Regulation of cellular senescence by eukaryotic members of the FAH superfamily – A role in calcium homeostasis?

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\textbf{A B S T R A C T}

Fumarylacetoacetate hydrolase (FAH) superfamily members are commonly expressed in the prokaryotic kingdom, where they take part in the committing steps of degradation pathways of complex carbon sources. Besides FAH itself, the only described FAH superfamily members in the eukaryotic kingdom are fumarylacetoacetate hydrolase domain containing proteins (FAHD) 1 and 2, that have been a focus of recent work in aging research. Here, we provide a review of current knowledge on FAHD proteins. Of those, FAHD1 has recently been described as a regulator of mitochondrial function and senescence, in the context of mitochondrial dysfunction associated senescence (MiDAS). This work further describes data based on bioinformatics analysis, 3D structure comparison and sequence alignment, that suggests a putative role of FAHD proteins as calcium binding proteins.

1. Introduction

1.1. Identification of FAHD1 as regulator of mitochondrial function

In 1959 and 1974, Corwin and Wojtczak identified a mitochondrial oxaloacetate decarboxylase from rat liver (Corwin, 1959; Anna and Wojtczak, 1974). This was about 60 years ago, and until recently the identity of the enzyme remained unclear. In 2007 high resolution 2D gels of mitochondrial preparations from young and senescent human umbilical vein endothelial cells (HUVEC) were prepared using the ProteoTope\textsuperscript{a} technique (Grobe et al., 2007). This revealed an age-related difference in isoelectric point of about 0.4 pI units for two protein spots (\#1756 and \#1780/1784) (Grobe et al., 2007; Etemad et al., 2019), suggesting differences in post-translational modification of the associated protein with cellular senescence. Mass spectrometric analysis identified the protein as fumarylacetoacetate hydrolase domain containing protein 1 (FAHD1) (Pircher et al., 2011). In 2011 and 2015, Pircher et al. were able to identify FAHD1 as acylpyruvate hydrolase (ApH) and oxaloacetate decarboxylase (ODx), which is localized in mitochondria (Pircher et al., 2011) and belongs to the broad FAH superfamily of enzymes (Pircher et al., 2011; Kang et al., 2011; Hong et al., 2020; Pircher et al., 2015; Timm et al., 1999; Bateman et al., 2001). The localization of FAHD1 in mitochondria (Pircher et al., 2011) and its ODx activity rendered a model of FAHD1 acting as regulator of oxaloacetate levels in the TCA cycle (Etemad et al., 2019; Pircher et al., 2015; Jansen-Duerr et al., 2016), which was accompanied by the description of the FAHD1 catalytic mechanism (Weiss et al., 2018a). Work with the model organism Caenorhabditis elegans provided first support for this hypothesis, as deletion of fahd-1 induced severe mitochondrial dysfunction and impaired locomotion activity (Taferner et al., 2015). Recent work linked FAHD-1 activity to serotonin signaling in the nematode (Baraldo et al., 2019). Work with human endothelial cells (HUVEC) displayed that depletion of FAHD1 inhibits mitochondrial electron transport chain (ETC) and induces cellular senescence in human endothelial cells (Petit et al., 2017). This enabled the hypothesis of FAHD1 being a regulator of cellular senescence via regulation of the mitochondrial ETC (Etemad et al., 2019) in the context of mitochondrial dysfunction associated senescence (MiDAS) described previously by us (Steckl et al., 2006) and others (Wiley et al., 2016).

Oxaloacetate decarboxylases are mainly known from prokaryotic organisms, where membrane-bound (Lietzau and St Maurice, 2014) and soluble variants exist (Klafl and Eikmanns, 2010). The membrane-bound variants generally depend on sodium ions and biotin, whereas the soluble variants depend on bivalent metal cations (Weiss et al.,...
chains are provided for binding of divalent cations (the FAH superfamily of proteins, highly conserved carboxylate side activity is now well understood in the eukaryotic members of the superfamily (Weiss et al., 2018a), even raised the idea of the eukaryotic FAHD1 being a hybrid of related prokaryotic precursor proteins (Weiss et al., 2018b). Recent work by Hong et al. (Hong et al., 2020) supports this idea via a phylogenetic tree analysis of FAH superfamily enzymes.

However, the exact role of FAH and FAHD1 proteins, and of FAHD1 in particular, is not fully revealed to date. Here, we provide a review of collected data on FAH and FAH superfamily members, describing FAH and FAH superfamily members as a regulator of the TCA cycle flux in the context of mitochondrial dysfunction associated senescence. We further present conclusive data obtained via bioinformatic analyses, in order to hypothesize a secondary role of FAH superfamily members as possible calcium binding protein. Published links between calcium metabolism, mitochondrial dysfunction, and cellular senescence are highlighted. This model will extend the role of FAH superfamily members as a regulator of the TCA cycle flux by suggesting multiple physiological functions of FAH superfamily enzymes in eukaryotes.

1.2. FAHD1 catalytic mechanism revealed by structural studies and site directed mutagenesis

FAH1 acts bi-functional as ApH and ODx (Weiss et al., 2018a). While ApH activity is common for the FAH superfamily of enzymes in prokaryotes (Hong et al., 2020), ODx activity is not common in the prokaryotic part of the family (except for individual members such as Cg1458 (Ran et al., 2013, 2011) in Corynebacterium glutanicum). ODx activity is now well understood in the eukaryotic members of the superfamily (Weiss et al., 2018b), in particular for FAH superfamily members, while the role of ApH activity in the metabolism of eukaryotes remains elusive.

The postulated mechanism for FAH1 catalytic activity (Weiss et al., 2018a) was substantiated by experimental data. Mutations of particular amino acids by replacement with alanine create enzymatic forms with strongly decreased ODx activity, which are often inactive for the hydrolysis of acylpyruvates (Weiss et al., 2018a). In all enzymes of the FAH superfamily of proteins, highly conserved carboxylate side chains are provided for binding of divalent cations (e.g. Mg$^{2+}$, Ca$^{2+}$, Mn$^{2+}$, Zn$^{2+}$, Cu$^{2+}$) (Hong et al., 2020; Weiss et al., 2018b). However, for execution of the specific catalytic functions FAH superfamily members prefer distinct metals. For FAH, Ca$^{2+}$ and Mg$^{2+}$ are functional metal ions. FAH1 shows highest catalytic activity with Mg$^{2+}$ and Mn$^{2+}$ as cofactors (Pircher et al., 2011). The metal cofactor (Mg$^{2+}$) is held in place by the side chains of the three amino acids E71, E73 and D102 (Weiss et al., 2018a). The substrates of FAH1, oxaloacetate (OAA) as well as acylpyruvates (Ap), adopt different forms in varying ratio depending on the prevailing pH-value. Under mitochondrial pH of about 7.8 Ap and OAA are competent to bind tightly in divalent binding mode to the cofactor Mg$^{2+}$ of FAH1. Upon this primary binding event of the substrate, FAH1 acquires catalytic competence through backbone-flip induced lid closure (Weiss et al., 2018a). This event structures the disordered region of the apo-enzyme and isolates the catalytic cavity from the mitochondrial environment. Structuring of the disordered region induces a short helical region (Weiss et al., 2018a).

