Mitochondria are one of the most exhaustively investigated organelles in the cell and most attention has been paid to the components of the mitochondrial electron transport chain (ETC) in the last 100 years. The ETC collects electrons from NADH or FADH₂ and transfers them through a series of electron carriers within multiprotein respiratory complexes (complex I to IV) to oxygen, therefore generating an electrochemical gradient that can be used by the F₁-F₀-ATP synthase (also named complex V) in the mitochondrial inner membrane to synthesize ATP. The organization and function of the ETC is a continuous source of surprises. One of the latest is the discovery that the respiratory complexes can assemble to form a variety of larger structures called super-complexes (SCs). This opened an unexpected level of complexity in this well-known and fundamental biological process. This review will focus on the current evidence for the formation of different SCs and will explore how they modulate the ETC organization according to the metabolic state. Since the field is rapidly growing, we also comment on the experimental techniques used to describe these SC and hope that this overview may inspire new technologies that will help to advance the field.
appear either as a monomer or as a dimer. Then, the transference of electrons between complexes occurs by the diffusion of the carriers CoQ and cyt c. In 2000, a study using Blue Native Gel Electrophoresis (BN-PAGE) demonstrated that complexes can interact and form large structures called super-complexes (SCs) recovering the idea of the solid model. Then it was proposed the existence of two types of SCs or respirasomes I + III₂ + IV (NADH-respirasome or N-respirasome) and III₂IV (CoQ respirasome or Q-respirasome) in a proportion of 2 N-respirasomes per Q-respirasome, to fit the overall 1 : 3 : 6 stoichiometries of complexes I : III : IV [7]. It was also postulated that those structures could be combined to form ‘respiratory strings’ as the physical concatenation of N- and Q-respirasomes forming long chains in the mitochondrial inner membrane [8-10]. The remaining observed associations like the abundant SC I + III₂ were considered broken parts of bigger entities. However, the existence of respiratory strings could not be experimentally supported and none of the hypothesized stoichiometry for the N- or Q-respirasomes has ever been documented. Instead, molecular [11-13] and structural evidence [14-17] supports the existence, of the N-respirasome as SC I + III₂ + IV (Figure 2a), that may further associate as megacomplex 2I + III₂ + 2IV [18] (Figure 2b). The Q-respirasome is observed as SC III₂ + IV and SC III₂ + IV₂ [11-13] (Figure 2a).
With quantitative variations between different sources of mitochondria, the variety of super assemblies between respiratory complexes observed by BN-PAGE is constant (Figure 2c). Recently the scientific debate has focused on the organization of COQ and cyto c pools [19], the impact of SC assembly for the activity of the

Figure 2. Respiratory complexes and SCs composition.
(a) Respiratory super-complexes are formed by different composition of complexes. Key factors are the expression of different subunit isoforms important for CIV dimerization (COX7A2, COX7A1, COX6A1, COX6A2), the assembly factors SCAF1 (important for CIII and IV interaction) and MCJ (inhibitor of CI and CIII assembly), MIM lipid composition (CL: cardiolipin, PE: phosphatidylethanolamine, PG: plasmalogen). (b) Respirasome alternative ternary interactions. N-respirasome may include a single copy of CIV2 instead of CIV. Moreover, the stoichiometry of megacomplex (2I + III2 + 2IV) is represented. The following pdb structures were used to develop this figure: CI (5xtd, Homo sapiens), CIII2 (5xte, Homo sapiens), CIV (5z62, Homo sapiens), CIV2 (1occ, Bos taurus), Loose N-respirasome (5j7y, Ovis aries), Tight N-respirasome (5j4z, Ovis aries) and Megacomplex (Homo sapiens, Bos taurus). (c) BN-PAGE of 2 h [35S]-methionine pulse labeled mtDNA encoded proteins wild-type mouse embryonic fibroblasts from and harvested after 24 h of chase.
complexes, the molecular mechanisms that drive the super assembly [2,20] and the bioenergetics and physiological role of SCs.

**How many super-complexes exist?**

The debate about the ETC organization [2] ended in 2008 with the proposal of the *plasticity model*, that considers the coexistence of complexes and SCs [21,22]. Thanks to a combination of genetic ablation of individual respiratory complexes and BN-PAGE, it was possible to identify SCs dependent on molecular interactions and discriminate them from spurious co-migration and breaking part of bigger structures [2,21,22]. In this model, CI, CII and CIV can be seen a monomer, CIII is always a dimer while and CIV can be also found as dimer. Further CI, CIII and CIV form pairs or trios with other respiratory complexes in significant proportions. Thus, besides the dimers of CIII and CIV and the N- and Q- respirasomes, additional binary associations to form SC I + III2, SC I + IV and SC I + IV2, whose function is uncertain, have been described [13] (Figure 2). Of note, the proportion of respiratory complexes appearing in free form or associated varies depending on the species [12,23–28], cell type [13,29–32], and physiological situation [24,34,35,28,33].

The first estimation of the distribution of complexes between free form and SCs was provided for bovine heart [36]. Thus, CI was mainly found in SCs with ≈15% in the free form [36], a proportion that is even lower in human cell lines [37] and human skeletal muscle [24]. For that reason, the existence of CI as an individual isolated complex was considered as artefact due to the solubilization of mitochondrial membrane with digitonin, a fundamental step for BN-PAGE analysis [7,38]. However, in rodents, bovine, zebrafish, Drosophila, fungus and plants, the persistent presence of free CI is observed [7,12,21,39–41] and can be modulated by different physiological stimuli [34,42]. Together with that, a careful kinetic analysis of CI assembly in human cells found that CI is fully assembled individually and quickly stabilized by super assembly [37]. All this data argues in favor of the existence of free CI.

