¹·s⁻¹]. These merge to a single 1268 expression only in closed systems. In open systems, internal transformations (catabolic flux, O₂ 1269 1270 consumption) are distinguished from external flux (such as O₂ supply). External fluxes of all substances 1271 are zero in closed systems. In a closed chamber O_2 consumption (internal flux of catabolic reactions k), 1272 $I_{\rm kO2}$ [pmol·s⁻¹], causes a decline of the amount of O₂ in the system, $n_{\rm O2}$ [nmol]. Normalization of these 1273 quantities for the volume of the system, V [L \equiv dm³], yields volume-specific O₂ flux, $J_{V,kO_2} = I_{kO_2}/V$ [nmol·s⁻¹·L⁻¹], and O₂ concentration, [O₂] or $c_{O_2} = n_{O_2}/V$ [µmol·L⁻¹ = µM = nmol·mL⁻¹]. Instrumental 1274 background O₂ flux is due to external flux into a non-ideal closed respirometer; then total volume-1275 specific flux has to be corrected for instrumental background O_2 flux— O_2 diffusion into or out of the 1276 1277 instrumental chamber. $J_{V,KO2}$ is relevant mainly for methodological reasons and should be compared with 1278 the accuracy of instrumental resolution of background-corrected flux, e.g., $\pm 1 \text{ nmol} \cdot \text{s}^{-1} \cdot \text{L}^{-1}$ (Gnaiger 2001). 'Metabolic' or catabolic indicates O_2 flux, J_{kO_2} , corrected for: (1) instrumental background O_2 1279 1280 flux; (2) chemical background O_2 flux due to autoxidation of chemical components added to the 1281 incubation medium; and (3) Rox for O_2 -consuming side reactions unrelated to the catabolic pathway k. 1282

6. Conversion of units

1286 Many different units have been used to report the O_2 consumption rate, OCR (Table 6). SI base units provide the common reference to introduce the theoretical principles (Figure 6), and are used with 1287 1288 appropriately chosen SI prefixes to express numerical data in the most practical format, with an effort 1289 towards unification within specific areas of application (**Table 7**). Reporting data in SI units—including 1290 the mole [mol], coulomb [C], joule [J], and second [s]—should be encouraged, particularly by journals 1291 which propose the use of SI units.

Table 6. Conversion of various formats and units used in respirometry and **ergometry.** e^{-} is the number of electrons or reducing equivalents. $z_{\rm B}$ is the charge number of entity B.

Format	1 Unit	•	Multiplication factor	<i>SI</i> -unit	Notes
<u>n</u>	ng.atom O·s ⁻¹	(2 e ⁻)	0.5	nmol O ₂ ·s ⁻¹	
<u>n</u>	ng.atom O·min ⁻¹	(2 e ⁻)	8.33	pmol O ₂ ·s ⁻¹	
<u>n</u>	natom O·min ⁻¹	(2 e ⁻)	8.33	pmol O ₂ ·s ⁻¹	
<u>n</u>	nmol O ₂ ·min ⁻¹	(4 e⁻)	16.67	pmol O ₂ ·s ⁻¹	
<u>n</u>	nmol O ₂ ·h ⁻¹	(4 e⁻)	0.2778	pmol O ₂ ·s ⁻¹	
<u>V</u> to <u>n</u>	mL O ₂ ·min ⁻¹ at STF	a D ^a	0.744	µmol O ₂ ·s ⁻¹	1
<u>e</u> to <u>n</u>	W = J/s at -470 kJ/m	nol O ₂	-2.128	µmol O ₂ ·s ⁻¹	
<u>e</u> to <u>n</u>	$mA = mC \cdot s^{-1}$	$(z_{\rm H^+} = 1)$	10.36	nmol H ⁺ ·s ⁻¹	2
<u>e</u> to <u>n</u>	$mA = mC \cdot s^{-1}$	$(z_{O_2} = 4)$	2.59	nmol O ₂ ·s ⁻¹	2
<u>n</u> to <u>e</u>	nmol $H^+ \cdot s^{-1}$	$(z_{\rm H^+} = 1)$	0.09649	mA	3
<u>n</u> to <u>e</u>	nmol O ₂ ·s ⁻¹	$(z_{O_2} = 4)$	0.38594	mA	3