Helix residues E33 and H30 form a well-known catalytically competent acid-base dyad which interacts through hydrogen bonding with an isolated water molecule in the catalytic center (Weiss et al., 2018a). To prepare for the break of the C$^2$–$^3$ bond, the enzyme has to provide a conformational control over the bound substrates via Q109. The corresponding mutation Q154A in Cg1458 (Ran et al., 2013) abolished ODx activity. R106 forms hydrogen bonds with E73 and Q109, which is a key feature for maintaining the tertiary structure of the binding pocket (Weiss et al., 2018a). K123 plays a significant role as proton source in the FAH1 catalytic mechanism. Accordingly, substitution of K123 by alanine creates inactive forms both for ApH and ODx activities (Weiss et al., 2018a).

Deliberate modulation of FAH1 catalytic activity by selective single-point mutation helps to further understand the role of FAH1 in mitochondria and prepares for future work with in vivo models. Comparing the activity of FAH1 with respect to the wild type in nematode and mouse will provide evidence for the postulated downstream effects. In parallel, current attempts to develop small molecules with the ability to increase or decrease FAH1 catalytic activity aim at translational strategies to fine tune FAH1 activity in particular physiological and pathological conditions.

2. FAH1 and FAHD2: unequal members of the eukaryotic FAH superfamily

2.1. FAH1 and FAHD2 proteins share the FAH fold

Homology search and sequence analysis of FAH1 with proteins encoded in the genome of mammals revealed a high level of 97 % sequence identity with a putatively cytosolic enzyme: FAH domain containing protein 2 (FAHD2), which is expressed in the human genome in two unrelated versions (a, and b). Both hFAHD2a and hFAHD2b are encoded on human chromosome 2 (GRCh38:CM000664.2) (Uhlen et al., 2015, 2005). hFAHD2a is transcribed in direct sense (95,402,721 – 95,416,616) and hFAHD2b in reverse (97,083,583 – 97,094,882).

We found 4 active transcripts for hFAHD2a and 2 active transcripts for hFAHD2b. In both cases two of the active transcripts encode the same protein information, which leads to three forms of hFAHD2a (Q96GK7, C9JGM0 and C9JSB6) and only one form of hFAHD2b (Q6P213) (Uhlen et al., 2015, 2005). Transcripts 2 and 3 of hFAHD2a (C9JGM0 and C9JSB6) do not include the FAH fold (see Fig. 1), so only transcript 1 of hFAHD2a (Q96GK7) and the one transcript of hFAHD2b (Q6P213) display homology with hFAH1 (Q6P587). We conclude that both hFAHD2a and hFAHD2b are homologs to FAH1. Of interest, sequence comparison of transcript 1 of hFAHD2a with hFAHD2b reveals a difference in only 6 amino acids. The question of why the human genome encodes two such similar proteins on different parts of the same chromosome remains elusive.

The protein structure of hFAHD2a and hFAHD2b is yet unreported, however, Swiss-Model (Waterhouse et al., 2018) homology modelling of the protein structure of hFAHD2a (transcript 1, Q96GK7) reveals a strong structural similarity with FAH1 (see panels A and B of Figure S3). All critical amino acids and structure motifs, that have been identified to be of importance for the catalytic activity of FAH1, are fully conserved (see Fig. 1). As a result of similarities with FAH1, Mg$^{2+}$ and Mn$^{2+}$ have been inferred as cofactors, and present data allows for the hypothesis of a similar enzymatic activity. Human FAH2 manifests an N-terminal part, which is not present in human FAH1 and which probably confers to the protein a strong hydrophobic character (see Fig. 1). In fact, this protein fragment also comprises TOM20 sites, which have been found via bioinformatics comparison of amino acid sequences (Holzknecht et al., 2018; Dorigati et al., 2018) (see Table 1 and section 2.3). The TargetP-2.0 (Almagro Artemes et al., 2019) server predicts the presence of a mitochondrial transit peptide (mTP) (see panel C of figure S3) around L14 of FAH2a and FAH2b, but not in the sequence of FAH1.

Human FAH2a was found to be highly expressed in tissue of liver, testes and thyroid (Uhlen et al., 2015, 2005), and seems to be overexpressed in cancer tissue compared to benign tissue in different types of cancer such as colorectal, breast, prostate, lung and liver cancer (Uhlen et al., 2015, 2005). Subcellular localization of FAH2a and FAH2b has yet to be investigated. While we have collected important information on FAH1 structure and activity, FAH2 is highly understudied. Scarce data is available for its catalytic activity, subcellular localization and expression (Fagerberg et al., 2014). A detailed functional characterization of FAH2a will be required to increase our
understanding of the overall role of FAHD proteins.

A survey of mitochondrial TCA cycle enzymes is given in Table 2, comparing the reported structure and predicted stability in solution at physiological conditions. Structure and general protein information has been obtained from the UniProt (Wasmuth and Lima, 2017) database. Theoretical pI and stability predictions have been computed using the ProtParam (Gasteiger et al., 2005) server. FAHD proteins are predicted to be unstable (Table 2, marked in red), however, FAHD1 is understood to form a soluble and catalytically active homodimer (Pircher et al., 2011, 2015; Weiss et al., 2018a; Manjasetty et al., 2004), whereas all other unstable proteins are part of larger protein complexes (Wasmuth and Lima, 2017) (Table 2, marked in green).

2.2. Subcellular localization of FAHD proteins: mitochondria and more?

Subcellular localization of FAHD1 was assessed via immunofluorescence by the Human Protein Atlas (Uhlen et al., 2005; Fagerberg et al., 2014; Uhlen et al., 2010). Using antibodies HPA043534 and CAB025530, FAHD1 was described to be localized primarily in mitochondria with a potential secondary localization in the nucleoplasm. The localization of potential interaction partners of FAHD1, as listed in the BioPlex (Huttlin et al., 2015) network (Table 3, Fig. 2; see also below), generally matches the data reported for FAHD1 subcellular localization, i.e., mitochondria and nucleoplasm; moreover, this annotation is also supported by information about localization and function of the interacting proteins, as gathered from the Human Protein Atlas (Uhlen et al., 2005; Fagerberg et al., 2014; Uhlen et al., 2010) and the UniProt (Wasmuth and Lima, 2017) database.