Regarding bovine heart, more than 40% CIII was found as III2 while more than 80% of CIV was found as a monomer [36]. Mouse and rat brain mitochondria show very low levels of free CI and a higher proportion of the respirasome compared with other tissues. In most cases, CIV appears as a monomer with a small fraction as dimers or associated to SCs, more abundant in heart and skeletal muscle than liver. Brown adipose tissue is characterized by high levels of the Q-respirasome (III2 + IV) [43]. Now, the consensus is that BN-PAGE has cataloged two major classes of SCs in animal cells: binary (SC I + III2 and III2 + IV) and ternary (SC I + III2 + IV) for which there is experimental evidence that co-migration is due to true interaction (Figure 2c), since the elimination of one complex modifies the migration of other(s) [21]. In addition, the significant increase in resolution allowed by cryo-slicing BN-PAGE samples combined with Mass Spectrometry analysis (named in general ‘complexome analysis’) led to the proposal of two novel binary SCs: I + IV and III2 + IV2 [11], and recently, similar SCs were described also in zebrafish [12] and in mice [13]. The ternary structures were named N-respirasomes because they contain the three complexes that could allow a full respiratory chain [7], and their capacity to respire from NADH was later confirmed [21]. Besides the more commonly reported SCs, a frequently observed association between I + III2 of very high molecular weight can be observed that may be compatible with 2(I + III2) or the consequence of interaction with underdetermined components [21]. In addition, a gigantic structure of ring-shape named megacomplex 2I + III2 + 2IV has also been reported [18]. The isolation of this megacomplex from HEK293T cells calls for caution regarding the relevance of this megacomplex in more physiological contexts (Figure 2a,b). Nowadays, it is still not completely clear whether CII could interact with other respiratory complexes. Indeed, in mammalian, it is mainly found in free form and barely co-migrating to other complexes in a BN-PAGE especially if the solubilization conditions are not stringent [21]. Another hypothesis, from *in silico* modeling, suggests that CII can fit into the ring of the megacomplex but experimental evidence is lacking [18]. Even if not part of any SC, CII has a role in the modulation of SC assembly under oxygen concentrations and NADH/FADH2 ratio [34]. On contrary, in pea shoot, CII has been described associated to a megacomplex II,III,IV2 [44] whose function and role are still unknown. Although this is out of the scope of this review the observation of CII migrating in high molecular positions in BN-PAGE may be more related to the postulated formation of associations with other enzymes of the TCA cycle called TCA metabolon [45].

The detection of co-migration of several respiratory complexes in the BN-PAGE gel by immunoblot or in gel activity is insufficient to demonstrate that they physically interact [21]. In this regard, the co-migration of I + IV2 and I + III2 may be misunderstood as respirasome [46], Visualization and dynamically tracking the ETC organization *in situ* have started to show results [47–51], but may still need improvement to be able to track endogenous levels of SCs.
Mechanisms and factors of SCs assembly

Experiments of genetic modulation of complex subunits, the use of detergent-free methods and the resolution of SCs structures by cryo-electron microscopy (cryo-EM) together with the discovery that SCs assembly is a genetic regulated process, definitively prove the existence and relevance of SCs.

In 2012–2013 several groups proposed candidate proteins responsible for the assembly between CIII and CIV [42,52–54]. They were named Rcf1 and Rcf2 in yeast [52–54] and HIGD1 and HIGD2 in mammals. Rcf1 and Rcf2 are members of the hypoxia-induced gene (domain) 1 Hig1 family, and later studies demonstrated that they are not required for SC assembly rather than for the stability, activity [55,56] and assembly of CIV [57,58]. Therefore, the only bona-fide SC assembly factor required for III2 and IV interaction is the subunit COX7A2L renamed as Super Complexe Assembly Factor 1 (SCAF1), discovered in 2013 [42]. Its absence does not affect any aspect of either individual CIV or CIII assembly, stability, or function [2] but affects the N- and Q- respirasome structures.

An unexpected finding, that later was pivotal for unveiling SCAF1 role, revealed that the more commonly used inbred mouse strain (C57BL/6 and their sub-strains) harbors an in-frame micro-deletion that renders SCAF1 non-functional [42,59,60]. Therefore, this strain lacks the Q-respirasome. The role of SCAF1 in the formation of N-respirasome turns to be more complex but it was recently solved [13,61]. Detailed complexome analysis showed that two different N-respirasomes can be formed with different subunit composition of CIV, ether with SCAF1 or with COX7A2 [13,61]. The N-respirasome with SCAF1 has a structural attachment between III2 and IV while the one with COX7A2 lacks physical interaction between complexes III and IV [13]. Both versions of the N-respirasome are functionally distinct, the presence of SCAF1 increases NADH-dependent respiration and reduces reactive oxygen species (ROS) production [13] resulting in better response to severe fasting and exercise in mice and zebrafish [12,13]. On the contrary, SCAF1 depletion in human embryonic kidney cells (HEK293T) does not apparently affect OXPHOS performance [61], probably because they barely express SCAF1 in basal conditions and almost lacks naturally SCAF1 containing respirasome [61].