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1 At standard temperature and pressure dry (STPD: 0 $^{\circ}$ C = 273.15 K and 1 atm = 101.325 kPa = 760 mmHg), the molar volume of an ideal gas, $V_{\rm m}$, and $V_{\rm m,O2}$ is 22.414 and 22.392 L·mol⁻¹, respectively. Rounded to three decimal places, both values yield the conversion factor of 0.744. 1299 For comparison at normal temperature and pressure dry (NTPD: 20 °C), $V_{m,O2}$ is 24.038 L·mol⁻¹. 1300 Note that the SI standard pressure is 100 kPa. 1301

2 The multiplication factor is $10^{6}/(z_{\rm B}\cdot F)$. 1302

3 The multiplication factor is $z_{\rm B} \cdot F/10^6$. 1303

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Name	Frequently used unit	Equivalent unit	Notes	
volume-specific flux, J_{V,O_2}	pmol·s ⁻¹ ·mL ⁻¹	nmol·s ⁻¹ ·L ⁻¹	1	
•	mmol·s ⁻¹ ·L ⁻¹	mol·s ⁻¹ ·m ⁻³		
cell-specific flow, $I_{O2/cell}$	pmol·s ⁻¹ ·10 ⁻⁶ cells	amol·s ⁻¹ ·cell ⁻¹	2	
-	pmol·s ⁻¹ ·10 ⁻⁹ cells	zmol·s ⁻¹ ·cell ⁻¹	3	
cell number concentration, C_{Nce}	10 ⁶ cells⋅mL ⁻¹	10 ⁹ cells·L ⁻¹		
mitochondrial protein concentration, C_{mtE}	0.1 mg·mL ⁻¹	0.1 g·L ⁻¹		
mass-specific flux, $J_{O_2/\underline{m}}$	pmol·s ⁻¹ ·mg ⁻¹	nmol·s ⁻¹ ·g ⁻¹	4	
catabolic power, P_k	µW·10 ⁻⁶ cells	pW·cell ⁻¹	1	
volume	1,000 L	m ³ (1,000 kg)		
	L	dm ³ (kg)		
	mL	$cm^{3}(g)$		
	μL	mm ³ (mg)		
	fL	μm ³ (pg)	5	
amount of substance concentration	$M = mol \cdot L^{-1}$	mol·dm ⁻³		
pmol: picomole = 10^{-12} mol	4 nmol: nanomole	$= 10^{-9} \text{ mol}$		
2 amol: attomole = 10^{-18} mol	5 fL: femtolitre =	10 ⁻¹⁵ L		
3 zmol: zeptomole = 10^{-21} mol				

1305 Table 7. Conversion of units with preservation of numerical values.

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1310 Although volume is expressed as m^3 using the SI base unit, the litre [dm³] is a conventional unit of volume for concentration and is used for most solution chemical kinetics. If one multiplies $I_{O2/Nce}$ by 1311 C_{Nce} , then the result will not only be the amount of O₂ [mol] consumed per time [s⁻¹] in one litre [L⁻¹], 1312 but also the change in O₂ concentration per second (for any volume of an ideally closed system). This 1313 is ideal for kinetic modeling as it blends with chemical rate equations where concentrations are typically 1314 1315 expressed in mol·L⁻¹ (Wagner et al. 2011). In studies of multinuclear cells—such as differentiated skeletal muscle cells—it is easy to determine the number of nuclei but not the total number of cells. A 1316 1317 generalized concept, therefore, is obtained by substituting cells by nuclei as the sample entity. This does 1318 not hold, however, for enucleated platelets.