A survey of predicted mitochondrial targeting sequences and their cleavage sites using the MitoFates (Fukasawa et al., 2015) server is given in Table 1. FAHD proteins display TOM20 binding sites, which have been found via bioinformatics comparison of amino acid sequences (Holzknecht et al., 2018; Dorigatti et al., 2018) (see Table 1). TOM20 subunits form a hydrophobic binding pocket in the outer mitochondrial membrane and are central components of the TOM receptor complex (Seki et al., 1995), that is responsible for the recognition and translocation of mitochondrial pre-proteins synthesized in the cytosol or close to the outer mitochondrial membrane (Lesnik et al., 2015) (see section 2.1).

Both FAHD1 and FAHD2 display sites for proteolytic cleavage of the targeting signal, performed by mitochondrial processing peptidase (MPP) and for cleavage of destabilizing N-terminal amino acid residues by intermediate cleaving peptidase 55 (ICP55), which is critical for stabilization of the mitochondrial proteome (Wasmuth and Lima, 2017) (see also Table 1). However, a possible cleavage of FAHD1 by MPP at amino acids N26 and Y27 would destroy the catalytic domain that is required for a functional protein (Weiss et al., 2018a), which appears unlikely. Cleavage of FAHD2 proteins by MPP and ICP55 is plausible, as also the TargetP-2.0 (Almagro Armenteros et al., 2019) server predicts the presence of a conserved mitochondrial transit peptide sequence (mTP CS) (see panel C of Figure S3) around L14 of FAHD2a and FAHD2b, but not in the sequence of FAHD1.

Fig. 1. Multiple sequence alignment of human FAHD2a, FAHD2b and FAHD1 isoforms. Human FAHD2 is expressed in two very similar, yet independent forms: FAHD2a and FAHD2b. Three active transcripts can be found for FAHD2a, and one for FAHD2b. Human FAHD1 is expressed in three isoforms. FAHD2a seems to be a hybrid form, consisting of a highly hydrophobic N-terminal sequence of 80 amino acids, fused to the actual FAHD protein. Transcripts 2 and 3 of FAHD2a translate to only the hydrophobic part, for which only transcript 1 of FAHD2a and FAHD2b translate to real FAHD proteins (see text). FAHD2a transcript 1 and FAHD2b differ in 6 amino acids marked with red boxes. FAHD proteins display TOM20 sites, which have been found via bioinformatics comparison of amino acid sequences (Holzknecht et al., 2018; Dorigatti et al., 2018), as well as sites for proteolytic cleavage of the targeting signal, performed by mitochondrial processing peptidase (MPP) and for cleavage of destabilizing N-terminal amino acid residues by intermediate cleaving peptidase 55 (ICP55), which is critical for stabilization of the mitochondrial proteome (Wasmuth and Lima, 2017) (see also Table 1). However, a possible cleavage of FAHD1 by MPP at amino acids N26 and Y27 would destroy the catalytic domain that is required for a functional protein (Weiss et al., 2018a), which appears unlikely. Cleavage of FAHD2 proteins by MPP and ICP55 is plausible, as also the TargetP-2.0 (Almagro Armenteros et al., 2019) server predicts the presence of a conserved mitochondrial transit peptide sequence (mTP CS) (see panel C of Figure S3) around L14 of FAHD2a and FAHD2b, but not in the sequence of FAHD1.
2.3. FAHD proteins are subject to differential mitochondrial import mechanisms

Proteins synthesized in the cytosol are imported into mitochondria via the general import pore (Lesnik et al., 2015; Walther and Rapaport, 2009), a multi-protein complex involving Tom5, Tom6, Tom7, Tom20, Tom22, Tom40, and Tom70. On the other hand, precursors of so-called signal-anchored proteins are imported to the mitochondria by a different mechanism (Ahting et al., 2005). Localization of FAHD1 in mitochondria despite the lack of a recognizable mitochondrial pre-sequence may suggest the presence of such a signal-anchor in FAHD1. The UniProt (Wasmuth and Lima, 2017) database lists curated (reviewed) entries of human proteins with signal-anchor motifs (keyword Signal-anchor KW-0735). BLASTp analysis of human FAHD1 and established signal anchor proteins displays significant sequence similarities with 8 entries, mapping to 4 proteins and their isoforms: Lactosylceramide

Table 1

| Enzyme | UniProt-Spec | Probability of pre-sequence | Mitochondrial pre-sequence | Cleavage site | Positions for TOM20 recognition motifs |
|--------|--------------|-----------------------------|-----------------------------|--------------|----------------------------------------|
| CS     | CSY_HUMAN    | 0.996                       | yes                         | 25(MPP)      | 7-11                                   |
| ACO    | ACON_HUMAN   | 0.995                       | yes                         | 19(MPP)      | 11-15                                  |
| IDH2   | IDHP_HUMAN   | 0.993                       | yes                         | 38(MPP), 39(kp55) | 4-8,58-62                             |
| IDH3   | IDH3A_HUMAN  | 0.961                       | yes                         | 26(MPP), 27(kp55) | 10-14,50-54                           |
|        | IDH3B_HUMAN  | 0.997                       | yes                         | 25(MPP), 33(oct1) | 10-14,31-35,63-67,79-74               |
|        | IDH3G_HUMAN  | 0.997                       | yes                         | 38(MPP)      | 2-6,12-16,17-71                       |
| OGDc   | OGD1_HUMAN   | 0.996                       | yes                         | 39(MPP), 40(kp55) | 8-12,90-93                           |
|        | OGD2_HUMAN   | 0.999                       | yes                         | 59(MPP), 67(oct1) | 8-4,57-61                             |
|        | OGD3_HUMAN   | 0.966                       | yes                         | 34(MPP), 35(kp55) | 7-11,13-17,18,22-90-94               |
| SUC(A/G)| SUC_A_HUMAN  | 0.421                       | yes                         | 23(MPP), 27(kp55) | 7-11,12-16,16-60                     |
|        | SUCB2_HUMAN  | 0.964                       | yes                         | 22(MPP), 33(kp55) | 7-11,24-28                            |
|        | SUCB3_HUMAN  | 0.826                       | yes                         | 52(MPP), 53(kp55) | 7-11,13-17                           |
| SDH    | SDHA_HUMAN   | 0.995                       | yes                         | 32(MPP), 40(oct1) | 7-11,13-17,18,22-90-94               |
|        | SDB_HUMAN    | 0.963                       | yes                         | 28(MPP)      | 7-11,13-17,18,22-90-94               |
|        | C50_HUMAN    | 0.992                       | yes                         | 51(MPP), 52(kp55) | 7-11,13-17,18,22-90-94               |
|        | DHSO_HUMAN   | 0.996                       | yes                         | 28(MPP)      | 7-11,13-17,18,22-90-94               |
| FH     | FUMH_HUMAN   | 1.000                       | yes                         | 44(MPP)      | 1-5,4-8,41-45,92-96                   |
| MDH2   | MDHM_HUMAN   | 0.999                       | yes                         | 16(MPP), 24(oct1) | 10-14                                  |
| FAHD1  | FAH1D_HUMAN  | 0.233                       | no                          | 26(MPP), 27(kp55) | 10-14                                  |
| FAHD2a | FAH2A_HUMAN  | 0.790                       | yes                         | 83(MPP), 84(kp55) | 34,38,80-84                           |
| FAHD2b | FAH2B_HUMAN  | 0.884                       | yes                         | 83(MPP), 84(kp55) | 34,38,80-84                           |