The available cryo-EM structures of the N-respirasome were obtained from a single tissue (heart) and lack of sufficient resolution to determine which subunit of the COX7A family (COX7A1, COX7A2 or COX7A2L/SCAF1) is present in the structure [14–16]. Proteomic analysis allows to discard the presence of COX7A1 since is preferentially found in complex for dimers in heart mitochondria [29] (Figure 2a). Therefore, it is uncertain which of the N-respirasomes (COX7A2 containing, SCAF1 containing or a mix of both) was analyzed [14–17]. Further improvement in resolution is required to address this point. Cryo-EM analysis also suggests the existence of two N-respirasomes with different degree of interaction of III2 and IV, the tight N-respirasome with contact between III2 and IV, and the loose N-respirasome and where no physical contact between III2 and IV is observed [14,16]. The gradual transformation of the tight form into the loose form in solution was confusing. The dynamicity of the III2 + IV interaction in the N-respirasome can justify also why the different areas of the is observed [14,16]. The gradual transformation of the tight form into the loose form in solution was confusing. The gradual transformation of the tight form into the loose form in solution was confusing.
The mechanism of I and III2 super assembly is mostly unknown, but several proteomic analyses strongly support that this interaction does not need any assembly factor [11,29,37]. Noteworthy, a protein called MCJ/DnaJC15 that interacts with complex I, reducing its activity and preventing its association into SC, has been proposed as negative regulator [64]. Several studies indicate that the lipid environment, specifically cardiolipin, determines the super assembly between CI and CIII [65–68]. In agreement with this, Tafazzin mutant, which affects cardiolipin synthesis, manifest the loss of SCs containing CI [67,69,70]. Also, mutations of Prohibitin [71] and Stomatin [72,73], impair the formation of CI and CIII interactions. The formation of SCs requires not only a favorable lipid environment and the participation of specific modulators, but also a well-defined cristae structure [74]. This was revealed by the loss of the SCs upon ablation of OPA1, a fundamental protein that regulates mitochondrial cristae formation, stability, and dynamics [75]. This multifaceted interaction between specific factors and the environment was illustrated by the deciphering of the mechanisms of SC formation induction by ER stress [33]. This induction required the ATF4 mediated activation of SCAF1 expression and the independent and parallel increasing of cristae density [33] (Figure 3).

**Figure 3. Metabolic adaptation of super-complexes.**
Super-complexes formation undergoes to adaptation upon different metabolic conditions. During development, the formation of super-complexes follows a genetically coordinated timing. Exercise improves super-complexes formation probably through a ROS/UPR/PERK mediated pathway. Endoplasmic reticulum stress response triggers SCs assembly through the PERK axis that activates both SCAF1 expression and cristae formation. The accumulative damage may be due to ROS increasing is responsible of super-complexes damage during aging and probably also in diabetes mellitus (DM) and heart failure. The massive dependency on anabolic reactions of cancer cells could be responsible of the super-complexes increase. Diet and in particular the ratio NADH/FADH2 modulate super-complexes distribution through RET. The different expression levels of SCAF1 in different tissues could be responsible of the variability in CIII2 + IV and CI + III2 + IV amount. Expression data obtained from Human Protein Atlas available from http://www.proteinatlas.org. The following pdb structures were used to develop this figure: CI (5xtd, Homo sapiens), CIII2 (5xte, Homo sapiens) and CIV (5z62, Homo sapiens).
**Functional relevance of SCs**

The findings that CI stability in mammals is impaired by the absence of CIII [76,77] and CIV or cyt c [78] led to the proposal that the SCs either stabilize the respiratory complexes [27,76,77] or serve as a platform where CI is assembled [38,79,80]. Recently, a series of experiments suggested that in the absence of either CIII or CIV, CI is quickly degraded due to the activation of retrograde electron transfer (RET) when the proportion of reduced CoQ is abnormally elevated [34] and it is rescued by the RET prevention [34] or the expression of an alternative oxidase capable of re-oxidizing CoQH2 (bypassing the role of CIII, cyt c and CIV together) [34]. Therefore, the stability of CI seems to be more dependent on the prevention of RET rather than on the actual formation of super-complexes. An additional and complementary explanation of the CI dependency on CIII was suggested by using human cell lines defective in CIII assembly [38]. These results suggested that the last step in the assembly in CI, the incorporation of the N-module (responsible for the NADH dehydrogenase activity) requires the previous interaction of the partially assembled CI with CIII [38]. Therefore, it is proposed that, in this cellular model, CI finish its assembly after the interaction with III2 [38,79].

In isolated I + III2, super assembly of CI and CIII impact on the capacity of CI to oxidize NADH [63] and diminishes the accessibility of external decylubiquinone (DQ) to CI [63], suggesting that the assembly of CI into SCs may partially protect it against RET and subsequently prevent its degradation. Along this line, the activation of SCs formation by the protein kinase R (PKR)-like endoplasmatic reticulum kinase (PERK), stabilizes CI and therefore supports the growth in galactose of cells harboring missense mutations [33]. In conclusion, the formation of SCs stabilizes CI, but this interaction does not seem to be mandatory to allow fully assembled and functional CI in all physiological scenarios. Thus, CI can be fully assembled independently of its interaction with CIII [13,34,37,38,76] or being its N-module added after super assembled with CIII [38,79].