For studies of cells, we recommend that respiration be expressed, as far as possible, as: $(1) O_2$ 1319 1320 flux normalized for a mitochondrial marker, for separation of the effects of mitochondrial quality and content on cell respiration (this includes FCRs as a normalization for a functional mitochondrial 1321 marker); (2) O₂ flux in units of cell volume or mass, for comparison of respiration of cells with different 1322 1323 cell size (Renner et al. 2003) and with studies on tissue preparations, and (3) O₂ flow in units of attomole (10^{-18} mol) of O₂ consumed in a second by each cell [amol·s⁻¹·cell⁻¹], numerically equivalent to 1324 [pmol·s⁻¹·10⁻⁶ cells]. This convention allows information to be easily used when designing experiments 1325 1326 in which O_2 flow must be considered. For example, to estimate the volume-specific O_2 flux in an instrument chamber that would be expected at a particular cell number concentration, one simply needs 1327 to multiply the flow per cell by the number of cells per volume of interest. This provides the amount of 1328 O_2 [mol] consumed per time [s⁻¹] per unit volume [L⁻¹]. At an O_2 flow of 100 amol·s⁻¹·cell⁻¹ and a cell 1329 density of 10^9 cells·L⁻¹ (10⁶ cells·mL⁻¹), the volume-specific O₂ flux is 100 nmol·s⁻¹·L⁻¹ (100 1330 $pmol \cdot s^{-1} \cdot mL^{-1}$). 1331

ET-capacity in human cell types including HEK 293, primary HUVEC and fibroblasts ranges from 50 to 180 amol·s⁻¹·cell⁻¹, measured in intact cells in the noncoupled state (see Gnaiger 2014). At 100 amol·s⁻¹·cell⁻¹ corrected for *Rox*, the current across the mt-membranes, I_{H^+e} , approximates 193 pA·cell⁻¹ or 0.2 nA per cell. See Rich (2003) for an extension of quantitative bioenergetics from the molecular to the human scale, with a transmembrane proton flux equivalent to 520 A in an adult at a catabolic power of -110 W. Modelling approaches illustrate the link between protonmotive force and currents (Willis *et al.* 2016).

1339 We consider isolated mitochondria as powerhouses and proton pumps as molecular machines to 1340 relate experimental results to energy metabolism of the intact cell. The cellular P»/O₂ based on oxidation

of glycogen is increased by the glycolytic (fermentative) substrate-level phosphorylation of 3 P»/Glyc 1341 1342 or 0.5 mol P» for each mol O₂ consumed in the complete oxidation of a mol glycosyl unit (Glyc). Adding 0.5 to the mitochondrial P»/O₂ ratio of 5.4 yields a bioenergetic cell physiological P»/O₂ ratio close to 1343 6. Two NADH equivalents are formed during glycolysis and transported from the cytosol into the 1344 1345 mitochondrial matrix, either by the malate-aspartate shuttle or by the glycerophosphate shuttle (Figure 2A) resulting in different theoretical yields of ATP generated by mitochondria, the energetic cost of 1346 which potentially must be taken into account. Considering also substrate-level phosphorylation in the 1347 TCA cycle, this high P»/O₂ ratio not only reflects proton translocation and OXPHOS studied in isolation, 1348 but integrates mitochondrial physiology with energy transformation in the living cell (Gnaiger 1993a). 1349 1350

1352 7. Conclusions

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Catabolic cell respiration is the process of exergonic and exothermic energy transformation in which scalar redox reactions are coupled to vectorial ion translocation across a semipermeable membrane, which separates the small volume of a bacterial cell or mitochondrion from the larger volume of its surroundings. The electrochemical exergy can be partially conserved in the phosphorylation of ADP to ATP or in ion pumping, or dissipated in an electrochemical short-circuit. Respiration is thus clearly distinguished from fermentation as the counterpart of cellular core energy metabolism. An O₂ flux balance scheme illustrates the relationships and general definitions (**Figures 1 and 2**).

Box 3: Recommendations for studies with mitochondrial preparations

- Normalization of respiratory rates should be provided as far as possible:
 - 1. *Biophysical normalization*: on a per cell basis as O₂ flow; this may not be possible when dealing with coenocytic organisms or tissues without cross-walls separating individual cells (*e.g.*, filamentous fungi, muscle fibers)
 - 2. *Cellular normalization*: per g protein; per cell- or tissue-mass as mass-specific O₂ flux; per cell volume as cell volume-specific flux
 - 3. *Mitochondrial normalization*: per mitochondrial marker as mt-specific flux.

