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Towards a functional definition of the mitochondrial human proteome



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The Human Proteome Organization (HUPO) has promoted in recent years a concerted action aimed at the full characterization of the human proteome. Two measures have been put forward in parallel. First, the chromosome-centric initiative (C-HPP) aimed at joining the efforts of several groups with a country-based strategy to unravel the human proteome to reduce to a minimum, and possibly to eliminate, the number of proteins that did not show evidence at the protein level in the dedicated database such as NeXtProt [1,2]. Second, the "biology and disease driven" initiative (B/D-HPP) aimed at functional clustering of proteins around a significant biological issue [3]. To address some of these gaps, several consortia were formed, focusing on the human eye, diabetes, pediatric diseases, liver, brain, cancer, infectious diseases, autoimmune disorders, epigenetics, among others. A great contribution to these initiatives was provided by the technological pillars: the Human Proteome Atlas (HPA), the bioinformatics, and the mass spectrometry technological advancement study groups [4].

The two HPP programs are strongly intersected; in fact, the effort to consolidate the current available tools to identify and

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quantify all the genome-encoded proteins is providing the fundamental framework to highlight novel molecular targets for the B/D efforts. The Italian HPP group did a great effort in bringing laboratories across the globe to take part in the Italian Proteomics Association (ItPA) project to unravel the mitochondrial human proteome project (mt-HPP) as part of both C-HPP and B/D-HPP initiatives [5].

Mitochondria are essential organelles for the life and death of cells, and several human disorders are associated to mitochondrial dysfunction. While some human diseases are strictly linked to mutations in the mitochondrial genome, many others are someway connected to mitochondrial functionality by defects in the sequence of proteins encoded by nuclear chromosomes that are imported into mitochondria through their mitochondrial transfer sequence (MTS) [6]. Mutations of cytosolic proteins that regulate the elimination of impaired mitochondria segregate with severe degenerative disorders, such as Parkinson's disease. For instance, a number of mutations in the PARK2, PINK1 and PARK7 genes have been shown to cause mitochondrial parkinsonism [7], although PARK2 and PARK7 gene products (parkin and DJ-1, respectively) do not belong, strictly speaking, to the mitochondrial proteome, as they are not imported into mitochondria. In fact, several recent reports highlighted the relevance of cytosolic

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molecular machineries and non-mitochondrial subcellular structures in the governance of mitochondrial functions. Although mitochondria are usually associated to their role in cellular energetics, they exert a number of functions that are not confined to ATP production. In particular, calcium ion homeostasis is regulated by a strict functional and spatial connection between mitochondria and the endoplasmic reticulum, and glycolysis has been shown to be coupled to ATP production through the spatial recognition of the glycolytic enzyme hexokinase by the outer membrane porin VDAC1 [8–10].

In recent years, several papers underlined the role of mitophagy, the disposal of dysfunctional mitochondria by the autophagic machinery on a whole-cell level. Parkin and PINK1 assumed a fundamental role in targeting mitochondria to the autophagosome through ubiquitination of outer surface membrane proteins, although this mechanism still seems to be controversial [11,12]. Additionally, the maintenance of a well-organized mitochondrial network was shown to be a peculiar actor in assuring full functionality of the mitochondrion [13]. To this purpose, a thorough assessment of the mitochondrial interactome should fill the gaps in our knowledge of the complex biological processes sustained by the mitochondrial proteome [14,15].

It is of great relevance to extend our present view of the mitochondrial proteome not only to those proteins that are encoded by the mitochondrial or nuclear genomes, but also to their interactors that take part in mitochondrial functionality, maintenance, dynamics and metabolism [16]. Thus, the integration of information obtained within the C-HPP and B/D-HPP actions with interactomics data deposited in the IntAct database [17] and protein annotation in the NeXtProt database is a priority (Fig. 1). To date, 13 proteins are listed in the NeXtProt database (September 2015 release) as mitochondrial-encoded, whereas another 1869 proteins were annotated as mitochondrial proteins. Among

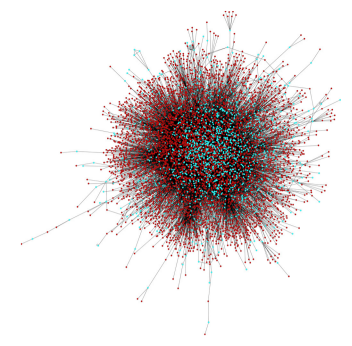


Fig. 2. The mitochondrial human proteome from a functional perspective. Cyan nodes represent proteins encoded by the mitochondrial genome or translocated to the mitochondrion. Protein identities were obtained by querying the NeXtProt database. Red nodes represent interactors of the cyan nodes as obtained by querying the IntAct database using the PSICQUIC query module embedded in Cytoscape.

them, 1277 entries were indicated to be evident at the protein level, with 592 remaining missing proteins. A search in the IntAct database allowed us to identify a broad panel of interactors that

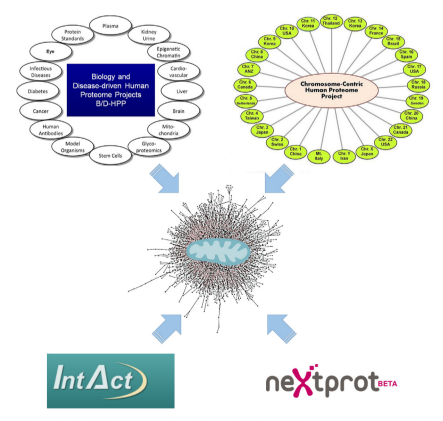


Fig. 1. A functional definition of the human mitochondrial proteome requires the integration of several efforts. From top-left, clockwise: functional studies from the B/D-HPP; a full coverage of the human proteome by C-HPP; protein evidence and functional annotation in the NeXtProt database; interactomics annotation from the IntAct database.

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