A second functional role proposed for SC is the organization of the electrons flux to enhance respiratory activity preventing electron traffic jams [34,42,81], and minimize ROS production [13,32,82]. CI and CIII containing SCs harbor CoQ in their structure that is sufficient to trigger the oxidation of NADH to cyt c [38]. Two questions need to be addressed: (i) is the CoQH2, generated by CI, released to the membrane milieu and mixed and equilibrated with the CoQH2 generated by other enzymes (like CII) before being oxidized by CIII? (ii) is the CoQH2 generated outside the SCs capable to be oxidized by CIII assembled to SCs together with CI. The two questions refer to the existence or not of metabolic channeling between complex I and III2 in SCs. Metabolic channeling can be enforced by building permanently sealed protein tunnels within the SCs connecting the different reaction centres or by the differential partitioning of the intermediate metabolites that prevent its free diffusion. Structural analysis from different groups demonstrates that the cavity that contain the CoQ binding sites of CI and CIII in the I + III2 SC is not sealed, and that CoQ would be able to diffuse out of the SC [14–16,63]. However, the kinetic of the flux of electrons from CII to CIII [42] or from NADH to two different AOX enzymes [13,83] is negatively affected by the formation or activity of SCs I + III2. These results are in discrepancy with those published by Fedor et al. [84]. The discrepancy may rely on the very different experimental conditions used. Thus, the former was observed by in vivo genetic manipulation of the I + III2 proportion and by in vivo expression of AOX [13,42,83]. On the other hand, Fedor et al. [84] used heart derived submitochondrial particles to whom bacterial expressed recombinant AOX was added in vitro. The same question applies to cyt c in the context of CIII and CIV containing SCs. In this case all reports are coincident in showing that electron transfer between III2 and IV within the mammalian Q-respirasome is more efficient than between free III2 and free IV [62] confirming in this way the postulated of the plasticity model [22,42]. Interestingly, this view is in full agreement with the observations of the role of the yeast respirasome [85,86]. For a detailed discussion on the functional segmentation of CoQ and cyt c see [19,87].

Several mechanisms can account for impact of SCs in the partial segmentation of the CoQ and cyt c pools. In the case of CoQ the lipid milieu surrounding the SC and the rest of the membrane can favor its turnover within the SC by partitioning, while cyt c can be retained in the SC by electrostatic forces [62,85,86]. Additionally, in both cases the proximity between the reaction centres caused by super assembly can favor its turnover within the SC.

**SCs under different metabolic conditions**

It is known but often neglected that the electron equivalents generated in catabolism that fuel the mitochondrial electron transport chain may have two different flavors: the soluble NADH, that would be oxidized through CI, or FADH2 that delivers electrons to the mitochondrial electron transport chain bypassing CI...
Very important, different substrates generate different proportions of FADH$_2$/NADH (F/N) electrons, and this ratio plays a fundamental role in ROS generation by mitochondria [81,88]. In particular, the higher the F/N ratio the higher the CoQH$_2$/CoQ ratio and this could facilitate, in conjunction with high membrane potential, the induction of RET. This phenomenon reduces, both in cell lines [34] and mouse liver [42], the proportion of CI assembled with CIII therefore increasing the fraction of CII available for FADH$_2$ enzymes to favor fatty acids oxidation [34]. This suggests that the dynamic modulation of SCs’ proportion is a mechanism to efficiently adapt to the available carbon sources (Figure 3). Thus, in cultured cells the proportion of CI in free form vs SCs is higher when mitochondria oxidize pyruvate than when they oxidize fatty acid [34]. Likewise, liver mitochondria from overnight fasted male mice reduce the proportion of CI containing SCs [42].

The observation that SC formation is modified in response to ER-stress [33] and the impact of calcium and sodium in the regulation of MIM fluidity and OXPHOS activity [89] reinforce the role of the super assembly in adapting the metabolic response of the mitochondria. In this context, mitochondrial shaping proteins are important not only in the organelle dynamics but also in regulating the architecture of the cristae to allow SCs formation [33,74,75]. Under this perspective, it is expected that the organization in SCs would respond to metabolic differences induced by cell type, physiological changes and environmental cues that impacts on metabolism. This assumption is corroborated of growing body of evidence where SCs formation is enhanced or reduced to face different metabolic requirements. A great example is physical activity that increases the assembly of SCs in both humans and rats [24,35]. A more causative link comes from the demonstration that wild-type SCAF1 is necessary to achieve maximum exercise performance both in mice [13] and zebrafish [12]. Without a description of a precise molecular mechanism, we can only speculate that it can be induced by the activation of the PERK-ATF4 axis [33] and the concomitant expression of mitochondrial remodeling enzymes like OPA1 [90] (Figure 3).

Heart failure, and ischemia/reperfusion are characterized by a decrease in respirasome proportions and OXPHOS capacity [91–93] leading to the hypothesis that targeting SC formation could be a promising therapeutic approach for these pathologies. The brain offers an extraordinary example of metabolic cooperation between cells. The high energetic demand of neurons is satisfied by an efficient cooperation between astrocytes and neurons [94]. The former is glycolytic and generates and releases lactate that is taken by neurons to perform oxidative phosphorylation. Thus, the proportion of free vs super assembled CI in mice astrocytes is higher than in neurons [32]. This is accompanied by lower respiration activity and more ROS formation in astrocytes, providing a functional link between the organization of SCs and mitochondrial metabolism [32].