With information on cell size and the use of multiple normalizations, maximum potential information
is available (Renner *et al.* 2003; Wagner *et al.* 2011; Gnaiger 2014). Reporting flow in a respiratory
chamber [nmol·s⁻¹] is discouraged, since it restricts the analysis to intra-experimental comparison of
relative (qualitative) differences.

- Catabolic mitochondrial respiration is distinguished from residual O₂ consumption. Fluxes in mitochondrial coupling states should be, as far as possible, corrected for residual O₂ consumption.
- 1377 • Different mechanisms of uncoupling should be distinguished by defined terms. The tightness of coupling relates to these uncoupling mechanisms, whereas the coupling stoichiometry varies as a 1378 1379 function the substrate type involved in ET-pathways with either three or two redox proton pumps operating in series. Separation of tightness of coupling from the pathway-dependent coupling 1380 stoichiometry is possible only when the substrate type undergoing oxidation remains the same for 1381 respiration in LEAK-, OXPHOS-, and ET-states. In studies of the tightness of coupling, therefore, 1382 simple substrate-inhibitor combinations should be applied to exlcude a shift in substrate competition 1383 1384 which may occur when providing physiological substrate cocktails.
- In studies of isolated mitochondria, the mitochondrial recovery and yield should be reported.
 Experimental criteria for evaluation of purity versus integrity should be considered. Mitochondrial markers—such as citrate synthase activity as an enzymatic matrix marker—provide a link to the tissue of origin on the basis of calculating the mitochondrial recovery, *i.e.*, the fraction of mitochondrial marker obtained from a unit mass of tissue. Total mitochondrial protein is frequently applied as a mitochondrial marker, which is restricted to isolated mitochondria.
- In studies of permeabilized cells, the viability of the cell culture or cell suspension of origin should be reported. Normalization should be evaluated for total cell count or viable cell count.
- Terms and symbols are summarized in Table 8. Their use will facilitate transdisciplinary communication and support further developments towards a consistent theory of bioenergetics and mitochondrial physiology. Technical terms related to and defined with normal words can be used as index terms in databases, support the creation of ontologies towards semantic information processing

1397 (MitoPedia), and help in communicating analytical findings as impactful data-driven stories. 'Making data available without making it understandable may be worse than not making it available 1398 at all' (National Academies of Sciences, Engineering, and Medicine 2018). Success will depend on 1399 taking next steps: (1) exhaustive text-mining considering Omics data and functional data; (2) network 1400 analysis of Omics data with bioinformatics tools; (3) cross-validation with distinct bioinformatics 1401 approaches; (4) correlation with functional data; (5) guidelines for biological validation of network 1402 data. This is a call to carefully contribute to FAIR principles (Findable, Accessible, Interoperable, 1403 Reusable) for the sharing of scientific data. 1404

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1407 Table 8. Terms, symbols, and units.1408

Term	Symbol	Unit	Links and comments
alternative quinol oxidase	AOX		Figure 2B
amount of substance B	n _B	[mol]	
ATP yield per O ₂	$Y_{\rm P \gg / O_2}$		$P \gg O_2$ ratio measured in any
5 - 1 - 2	1///02		respiratory state
catabolic reaction	k		Figure 1 and 3
catabolic respiration	$J_{ m kO_2}$	varies	Figure 1 and 3
cell number	$N_{ m ce}$	[x]	$N_{\rm ce} = N_{\rm vce} + N_{\rm dce}$; Table 5
cell respiration	$J_{ m rO_2}$	varies	Figure 1
cell viability index	VI		$VI = N_{\rm vce}/N_{\rm ce} = 1 - N_{\rm dce}/N_{\rm ce}$
charge number of entity B	$Z_{\rm B}$		Table 6; $z_{O_2} = 4$
Complexes I to IV	CI to CIV		respiratory ET Complexes; F
•			2B
concentration of substance B	$c_{\rm B} = n_{\rm B} \cdot V^{-1};$	$[B] [mol \cdot m^{-3}]$	Box 2
coupling control state	CCS		Section 2.4.1
dead cell number	$N_{ m dce}$	[x]	non-viable cells, loss of plass
			membrane barrier function; 7
electric format	<u>e</u>	[C]	Table 6
electron transfer system	ETS		state; Figure 2B, Figure 4
flow, for substance B	$I_{ m B}$	$[mol \cdot s^{-1}]$	system-related extensive qua
		-	Figure 6
flux, for substance B	$J_{ m B}$	varies	size-specific quantitiy; Figur
inorganic phosphate	$\mathbf{P}_{\mathbf{i}}$		Figure 2C
inorganic phosphate carrier	PiC		Figure 2C
intact cell number,			-
viable cell number	$N_{ m vce}$	[X]	viable cells, intact of plasma
			membrane barrier function; T
LEAK state	LEAK		state; Table 1, Figure 4
mass format	<u>m</u>	[kg]	Table 4, Figure 6
mass of sample X	m_X	[kg]	Table 4
mass, dry mass	$m_{ m d}$	[kg]	mass of sample <i>X</i> ; Figure 6
			(frequently called dry weight
mass, wet mass	$m_{ m w}$	[kg]	mass of sample <i>X</i> ; Figure 6
			(frequently called wet weight
mass of object X	$M_X = m_X \cdot N_X$		mass of entity X; Table 4
MITOCARTA		https://	/www.broadinstitute.org/scient
			community/science/programs
			olic-disease-
			program/publications/mitoca
			carta-in-0