FAHD2.

2.3. FAHD proteins are subject to differential mitochondrial import mechanisms

Proteins synthesized in the cytosol are imported into mitochondria via the general import pore (Lesnik et al., 2015; Walther and Rapaport, 2009), a multi-protein complex involving Tom5, Tom6, Tom7, Tom20, Tom22, Tom40, and Tom70. On the other hand, precursors of so-called signal-anchored proteins are imported to the mitochondria by a different mechanism (Ahting et al., 2005). Localization of FAHD1 in mitochondria despite the lack of a recognizable mitochondrial pre-sequence may suggest the presence of such a signal-anchor in FAHD1. The UniProt (Wasmuth and Lima, 2017) database lists curated (reviewed) entries of human proteins with signal-anchor motifs (keyword Signal-anchor KW-0735). BLASTp analysis of human FAHD1 and established signal anchor proteins displays significant sequence similarities with 8 entries, mapping to 4 proteins and their isoforms: Lactosylceramide

Table 2

| Enzyme | Name                     | UniProt-ID | UniProt-Spec | Theoretical pI | Instability index | Stability | Part of a complex | Structure |
|--------|--------------------------|------------|--------------|----------------|-------------------|-----------|-------------------|-----------|
| CS     | Citrate synthase         | Q73590     | CSY_HUMAN    | 8.45           | 22.40             | stable     | no                | homodimer |
| ACO    | Aconitate hydratase      | Q97998     | ACON_HUMAN   | 7.36           | 34.70             | stable     | no                | monomer   |
| IDH2   | Isocitrate dehydrogenase | P48735     | IDHP_HUMAN   | 8.88           | 29.77             | stable     | no                | homodimer |
| IDH3   | Isocitrate dehydrogenase | P50213     | IDH3A_HUMAN  | 6.46           | 41.24             | unstable   | yes (IDH3)       | complex   |
|        | (NADP) subunit alpha     | P43837     | IDH3B_HUMAN  | 8.64           | 36.88             | stable     | yes (IDH3)       | complex   |
|        | Isocitrate dehydrogenase | P51553     | IDH3G_HUMAN  | 8.75           | 45.59             | unstable   | yes (IDH3)       | complex   |
|        | (NADP) subunit beta      | P02318     | OGD1_HUMAN   | 6.39           | 45.17             | unstable   | yes (OGDC)       | complex   |
| OGDc   | Dihydrolipoyl-succinate  | P36957     | OGD2_HUMAN   | 9.10           | 50.53             | unstable   | yes (OGDC)       | complex   |
|        | dehydrogenase            | P39257     | DLDH_HUMAN   | 9.75           | 28.07             | stable     | yes (OGDC)       | complex   |
| SUC(A/G)| Succinate-CoA ligase     | P55937     | SUCA_HUMAN   | 9.01           | 41.30             | unstable   | yes (SUCA/SUCG)  | complex   |
|        | (ADP/GDP-forming) subunit alpha | Q96999 | SUCB2_HUMAN | 6.15           | 32.54             | stable     | yes (SUCG)       | complex   |
|        | Succinate-CoA ligase     | P89287     | SUCA1_HUMAN  | 7.05           | 41.13             | unstable   | yes (SUCA)       | complex   |
| SDC    | Succinate dehydrogenase  | P31040     | SDHA_HUMAN   | 7.06           | 37.04             | stable     | yes (SDH)        | complex   |
|        | (ubiquinone) flavin       | P29152     | SDHB_HUMAN   | 7.03           | 60.13             | unstable   | yes (SDH)        | complex   |
|        | substrate                | P99643     | CSO_HUMAN    | 9.74           | 47.79             | unstable   | yes (SDH)        | complex   |
|        | Succinate dehydrogenase  | P14521     | DHSD_HUMAN   | 8.92           | 33.20             | stable     | yes (SDH)        | complex   |
| FH     | Fumarase hydratase       | P07954     | FUMH_HUMAN   | 8.85           | 28.59             | stable     | no                | homodotramer |
| MDH2   | Malate dehydrogenase     | P40256     | MOHY_HUMAN   | 8.92           | 31.92             | stable     | no                | homodimer |
| FAHD1  | Fumaratelyase oxidoreductase domain containing protein 1 | Q6PS87 | FAH1D_HUMAN | 6.98 | 42.36 | unstable | likely | homodimer |
| FAHD2a | Fumaratelyase oxidoreductase domain containing protein 2a | Q9GXX7 | FAH2A_HUMAN | 6.48 | 41.26 | unstable | unknown | unknown |
| FAHD2b | Fumaratelyase oxidoreductase domain containing protein 2b | Q4P233 | FAH2B_HUMAN | 7.64 | 40.43 | unstable | unknown | unknown |
alpha-2,3-sialyltransferase (Q9UNP4, Q9UNP4−2, Q9UNP4−3), Beta-1,4-galactosyltransferase 7 (Q9UBV7), Adipocyte plasma membrane-associated protein (Q9HDC9, Q9HDC9−2), and Membrane metalloendopeptidase-like 1 (Q495T6, Q495T6−2). Alignment displays sequence identity in the amino acid ranges 1–24, 26–84, 27–131 and 185–207 of human FAHD1. For details on the dataset and computation see supplementary material.