SCs organization is rearranged during immune response in macrophages. Mouse macrophages activated by live E. coli decrease the SCs containing CI and increase free CIII. Consequently, there is an increase in the activity of CII and glycerol-3-phosphate dehydrogenase that allows the use of FADH$_2$ as substrate [30]. Along the same line, it was reported that an incremental increase in CII activity in LPS-activated mouse macrophages together with a decrease in the NAD$^+$/NADH ratio activated RET, thereby producing ROS [95]. This mechanism potentiates the inflammatory state. While the number of studies on immune metabolism and SCs association are growing, little is known about the role of mitochondria in development and if and how the organization of SCs affects organogenesis is still an open question. Recently, a comprehensive analysis in zebrafish demonstrated that the SCs appear at early stages of embryonic development, and their number and associations increase progressively [96]. Few studies in mammals showed that SCs formation increases during adipogenic differentiation of human mesenchymal stem cells and, in embryonic mouse heart, SC formation starts at E11.5, concomitant with a burst in OXPHOS activity [96]. Further studies corroborate that mouse neonatal cardiomyocytes assemble less SCs than adult cells, supporting the idea that specific metabolic requirements induce the formation of SCs [29]. Interestingly, in cardiomyocytes lacking the transcriptional repressor CTCF, SCs formation is blunted accompanied with a disruption of the cardiac developmental program [96]. On the contrary, with age, SCs decrease at least in the brain [97] and heart [98] (Figure 3) suggesting that SCs formation goes together with the higher energetic demand of the tissues.

Two important pathologies are associated with changes in SCs association: diabetes and cancer. A unique study reported that SCs assembly is reduced in the rectus abdominis muscle of diabetes patients and this correlates with poor mitochondrial function [99], but the precise molecular links and whether the pathological condition can be ameliorated by compensating for the loss of SCs remains unclear. One therapeutic intervention could be a specific diet or the addition to food derived components. An example that shed light on this possibility, is hydroxytyrosol [100], a component of olive oil, that burst the formation of SC in rat muscles. Another example is dietary fatty acids whose consumption modifies the composition of the mitochondrial
Despite of evidence that they can modulate mitochondrial functions and complexes activities, there are no studies regarding the modulation of SC assembly by modulation of dietary fatty acids. We dare to anticipate that it is going to be an important field of research with potential applicability.

Cancer is a complex pathological condition characterized by altered metabolism, genetic mutations, distorted cell cycle, uncontrolled immunomodulatory factors, and disorganized tissue architecture. Nowadays, great effort is made in elucidating the heterogenous metabolic profiles found in cancer together with the dynamic rearrangement during tumor progression to accommodate and adapt to environmental cues. In this complex landscape, mitochondria and more specifically, ETC are critical players. However, the potential relevance of the SCs dynamics has not been systematically investigated. Interestingly, SCAF1/COX7A2L was discovered as a strong estrogen induced gene. Thus, tamoxifen resistance, a selective estrogen receptor modulator widely used to treat breast cancer, has been connected to SC formation. Her2high tumors, characterized by tamoxifen resistance, have more SCs and more CI dependent respiration. Others reported a correlation between reduction in SCs and elevated resistance. In favor of the pro-tumoral effect, it was estimated that expression of SCAF1 is an unfavorable prognosis marker for liver cancer. In agreement with that, it was reported that the deficiency in MCJ/DnaJC15, which negatively regulates SC formation, is associated with multidrug resistance in mouse and human breast cancer. SCAF1 was found overexpressed in clinical breast and endometrial cancer. Thus, it increases the stabilization of SCs together with a modulation of the metabolism of the TCA cycle intermediates that make tumor cells more resistant to hypoxia and this effect is dampened by the silencing of 2-oxoglutarate dehydrogenase complex. These results clearly show that the formation of SCs is strictly related to the modulation of metabolism especially in oestrogen dependent tumors. More studies are needed to understand the molecular mechanisms and the metabolic adaptation in response to modulation of SCs in tumorigenic conditions. In this direction, SCs is also indirectly regulated by p53 through DPYSL4, belonging to the collapsing response mediator family. DPYSL4 localizes with SCs in BN-PAGE suggesting that it may function in assembly or stability of SCs but further studies are needed to understand its mechanism of action and why SCs formation impact tumor growth in a lung-metastasis model.

**Perspectives**

- After a strong debate on the existence and function of SCs, a growing body of evidence confirm that SCs exist, and are essential elements for metabolism. Therefore, they are key elements in understanding and eventually manipulating metabolism.

- The assembly of SCs optimizes and organizes the flux of electrons, controls ROS production and modulates the activity of complexes. In vivo they confer metabolic advantage. However, we are just starting to understand these processes and many critical questions remain to be answered.

- In the future, studies on SCs dynamics, regulation and physiological implications will be pivotal to understand metabolism and develop strategies for therapeutic purposes. Moreover, an improvement of the techniques to track their dynamic composition in vivo will be mandatory.

**Competing Interests**

The authors declare that there are no competing interests associated with the manuscript.

**Author Contributions**

The authors contributed equally to all aspects of the article.
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Abbreviations
BN-PAGE, Blue Native Gel Electrophoresis; ETC, electron transport chain; OXPHOS, oxidative phosphorylation; PERK, (PKR)-like endoplasmic reticulum kinase; RET, retrograde electron transfer; ROS, reactive oxygen species; SCAF1, Super Complex Assembly Factor 1; SCs, super-complexes.