MitoPedia		$\Pi U U . / / W V$	ww.bioblast.at/index.php/MitoPedia
mitochondria or mitochondrial	mt		Box 1
mitochondrial DNA	mtDNA		Box 1
mitochondrial concentration	$C_{mtE} = mtE \cdot V^{-1}$	[mtEU·m ⁻³]	
mitochondrial content	mtE_X		$mtE_X = mtE \cdot N_X^{-1}$; Table 4
mitochondrial		[]	
elementary component	mtE	[mtEU]	quantity of mt-marker; Table 4
mitochondrial elementary unit	mtEU	varies	specific units for mt-marker; Table 4
mitochondrial inner membrane	mtIM	ven tes	MIM is widely used; the first M is
	literivi		replaced by mt; Figure 2; Box 1
mitochondrial outer membrane	mtOM		MOM is widely used; the first M is
intoenondriai outer memorane	Intow		replaced by mt; Figure 2; Box 1
mitochondrial recovery	Y_{mtE}		fraction of <i>mtE</i> recovered in sample
Intochondrial recovery	1 mtE		from the tissue of origin
mit cohon dui al vi al d	V		-
mitochondrial yield	$Y_{mtE/\underline{m}}$		mt-yield per tissues mass; $Y_{mtE/\underline{m}} =$
1 6		r 11	$Y_{mtE} \cdot D_{mtE}$
molar format	<u>n</u>	[mol]	Table 6
negative	neg	- 2-	Figure 4
number concentration of X	$C_{\underline{N}X}$	[x·m⁻³]	Table 4
number format	<u>N</u>	[X]	Table 4, Figure 6
number of entities X	N_X	[X]	Table 4, Figure 6
number of entity B	$N_{ m B}$	[X]	Table 4
oxidative phosphorylation	OXPHOS		state; Table 1, Figure 4
oxygen concentration	$c_{\rm O2} = n_{\rm O2} \cdot V^{-1}$	[mol·m ⁻³]	[O ₂]; Section 3.2
oxygen flux, in reaction r	$J_{ m rO2}$	varies	Figure 1
pathway control state	PCS		Section 2.2
permeabilized cell number	$N_{ m pce}$	[x]	experimental permeabilization of
-			plasma membrane; Table 5
phosphorylation of ADP to ATP	P»		Section 2.2
$P \gg O_2$ ratio	$P \gg O_2$		mechanistic $Y_{P \gg / O_2}$, calculated from
			pump stoichiometries; Figure 2B
positive	pos		Figure 4
proton in the negative compartment	•		Figure 4
proton in the positive compartment			Figure 4
protonmotive force	pmf	[V]	Figures 1, 2A and 4; Table 1
rate of electron transfer in ET state	E	[1]	ET-capacity; Table 1
rate of LEAK respiration			Table 1
rate of oxidative phosphorylation	P		OXPHOS capacity; Table 1
rate of residual oxygen consumption			Table 1, Figure 1
residual oxygen consumption	ROX		state; Table 1
respiratory supercomplex	$SC I_n III_n IV_n$		supramolecular assemblies
			composed of variable copy numbers
	D D	1	(<i>n</i>) of CI, CIII and CIV; Box 1
specific mitochondrial density	$D_{mtE} = mtE \cdot m_X$	·[mtEU·kg ⁻¹]	l able 4
substrate-uncoupler-inhibitor-	~~~~		
titration protocol	SUIT	- 2	##
volume	V	[m ⁻³]	Table 7
volume format	\underline{V}	[m ⁻³]	Table 6