This data may suggest a possible mechanism by which FAHD1 is synthetized in the cytosol and incorporated into mitochondria as a signal-anchored protein. The aforementioned predicted sites for cleavage of the FAHD1 sequence by MPP and ICP55 (see above) provide additional support for this theory. However, a possible cleavage by MPP at amino acids N26 and Y27 (see Table 1) would destroy the catalytic domain that is required for a functional protein (Weiss et al., 2018a) (see above), which appears unlikely. Hence, additional studies about processing of FAHD1 polypeptides during mitochondrial import seem warranted.

2.4. Potential interaction partners of FAHD proteins

Certain proteins have been listed in previous versions of the BioPlex (Huttlin et al., 2015) network, but have been removed in newer versions, probably reflecting a more stringent use of the COMPASS software (Huttlin et al., 2015) in more recent studies. Taking these changes into account, the probability of interaction partners may be ranked, preferring proteins that are listed in newer versions over proteins that were dropped in newer versions. Accordingly, the most probable binding partners of FAHD proteins are depicted as a bubble chart diagram in Fig. 2, each outer circle representing a lower ranking than the inner circles. The following proteins have been identified as potential FAHD1 interaction partners (see Fig. 2), some of which are also reported to interact with FAHD2:

### Table 3

Potential interaction partners of FAHD proteins, as listed in the BioPlex (Huttlin et al., 2015) network of different versions. Highlighted in gray are proteins that are listed in the newest versions 2 and 3 of the network. Other proteins were listed in early versions of the network but removed in the latest stable version 3. Localization and description of the proteins was gathered from the Human Protein Atlas (Uhlen et al., 2005; Fagerberg et al., 2014; Uhlen et al., 2010) and the UniProt (Wasmuth and Lima, 2017) database.

| Enzyme | Interaction | BioPlex | Localization (Human Protein Atlas, UniProt) | Description |
|--------|-------------|---------|---------------------------------------------|-------------|
| ARL2   |            | 3.0     | Nucleoplasm, Nuclear, Golgi apparatus, focal adhesion sites, Cytosol, Nucleoplasm | ADP ribosylation factor 1 (GTPase-activating protein 2, GTPase binding domain containing 1) |
| PTBD1  |            | 3.0     | Nucleoplasm, Nuclear, Mitochondria | Carnitine palmitoyltransferase 2 |
| CPT2   |            | 3.0, 2.0, 1.0 | Nucleoplasm, Nuclear, Mitochondria | Carnitine palmitoyltransferase 2 |
| DHPS4  |            | 3.0     | Vesicles, Nuclear membrane | Dehydrogenase/reductase/Fatty acid amidohydrolase 4 |
| DHO1   |            | 2.0, 1.0 | Mitochondria | Dehydrogenase E1 and transketolase domain containing 1 |
| FSD1   |            | 2.0, 1.0 | Nucleus | Fibrinogen type III and SPRY domain containing 1 |
| INHA   |            | 3.0     | Vesicles | Inhibin alpha subunit |
| CLUH   |            | 3.0, 2.0, 1.0 | Vesicles, Nuclear bodies | Clustered mitochondrial homolog |
| MTFD1  |            | 3.0     | Nucleoplasm, Mitochondria | Mitochondrial transcription termination factor 3 |
| NDP56  |            | 3.0, 2.0, 1.0 | Mitochondria | Nuclear ubiquitinyltransferase subunit 56 |
| ORI10H3 |            | 2.0, 1.0 | Cell membrane | Olfactory receptor family 10 subfamily H member 3 |
| PNP71  |            | 3.0, 2.0, 1.0 | Mitochondria | Polypeptide nucleolus/miniglobin 1 |
| UBR3   |            | 3.0, 2.0, 1.0 | Nucleoplasm, Nuclear | Ubiquitin protein ligase E3 component n-recogpin 3 (putative) |

![A) B) C)](image_url)

**Fig. 2.** Predicted interaction partners of FAHD proteins.

The most probable interaction partners of FAHD proteins according to data analysis by the BioPlex (Huttlin et al., 2015) network, are depicted as a bubble chart diagram. Certain proteins have been listed in previous versions of the BioPlex (Huttlin et al., 2015) network, but have been removed in newer versions. Taking these changes into account, the probability of interaction partners may be ranked, preferring proteins that are listed in newer versions over proteins that were dropped in newer versions. In each panel, outer circles represent a lower ranking compared with the inner circles.
Carnitine palmitoyltransferase 2 (CPT2) is part of the carnitine shuttle system that is required for the import of palmitic acid into the mitochondrial matrix. CPT2 is localized at the matrix side of the inner mitochondrial membrane and required for the import of fatty acids into mitochondria (UniProt (Wasmuth and Lima, 2017)). Clustered mitochondria homolog (CLUH) is an mRNA-binding protein which is thought to ascertain proper cytoplasmic distribution of mitochondria. CLUH specifically binds mRNAs of nuclear-encoded mitochondrial proteins in the cytoplasm and regulates the transport and/or translation of these transcripts close to mitochondria, playing a role in mitochondrial biogenesis (UniProt (Wasmuth and Lima, 2017)). NADH dependent ubiquinone oxidoreductase subunit S6 (NDUFS6) is an accessory subunit of the mitochondrial membrane respiratory chain NADH dehydrogenase (Complex I) (UniProt (Wasmuth and Lima, 2017)). Polyribonucleotide nucleotidytransferase 1 (PNPT1) as an RNA-binding protein is implicated in numerous RNA metabolic processes. It catalyzes the phosphorylation of single-stranded polyribonucleotides processively in the 3′→5′ direction (UniProt (Wasmuth and Lima, 2017)). Putative ubiquitin protein ligase E3 component n-recognin 3 (UBR3) is an E3 ubiquitin-protein ligase which is a component of the N-end rule pathway, leading to ubiquitination and subsequent degradation of its target proteins (UniProt (Wasmuth and Lima, 2017)). Bola family member 3 (BOLA3) acts as a mitochondrial iron-sulfur (Fe-S) cluster assembly factor that facilitates Fe-S cluster insertion into a subset of mitochondrial proteins (UniProt (Wasmuth and Lima, 2017)). Heat shock protein family D (Hsps60) member 1 (HSPD1) is a chaperonin implicated in mitochondrial protein import and macromolecular assembly (UniProt (Wasmuth and Lima, 2017)).

Based on this dataset, we hypothesize a possible relation of FAHD proteins with fatty acid beta-oxidation and RNA metabolic processes. A possible association of FAHD1 with Complex I would support our model of FAHD1 acting as regulatory enzyme in the context of mitochondrial dysfunction associated senescence (MiDAS) described by us (Stöckl et al., 2020) and others (Wiley et al., 2016). However, more experimental data is required in order to probe for such connections.