References
1 Chandell, N.S. (2015) Evolution of mitochondria as signaling organelles. Cell Metab. 22, 204–206 https://doi.org/10.1016/j.cmet.2015.05.013
2 Enríquez, J.A. (2016) Supramolecular organization of respiratory complexes. Annu. Rev. Physiol. 78, 533–561 https://doi.org/10.1146/annurev-physiol-021115-105031
3 Lobo-Jarne, T. and Ugalde, C. (2016) Respiratory chain supercomplexes: structures, function and biogenesis. Semin. Cell Dev. Biol. 76, 179–190 https://doi.org/10.1016/j.semcdb.2017.07.021
4 Kellin, D. and Hartree, E.F. (1947) Activity of the cytochrome system in heart muscle preparations. Biochem. J. 41, 500–502 https://doi.org/10.1042/bj0410500
5 Chance, B., Williams, G.R. and Hollunger, G. (1963) Inhibition of electron and energy transfer in mitochondria. I. Effects of Amytal, thiopental, rotenone, progesterone, and methylene glycol. J. Biol. Chem. 238, 418–431 https://doi.org/10.1016/S0021-9258(19)84014-0
6 Hackenbrock, C.R., Chazotte, B. and Gupte, S.S. (1986) The random collision model and a critical assessment of diffusion and collision in mitochondrial electron transport. J. Bioenerg. Biomembr. 18, 331–368 https://doi.org/10.1007/BF00743010
7 Schägger, H. and Pfeiffer, K. (2000) Supramolecular organization of the respiratory chains of yeast and mammalian mitochondria. EMBO J. 19, 1777–1783 https://doi.org/10.1093/emboj/19.8.1777
8 Wittig, I., Carrozzo, R., Santorelli, F.M. and Schägger, H. (2006) Supramolecular organization of the respiratory supercomplexes. Biochim. Biophys. Acta Bioenergetics 1757, 1066–1072 https://doi.org/10.1016/j.bbabio.2006.05.006
9 Buttema, J.B., Braun, H.-P., Boekema, E.J. and Koufi, R. (2009) Megacomplex organization of the mitochondrial oxidative phosphorylation system by structural analysis of respiratory supercomplexes from potato. Biochim. Biophys. Acta Bioenergetics 1787, 60–67 https://doi.org/10.1016/j.bbabio.2008.10.010
10 Wittig, I. and Schägger, H. (2009) Supramolecular organization of the mitochondrial respiratory chain in mammalian mitochondria. Biochim. Biophys. Acta Bioenergetics 1787, 672–680 https://doi.org/10.1016/j.bbabio.2008.12.016
11 Müller, C.S., Bildt, W., Haupt, A., Ellenrieder, L., Becker, T., Hunte, C. et al. (2016) Cryo-slicing blue native mass spectrometry (cSN-MS), a novel high resolution technology for complexome profiling. Mol. Cell. Proteomics 15, 669–681 https://doi.org/10.1074/mcp.M115.054080
12 García-Poyatos, C., Cogliati, S., Calvo, E., Hershman-Sazonov, P., Lagarrigue, S., Magni, R. et al. (2020) SCAF1 promotes respiratory supercomplexes and metabolic efficiency in zebrafish. EMBO Rep. 21, e50287 https://doi.org/10.15252/embr.202050287
13 Calvo, E., Cogliati, S., Hershman-Sazonov, P., Loureiro-López, M., Guàrdia, A., Casuso, R. et al. (2020) Functional role of respiratory supercomplexes in mice: SCAF1 relevance and segmentation of the qpool. Sci. Adv. 6, eaaz7509 https://doi.org/10.1126/sciadv.aaz7509
14 Letts, J.A., Fricke, A., and Sazonov, L.A. (2016) The architecture of respiratory supercomplexes. Nature 537, 644–648 https://doi.org/10.1038/nature19774
15 Gu, J., Wu, M., Guo, R., Yan, K., Lei, J., Gao, N. et al. (2016) The architecture of the mammalian respirasome. Nature 537, 639–643 https://doi.org/10.1038/nature19359
16 Sousa, J.S., Mills, D.J., Vonck, J. and Kühlbrandt, W. (2016) Functional asymmetry and electron flow in the bovine respiratory supercomplex. eLife 5, e21290 https://doi.org/10.7554/eLife.21290
17 Wu, M., Gu, J., Guo, R., Huang, Y. and Yang, M. (2016) Structure of mammalian respiratory supercomplex II112V1. Cell 167, 1598–1609.e10 https://doi.org/10.1016/j.cell.2016.11.012
18 Guo, R., Zong, S., Wu, M., Gu, J. and Yang, M. (2017) Architecture of human mitochondrial respiratory megacomplex II112V2. Cell 170, 1247–1257. e12 https://doi.org/10.1016/j.cell.2017.07.050
19 Hershman-Sazonov, P. and Enríquez, J.A. (2021) Functional segmentation of CoQ and cyt c pools by respiratory supercomplex superassembly. Free Radic. Biol. Med. 167, 232–242 https://doi.org/10.1016/j.freeradbiomed.2021.03.010
20 Mitenkov, D., Blaza, J.N., Larsson, N.-G. and Hirst, J. (2017) The enigma of the respiratory chain supercomplex. Cell Metab. 25, 765–776 https://doi.org/10.1016/j.cmet.2017.03.009
21 Acín-Peréz, R., Fernández-Silva, P., Peleato, M.L., Pérez-Martos, A. and PhD, J.A.E. (2008) Respiratory active mitochondrial supercomplexes. Mol. Cell 32, 529–539 https://doi.org/10.1016/j.molcel.2008.10.021
22 Acín-Peréz, R. and Enríquez, J.A. (2014) The function of the respiratory supercomplexes: the plasticity model. Biochim. Biophys. Acta Bioenergetics 1837, 444–450 https://doi.org/10.1016/j.bbabio.2013.12.009
23 Bundgaard, A., James, A.M., Harbou, M.E., Murphy, M.P. and Fago, A. (2020) Stable mitochondrial complexes II2 supercomplex interactions in reptiles versus homoeothermic vertebrates. J. Exp. Biol. 223, jeb223776 https://doi.org/10.1242/jeb.223776
24 Greggio, C., Jha, P., Kukarni, S.S., Lagarrigue, S., Broxley, N.T., Bottant, M. et al. (2017) Enhanced respiratory chain supercomplex formation in response to exercise in human skeletal muscle. Cell Metab. 25, 301–311 https://doi.org/10.1016/j.cmet.2016.11.004
25 Eubel, H. (2003) New insights into the respiratory chain of plant mitochondria. supercomplexes and a unique composition of complex II. Plant Physiol. 133, 274–286 https://doi.org/10.1104/pp.103.024620
26 Marques, I., Dencher, N.A., Videira, A. and Krause, F. (2007) Supramolecular organization of the respiratory chain in Neurospora crassa mitochondria. 