Experimentally, respiration is separated in mitochondrial preparations from the interactions with the fermentative pathways of the intact cell. OXPHOS analysis is based on the study of mitochondrial preparations complementary to bioenergetic investigations of intact cells and organisms—from model organisms to the human species including healthy and diseased persons (patients). Different mechanisms of respiratory uncoupling have to be distinguished (**Figure 3**). Metabolic fluxes measured in defined 1507 coupling and pathway control states (Figures 5 and 6) provide insights into the meaning of cellular and1508 organismic respiration.

1509 The optimal choice for expressing mitochondrial and cell respiration as O_2 flow per biological 1510 sample, and normalization for specific tissue-markers (volume, mass, protein) and mitochondrial 1511 markers (volume, protein, content, mtDNA, activity of marker enzymes, respiratory reference state) is 1512 guided by the scientific question under study. Interpretation of the data depends critically on appropriate 1513 normalization (**Figure 6**).

MitoEAGLE can serve as a gateway to better diagnose mitochondrial respiratory adaptations and defects linked to genetic variation, age-related health risks, sex-specific mitochondrial performance, lifestyle with its effects on degenerative diseases, and thermal and chemical environment. The present recommendations on coupling control states and rates, linked to the concept of the protonmotive force, are focused on studies with mitochondrial preparations (**Box 3**). These will be extended in a series of reports on pathway control of mitochondrial respiration, respiratory states in intact cells, and harmonization of experimental procedures.

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1812 S1. Manuscript phases and versions - an open-access apporach1813

1814 This manuscript on 'Mitochondrial respiratory states and rates' is a position statement in the frame of COST Action
 1815 CA15203 MitoEAGLE. The list of co-authors evolved beyond phase 1 in the bottom-up spirit of COST.

1816 The global MitoEAGLE network made it possible to collaborate with a large number of co-authors to reach 1817 consensus on the present manuscript. Nevertheless, we do not consider scientific progress to be supported by 1818 'declaration' statements (other than on ethical or political issues). Our manuscript aims at providing arguments for 1819 further debate rather than pushing opinions. We hope to initiate a much broader process of discussion and want to 1820 raise the awareness on the importance of a consistent terminology for reporting of scientific data in the field of 1821 bioenergetics, mitochondrial physiology and pathology. Quality of research requires quality of communication. 1822 Some established researchers in the field may not want to re-consider the use of jargon which has become 1823 established despite deficiencies of accuracy and meaning. In the long run, superior standards will become accepted. 1824 We hope to contribute to this evolutionary process, with an emphasis on harmonization rather than standardization.

- 1825 Phase 1 The protonmotive force and respiratory control
 1826 http://www.mitoeagle.org/index.php/The protonm
 - http://www.mitoeagle.org/index.php/The_protonmotive_force_and_respiratory_control
 - 2017-04-09 to 2017-09-18 (44 versions)
 - 2017-09-21 to 2018-02-06 (44+21 versions) <u>http://www.mitoeagle.org/index.php/MitoEAGLE_preprint_2017-09-21</u> 2017-11-11: Print version (16) for MiP2017/MitoEAGLE conference in Hradec Kralove
 - *Phase 2* Mitochondrial respiratory states and rates: Building blocks of mitochondrial physiology Part 1 <u>http://www.mitoeagle.org/index.php/MitoEAGLE_preprint_2018-02-08</u>
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1835 Phase 4 Journal submission

CELL METABOLISM, aiming at indexing by *The Web of Science* and *PubMed*.