3. FAHD proteins may play an unanticipated role in calcium homeostasis

3.1. Calcium in mitochondria

Calcium plays a key role in many vital processes, such as bone homeostasis, signal processing in neurons (inclusive serotonin effects), cell death and survival. Deterioration of calcium homeostasis is associated with aging (Herraiz-Martínez et al., 2015; Veldurthy et al., 2016), and both directly (Herraiz-Martínez et al., 2015) and indirectly linked to cholesterol homeostasis (van der Wulp et al., 2013; Wang et al., 2017). Serotonin levels and calcium homeostasis are linked to bone loss and type 2 diabetes (Erjavec et al., 2016). Vitamin D is associated to bone health and is an essential cofactor for calcium binding in the bone, which becomes even more important with aging (Veldurthy et al., 2016; Oudshoorn et al., 2009). The major calcium reservoir in cells is the endoplasmic reticulum. Mitochondrial calcium content is tightly regulated in most if not all eukaryotic cells.

Calcium uptake into and release from mitochondria is important in regulating a variety of cellular physiological functions (Takeuchi et al., 2015). Calcium handling by mitochondria is involved in energy production, in buffering and shaping cytosolic calcium, and in determining cell fate by triggering or preventing apoptosis (Contreras et al., 2010). Mitochondrial Ca²⁺ uptake is mainly mediated by a mitochondrial Ca²⁺ uniporter (MCU) driven by membrane potential (Perocchi et al., 2010), as well as by 2 H⁺ – Ca²⁺ exchange (Finkel et al., 2015). Mitochondrial Ca²⁺ is mainly released by a 3 Na⁺ – Ca²⁺ exchanger (NCLX) (Carafoli, 1974), but also by an active 2 H⁺ – Ca²⁺ exchange that has a dominant effect on release of Ca²⁺ from mitochondria in tissues in which mitochondrial NCLX activity is low (Takeuchi et al., 2015; Gunter and Pfeiffer, 1990). Calcium-binding mitochondrial carrier proteins (e.g. SL2C5A12, SL2C5A23, and SL2C5A24) are reported to facilitate the calcium-dependent exchange of cytoplasmic metabolites across the mitochondrial inner membrane. However, there is scarce data on mitochondrial calcium binding proteins, except for mitochondrial ATP synthase F1-beta-subunit (Hubbard and McHugh, 1996), and for the predominantly mitochondrial protein HAX1 (Balcerak et al., 2017).

Of note, uptake of Ca²⁺ requires co-transport of an inner mitochondrial membrane permeable anion such as acetate or phosphate (Starkov, 2010), and the accumulated Ca²⁺ forms a detectable precipitate (Chinopoulos and Adam-Vizi, 2010) in the matrix of mitochondria in an apparently spontaneous process (Starkov, 2010). The granules contain significant amounts of carbon and nitrogen, indicating the presence of yet unidentified protein(s), that are suggested to serve as nucleation centers, facilitating formation of the Ca²⁺ precipitate (Starkov, 2010). This precipitate is suggested to be in pH equilibrium with the inner mitochondrial matrix, and eventually slowly released back into the cytosol (Starkov, 2010; Chinopoulos and Adam-Vizi, 2010).

During cellular activation Ca²⁺ levels in the mitochondrial matrix may reach up to μmol/L levels (Ivanivnik and Macleod, 2013). High levels of intracellular Ca²⁺ activate mitochondrial NADP dependent isocitrate dehydrogenase (IDH2) and the 2-oxoglutarate dehydrogenase complex (OGDC), as well as pyruvate dehydrogenase phosphatase (Pelley, 2007), which in turn activates the pyruvate dehydrogenase complex (PDC) (Pelley, 2007) to create acetyl-CoA to be used by citrate synthase (CS). These changes increase the reaction rate of many of the steps in the TCA cycle, and therefore increase flux throughout the pathway.

3.2. Endoplasmic reticulum and mitochondria direct the role of calcium in cellular senescence

Published links between calcium signaling and cellular senescence are summarized in a recent review by Martin and Bernard (Martin and Bernard, 2018), summarizing how calcium critically controls many molecular processes and cellular functions (Martin and Bernard, 2018; Humeau et al., 2018; Parys and Bultynck, 2018). In particular, knockdown of the mitochondrial calcium uniporter was reported to foster escape from senescence (Martin and Bernard, 2018). Elevation of intracellular calcium levels has been observed in response to different types of senescence-inducing stresses (telomere shortening, oncogene activation, roteneone or oxidative stress) in several cell types (Martin and Bernard, 2018). High concentrations of intracellular calcium are sustained during senescence (Martin and Bernard, 2018; Farfariello et al., 2015). This increase in calcium concentration has been attributed to calcium influx through plasma membrane calcium channels or to calcium release from the endoplasmic reticulum, depending on the context (Martin and Bernard, 2018; Giorgio et al., 2018). The endoplasmic reticulum was reported by many studies to play a key role in the regulation of calcium levels, cross-talking with mitochondria (Wiel et al., 2014; Gutiérrez and Simmen, 2018; Carreras-Sureda et al., 2018; Pitts and Hoffmann, 2018), i.e., endoplasmic reticulum and mitochondria can be spatially and functionally coupled through mitochondria-associated endoplasmic reticulum membranes which favor the transfer of calcium from the endoplasmic reticulum to mitochondria (Patergnani et al., 2011). Endoplasmic reticulum chaperones tweak the mitochondrial calcium rheostat to control metabolism and cell death (Gutiérrez and Simmen, 2018). The main endoplasmic reticulum calcium release channels, inositol 1,4,5-trisphosphate receptors (ITPRs), were originally proposed as suppressors of autophagy (Bootman et al., 2018). In particular, calcium release through ITPR2 channels was reported to lead to mitochondrial calcium accumulation and senescence (Wiel et al., 2014). Calcium released from the endoplasmic reticulum in response to senescence-inducing stresses mainly exerts its effects through reactive oxygen species (Carreras-Sureda et al., 2018).
human mammary epithelial cells and primary human fibroblasts, on-cogene activation and telomere shortening may also trigger calcium release from endoplasmic reticulum stores through the activation of the PLC/IP3/IP3R pathway (Martin and Bernard, 2018).