27 Stroh, A., Anderka, O., Pfeiffer, K., Yagi, T., Finel, M., Ludwig, B. et al. (2004) Assembly of respiratory complexes I, III, and IV into NADH dehydrogenase supercomplex stabilizes complex I in Paracoccus denitrificans. 

28 Arribat, I., Greppler, D., Lagarrigue, S., Richard, J., Gachet, M., Gut, P. et al. (2019) Mitochondria in embryogenesis: an organellogenesis perspective. 

29 Cogliati, S., Calvo, E., Loureiro, M., Guaras, A.M., Nieto-Arellano, R., Garcia-Poyatos, C. et al. (2016) Mechanism of super-assembly of respiratory complexes III and IV. Nature 539, 579–582. https://doi.org/10.1038/nature16517

30 Guadarrama, J., Acín-Pérez, R., Martínez-Cánovas, S., Enamorado, M., Ugolini, M., Nistal-Villín, E. et al. (2016) Mitochondrial respiratory chain adaptations in macrophages contribute to antibacterial host defense. Nat. Immunol. 17, 1037–1045. https://doi.org/10.1038/ni.3509

31 Lapuente-Brun, E., Moreno-Loshuertos, R., Acín-Pérez, R., Latorre-Pellicer, A., Colás, C., Balsa, E. et al. (2013) Supercomplex assembly determines supercomplex stability. 

32 Lopez-Fabuel, F., Riesebrother, J., Carabias-Carrasco, M., Almeida, A. and Bulafos, J.P. (2017) Mitochondrial complex I activity is conditioned by supercomplex I-III2-IV assembly in brain cells: relevance for Parkinson’s disease. Neurochem. Res. 42, 1676–1682. https://doi.org/10.1007/s11064-017-2191-2

33 Zhu, F., Yang, Z., Wang, F., Li, D., Cao, H., Tian, Y. et al. (2020) 4-Dimensional observation ER-mitochondria interaction in living cells under nanoscopy microscopy. Sci. Rep. 10, 4571. https://doi.org/10.1038/s41598-020-59585-7

34 Protasoni, M., Pérez-Pérez, R., Lobo-Jarne, T., Harbour, M.E., Ding, S., Peñas, A. et al. (2020) Respiratory supercomplexes act as a platform for supercomplex stability. 

35 Vukotic, M., Oeljeklaus, S., Wiese, S., Vögtle, F.N., Meisinger, C., Meyer, H.E. et al. (2012) Rcf1 mediates cytochrome oxidase assembly and regulates supercomplex stability. 

36 Schägger, H. and Pfeiffer, K. (2001) The ratio of oxidative phosphorylation complexes I–V in bovine heart mitochondria and the composition of respiratory chain supercomplexes. J. Biol. Chem. 276, 37861–37867. https://doi.org/10.1074/jbc.M106474200

37 Guerrero-Castillo, S., Baertling, F., Kownatzki, D., Wessels, H.J., Arnold, S., Brandt, U. et al. (2017) The assembly pathway of mitochondrial respiratory chain complex I. Cell Metab. 25, 126–139. https://doi.org/10.1016/j.cmet.2016.09.002

38 Cogliati, S., Calvo, E., Loureiro, M., Guaras, A.M., Nieto-Arellano, R., Garcia-Poyatos, C. et al. (2016) Mechanism of super-assembly of respiratory complexes III and IV. Nature 539, 579–582. https://doi.org/10.1038/nature16517

39 Guadarrama, J., Acín-Pérez, R., Martínez-Cánovas, S., Enamorado, M., Ugolini, M., Nistal-Villín, E. et al. (2016) Mitochondrial respiratory chain adaptations in macrophages contribute to antibacterial host defense. Nat. Immunol. 17, 1037–1045. https://doi.org/10.1038/ni.3509

40 Lapuente-Brun, E., Moreno-Loshuertos, R., Acín-Pérez, R., Latorre-Pellicer, A., Colás, C., Balsa, E. et al. (2013) Supercomplex assembly determines supercomplex stability. 

41 Zhu, F., Yang, Z., Wang, F., Li, D., Cao, H., Tian, Y. et al. (2020) 4-Dimensional observation ER-mitochondria interaction in living cells under nanoscopy microscopy. Sci. Rep. 10, 4571. https://doi.org/10.1038/s41598-020-59585-7

42 Lopez-Fabuel, F., Riesebrother, J., Carabias-Carrasco, M., Almeida, A. and Bulafos, J.P. (2017) Mitochondrial complex I activity is conditioned by supercomplex I-III2-IV assembly in brain cells: relevance for Parkinson’s disease. Neurochem. Res. 42, 1676–1682. https://doi.org/10.1007/s11064-017-2191-2

43 Moreno-Loshuertos, R. and Férnandez-Silva, P. (2021) Tissue specific modulation of respiratory chain supercomplex organization. J. Biol. Chem. 297, 26453–26461. https://doi.org/10.1074/jbc.M120075200