1839 **S2. Authors** 1840

This manuscript developed as an open invitation to scientists and students to join as co-authors, to provide a
 balanced view on mitochondrial respiratory control and a consensus statement on reporting data of mitochondrial
 respiration in terms of metabolic flows and fluxes.

1844 Co-authors are added in alphabetical oder based upon a first draft written by the corresponding author, who
1845 edited all versions. *Co-authors confirm to have read the final manuscript, possibly have made additions or*1846 suggestions for improvement, and agree to implement the recommendations into future manuscripts, presentations
1847 and teaching materials.

We continue to invite comments and suggestions, particularly if you are an early career investigator adding
an open future-oriented perspective, or an established scientist providing a balanced historical basis. Your critical
input into the quality of the manuscript will be most welcome, improving our aims to be educational, general,
consensus-oriented, and practically helpful for students working in mitochondrial respiratory physiology.

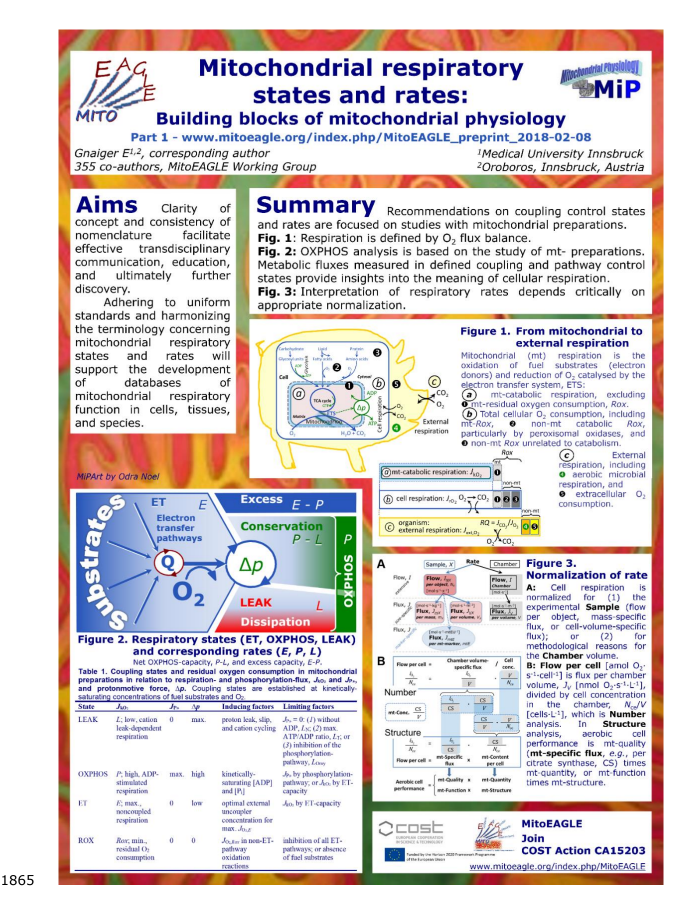
To join as a co-author, please feel free to focus on a particular section, providing direct input and references,
and contributing to the scope of the manuscript from the perspective of your expertise. Your comments will be
largely posted on the discussion page of the MitoEAGLE preprint website.

1855 If you prefer to submit comments in the format of a referee's evaluation rather than a contribution as a co-1856 author, we will be glad to distribute your views to the updated list of co-authors for a balanced response. We would 1857 ask for your consent on this open bottom-up policy.

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1860 S3. Joining COST Actions1861

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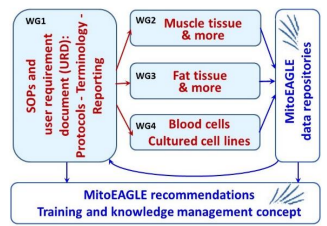


Mitochondrial fitness mapping - Quality management network

Mission of the global MitoEAGLE network

in collaboration with the Mitochondrial Physiology Society, MiPs

- Improve our knowledge on mitochondrial function in health and disease with regard to Evolution, Age, Gender, Lifestyle and Environment
- Interrelate studies across laboratories with the help of a MitoEAGLE data management system
- Provide standardized measures to link mitochondrial and



physiological performance to understand the myriad of factors that play a role in mitochondrial physiology

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