3.3. FAHD proteins are highly expressed in Ca²⁺ rich and Ca²⁺ regulating tissues

Calcium is the most abundant mineral in the human body, with Ca²⁺ concentration in plasma ranging between 2.1 and 2.6 mmol/L. (Minisola et al., 2015), while higher calcium levels are defined as hypercalcemia (Minisola et al., 2015). While about 99 % of the body’s calcium is stored in the bone, about 1 % can be found in the blood serum, referred to as free calcium. The level of free calcium must remain within a very narrow concentration range to support vital physiological functions (Minisola et al., 2015). Cells absorb Ca²⁺ across the brush border of the enterocyte cell membrane by a mechanism that requires energy and vitamin D as an essential cofactor (Veldurthy et al., 2016), and vitamin D deficiency has been related to calcium homeostasis and aging (Oudshoorn et al., 2009; Kuro-o et al., 1997; Urakawa et al., 2006).

The absorption of calcium from food is performed by acid secretion from the stomach that converts calcium from various sources to Ca²⁺ salt which is then absorbed primarily in the duodenum. This mechanism is mainly influenced by conditions within the lumen of the small intestine. The thyroid gland releases calcitonin when levels of serum calcium are too high, which slows down the process of calcium release in the bone. The parathyroid gland produces parathyroid hormone when levels of serum calcium become too low, which in turn stimulates the release of calcium from the bones into the bloodstream. Hypocalcemia is mainly caused by malfunctions in the parathyroid gland. On the other hand, about 99 % of free calcium is reabsorbed by the kidney. Also, Ca²⁺ interferes with the absorption of iron (Fe²⁺) in the liver, so Ca²⁺ may accumulate in the liver (Ruchay, 2016). Of note, calcium homeostasis is highly important for the heart, and aging of the heart is associated with a decrease of calcium levels in the heart tissue (Herraz-Martínez et al., 2015).

Table 4 summarizes the data on FAHD expression in human tissues, as listed in the Human Protein Atlas (Uhlen et al., 2005; Fagerberg et al., 2014; Uhlen et al., 2010). It is striking that FAHDI is highly expressed in tissues that are associated with calcium metabolism and the regulation of calcium homeostasis. FAHD proteins are generally high in the parathyroid gland, stomach, and kidney. FAHDI levels are also high in the adrenal gland, small intestine and duodenum. Levels of FAHDIa and FAHDIb are high in the liver, thyroid gland and salivary gland, where levels of FAHDI are high as well. There are several studies connecting these organs to calcium homeostasis and regulation (Brown and Vaidya, 2014; Ambudkar, 2016). The nasopharynx (displaying high levels of FAHDI) is usually not associated with calcium regulation, however, there is a recent documentation of a rare case of nasopharynx carcinoma because of hypercalcemia (Chaudhary and Sah, 2020). In contrast, detected FAHDI protein levels are generally low in tissues that are not associated to calcium homeostasis Table 5.

3.4. Indirect evidence for calcium binding of FAHDI proteins

IonCom (Zheng et al., 2019; Hu et al., 2016) analysis for human FAHDI was performed to obtain information on predicted ion binding sites (see Table 4). This analysis was done by aligning deep neural-network based contact maps based on the 3D PDB structural data of human FAHDI (6FOH). Potential binding sites have been predicted for Zn²⁺, Ca²⁺, Mg²⁺, Na⁺, K⁺, PO₄³⁻. No binding sites have been predicted for Cu²⁺, Fe²⁺/³⁺, Mn²⁺, CO₂⁻, NO₂⁻, SO₄²⁻. The experimentally verified binding motif for Mg²⁺ in the catalytic domain (Weiss et al., 2018a) was successfully predicted by the algorithm. This is considered as a trustful quality control. Other binding sites are reported for Zn²⁺ and for Ca²⁺, as well as for PO₄³⁻.

Calcium-binding proteins participate in calcium cell signaling pathways by binding of calcium ions, thereby regulating the levels of free Ca²⁺ in the cytosol of the cell. Free calcium in the mitochondrial matrix can vary widely (100–800 nmol/L) (Finkel et al., 2015), depending on the extra-mitochondrial calcium level. Many different calcium-binding proteins exist, that are known to be heterogeneous, among them a group of proteins known as the EF-hand superfamily (Ishida and Vogel, 2013). The EF hand is a helix-loop-helix structural domain or motif found in a large family of calcium-binding proteins (Nakayama and Krebsinger, 1994). None of the reported EF-motifs (Ishida and Vogel, 2013) was fully identified in the sequence of FAHDI, but BLASTp analysis detected the amino acid sequence 142-DPHALK-147 in FAHDI that would partly match one of the reported EF-hand motifs (Ishida and Vogel, 2013) (SGREGDKHKLKKSE). BLASTp analysis of human FAHDI was performed against known EF-hand domain-containing proteins (see Fig. 3D; see supplementary material for details on the dataset and computation). Among the screened proteins, human Zinc finger ZZ-type and EF-hand domain-containing protein 1 (ZZEF1, UniProt (Wasmuth and Lima, 2017)-ID O43149) displays significant sequence identity with human FAHDI isoform 1 (UniProt (Wasmuth and Lima, 2017)-ID Q6P587). The N-terminal motif is succeeded by a flexible loop region that is typical for FAH superfamily enzymes and participates in the catalytic mechanism (Weiss et al., 2018a) (see Fig. 3A). AllostERIC regulation may be anticipated.

Similar data analysis has been performed for known zinc binding proteins, focusing on the LIM domain (PDB: 1X62), the Zinc Finger 3 motif (PDB: 1VA3), the coiled-coil Zn hook (PDB: 1LBG) and LCK fragments (PDB: IQ68). Among the four screened motifs, the Zinc Finger 3 motif and the coiled-coil Zn hook showed significant sequence identity with FAHDI in BLASTp analysis (see Fig. 3B and C). The two representative structures are Zinc-hook domain-containing protein RAD50 (Hopfer et al., 2002) (see Fig. 3B) and Transcription factor Sp1 (Oka et al., 2004) (see Fig. 3C). The Rad50 zinc-hook is a structure joining Mre11 complexes that are central to chromosomal maintenance, and functions in homologous recombination, telomere maintenance and sister chromatid association (Hopfer et al., 2002). SP1 is a transcription factor that can activate or repress transcription in response to physiological and pathological stimuli (Oka et al., 2004). It positively regulates the transcription of the core clock component ARNTL/BMAL1 (Oka et al., 2004) and plays an essential role in the regulation of FGF5 gene expression (Oka et al., 2004). Although a local sequence similarity does not imply similar protein function in general, these data complement the data of possible FAHDI interaction partners (see above) and contribute to the hypothesis of a potential relation of FAHDI proteins with RNA metabolism.