44 Ukolova, I. (2014) The composition of pea mitochondrial supercomplexes under cold conditions. Biochim. Biophys. Acta Bioenergetics 1837, e32 https://doi.org/10.1016/j.ubioc.2014.05.311

45 Wu, F. and Minteer, S. (2015) Kreb’s cycle metabolism: structural evidence of substrate channeling revealed by cross-linking and mass spectrometry. Angew. Chem. Int. Ed. Engl. 54, 1851–1854. https://doi.org/10.1002/anie.201400336

46 Sun, D., Li, B., Qiu, R., Fang, H. and Lyu, J. (2016) Cell type-specific modulation of respiratory chain supercomplex organization. Int. J. Mol. Sci. 17, 926. https://doi.org/10.3390/ijms17060926

47 Shtukenberg, M.V., Dykov, D., Busch, K., Streeker, V., Wittig, I. and Breder-Hahn, J. (2010) Determination of protein mobility in mitochondrial membranes of living cells. Biochim. Biophys. Acta Bioenergetics 1798, 2022–2032. https://doi.org/10.1016/j.bbamem.2010.07.016

48 Mustier, B., Kohl, W., Wittig, I., Streeker, V., Joos, F., Haase, W. et al. (2010) Respiratory chain complexes in dynamic mitochondria display a patchy distribution in live cells. PLoS ONE 5, e11910 https://doi.org/10.1371/journal.pone.0011910

49 Appelhans, T., Richter, C.P., Wilkening, V., Hess, S.T., Pfeiffer, J. and Busch, K.B. (2012) Nanoscale organization of mitochondrial microcompartments revealed by combining tracking and localization microscopy. Nano Lett. 12, 610–616. https://doi.org/10.1021/nl203434a

50 Wilkening, V., Kohl, W. and Busch, K. (2013) Restricted diffusion of OXPHOS complexes in dynamic mitochondria delays their exchange between cristae and engenders a transitory mosaic distribution. J. Cell Sci. 126, 103–116. https://doi.org/10.1242/jcs.108852

51 Rieger, B., Shalataev, D.N., Söhnle, A.-C., Kohl, W., Duwe, P., Mulkidjanian, A.Y. et al. (2017) Lifetime imaging of GFP at CoxVIIIa reports respiratory supercomplex assembly in live cells. Sci. Rep. 7, 46055 https://doi.org/10.1038/srep46055

52 Strogglová, V., Fumuss, A., Robb-McGrath, M., Garlich, J. and Stuart, R.A. (2012) Rcf1 and Rcf2, members of the hypoxia-induced gene 1 protein family, are critical components of the mitochondrial cytochrome b1-cytochrome c oxidase supercomplex. Mol. Cell. Biol. 32, 1363–1373. https://doi.org/10.1128/MCB.06369-11

53 Vukotic, M., Oeljeklaus, S., Wiese, S., Vogtle, F.N., Meisinger, C., Meyer, H.E. et al. (2012) Rcf1 mediates cytochrome oxidase assembly and respiration rate formation, revealing heterogeneity of the enzyme complex. Cell Metab. 15, 336–347. https://doi.org/10.1016/j.cmet.2012.01.016

54 Chen, Y.-C., Taylor, E.B., Dephoure, N., Heo, J.-M., Tonhato, A., Papandreou, I. et al. (2012) Identification of a protein mediating respiratory supercomplex stability. Cell Metab. 15, 348–360. https://doi.org/10.1016/j.cmet.2012.02.006

55 Garlich, J., Streeker, V., Wittig, I. and Stuart, R.A. (2017) Mutational analysis of the RQRQ motif in the yeast Hig1 type 2 protein Rcf1 reveals a regulatory role for the cytochrome c oxidase complex. J. Biol. Chem. 292, 5216–5226. https://doi.org/10.1074/jbc.M116.758045
116 Watanabe, T., Inoue, S., Hiroi, H., Orimo, A., Kawashima, H. and Muramatsu, M. (1998) Isolation of estrogen-responsive genes with a CpG island library. Mol. Cell. Biol. 18, 442–449 https://doi.org/10.1128/MCB.18.1.442

117 Rohlenova, K., Sachaphibulkij, K., Stursa, J., Bezawork-Geleta, A., Blecha, J., Endaya, B. et al. (2017) Selective disruption of respiratory supercomplexes as a New strategy to suppress Her2 high breast cancer. Antioxid. Redox Signal. 26, 84–103 https://doi.org/10.1089/ars.2016.6677

118 Tomková, V., Sandoval-Acuña, C., Torrealba, N. and Truksa, J. (2019) Mitochondrial fragmentation, elevated mitochondrial superoxide and respiratory supercomplexes disassembly is connected with the tamoxifen-resistant phenotype of breast cancer cells. Free Radic. Biol. Med. 143, 510–521 https://doi.org/10.1016/j.freeradbiomed.2019.09.004

119 Ikeda, K., Hiroe-Inoue, K., Suzuki, T., Hobo, R., Nakasato, N., Takeda, S. et al. (2019) Mitochondrial supercomplex assembly promotes breast and endometrial tumorigenesis by metabolic alterations and enhanced hypoxia tolerance. Alat. Commun. 10, 4108 https://doi.org/10.1038/s41467-019-12124-6

120 Nagano, H., Hashimoto, N., Nakayama, A., Suzuki, S., Miyabayashi, Y., Yamato, A. et al. (2018) p53-inducible DPYSL4 associates with mitochondrial supercomplexes and regulates energy metabolism in adipocytes and cancer cells. Proc. Natl Acad. Sci. U.S.A. 115, 8370–8375 https://doi.org/10.1073/pnas.1804243115