The data of IonCom (Zheng et al., 2019; Hu et al., 2016) analysis suggesting Zn²⁺ and Ca²⁺ binding of FAHDI seems to match with the BLASTp alignment of FAHDI and zinc or calcium binding proteins, although no complete binding motif (ZZ-type, EF-hand, LIM domain, Zinc-hook, ...) could be identified in the FAHDI sequence.

FAHDI shows highest ApH-activity with Mg²⁺ and Mn²⁺ as cofactors, whereas Ca²⁺- and Zn²⁺-bound enzyme displays strongly reduced catalytic activity (Pircher et al., 2011). ODx activity of FAHDI prefers the same metals as ApH. Such findings implicate that distinct divalent metal ions, such as Ca²⁺ and Zn²⁺, may be prone to inhibit the catalytic activity of FAH superfamily proteins. High levels of calcium would reduce FAHDI’s enzymatic activity by contest of cofactor Mg²⁺ and competing Ca²⁺ ions. We further tested if there is a potential contest of the cofactors that may be associated to Ca²⁺ regulation. When catalytic activity of recombinant human FAHDI (Weiss et al., 2019) was tested in in vitro assays against cofactor concentrations, we observed a significant decrease of ODx activity with increasing Ca²⁺ concentrations (A. Weiss et al., unpublished). We propose a model where FAHDI is regulated by a contest of cofactor Mg²⁺ and competing Ca²⁺ ions, and its catalytic ODx activity is decreased by increased Ca²⁺.
levels (see Fig. 4). In consequence, decreased Ca^{2+} levels would decrease oxaloacetate levels by activation of FAHD1 (in the presence of Mg^{2+}).

3.5. FAHD1 effects on serotonin signaling – a link to Ca^{2+} signaling?

We could show that egg laying behavior is altered in fahd-1 depleted Caenorhabditis elegans (Taferner et al., 2015; Baraldo et al., 2019). Whereas wild-type animals do not lay eggs when put in a hypertonic salt solution and commence egg-laying only after serotonin-treatment, fahd-1 (-/-) worms did not cease egg-laying under these unfavorable conditions (Taferner et al., 2015; Baraldo et al., 2019) nor did they increase their egg-laying rate upon contact with exogenously applied serotonin (up to 10 mM) (Baraldo et al., 2019). It is known that egg-laying is an active process which is regulated by neuronal signals mediated by serotonin (and several other neurotransmitters) (Horvitz et al., 1982; Trent et al., 1983) and requires intact vulval musculature (Desai et al., 1988; Schinkmann and Li, 1992; Weinshenker et al., 1995). Altered egg-laying behavior in fahd-1 depleted worms was associated with a significant upregulation of the gene basl-1, that is predicted to have carboxylase activity and pyridoxal phosphate binding activity (WormBase, WBGene00015467#0−9f-10). BLASTp analysis of UniProt (Wasmuth and Lima, 2017) entry O45138 BAS-Like OS = Caenorhabditis elegans provided about 35 % sequence identity with UniProt (Wasmuth and Lima, 2017) entry P20711, the human protein aromatic-L-amino-acid decarboxylase (DDC, also PXLP-DDC or AADC). This protein catalyzes the decarboxylation of L-dopa to dopamine, and of 5-hydroxy-L-tryptophan to serotonin (EC:4.1.1.28). The catalytic activity of the human protein matches the reported activity of the nematode protein. Upregulation of basl-1 as a reaction to fahd-1 knockout would, therefore, indicate the increased production of serotonin from precursor metabolite 5-hydroxy-L-tryptophan. From these data we concluded that FAHD-1 in Caenorhabditis elegans modulates serotonin signaling (Baraldo et al., 2019).
Table 5
Ion ligand binding prediction using the IonCom (Zheng et al., 2019; Hu et al., 2016) analysis, by aligning deep neural-network based contact maps based on the PDB data of human FAHD1 (6FOH). Potential binding sites have been predicted for Zn^{2+}, Ca^{2+}, Mg^{2+}, Na^{+}, K^{+}, PO_{4}^{3−}. No binding sites have been predicted for Cu^{2+}, Fe^{2+/3+}, Mn^{2+}, CO_{3}^{2−}, NO_{2}^{−}, SO_{4}^{2−}.

| Zn^{2+} | Ca^{2+} | Mg^{2+} | Na^{+} | K^{+} | PO_{4}^{3−} |
|---------|---------|---------|--------|------|------------|
| G17     |         |         |        |      |            |
| K18     |         |         |        |      |            |
| C22     |         |         |        |      |            |
| V23     |         |         |        |      |            |
| G24     |         |         |        |      |            |
| R25     |         |         |        |      |            |
| S36     |         |         |        |      |            |
| F45     |         |         |        |      |            |
| S49     |         |         |        |      |            |
| E55     |         |         |        |      |            |
| H69     |         |         |        |      |            |
| E71     |         |         |        |      |            |
| E73     |         |         |        |      |            |
| C82     |         |         |        |      |            |
| V85     |         |         |        |      |            |
| Y97     |         |         |        |      |            |
| L101    |         |         |        |      |            |
| D102    |         |         |        |      |            |
| M103    |         |         |        |      |            |
| R106    |         |         |        |      |            |
| D107    |         |         |        |      |            |
| Q109    |         |         |        |      |            |
| C112    |         |         |        |      |            |
| W119    |         |         |        |      |            |
| K223    |         |         |        |      |            |
| F125    |         |         |        |      |            |
| T126    |         |         |        |      |            |
| C129    |         |         |        |      |            |
| S132    |         |         |        |      |            |
| L150    |         |         |        |      |            |
| N153    |         |         |        |      |            |
| E155    |         |         |        |      |            |
| E159    |         |         |        |      |            |
| D186    |         |         |        |      |            |
| G191    |         |         |        |      |            |
| T192    |         |         |        |      |            |
| D208    |         |         |        |      |            |
| E204    |         |         |        |      |            |
| I205    |         |         |        |      |            |
| A207    |         |         |        |      |            |
| S214    |         |         |        |      |            |
| E223    |         |         |        |      |            |

Calcium homeostasis in nematodes is involved in movement, fertility, egg-laying and growth of Caenorhabditis elegans (Bandyopadhyay et al., 2002), and it may in fact be a deteriorated calcium homeostasis that impacts the nematode’s egg-laying behavior, as was implied by others (Bandyopadhyay et al., 2002). Recent work on serotonin signaling and calcium homeostasis in different species showed diverse outcomes. Effects have been reported in studies of milk production and milk quality in dairy cows (Hernández-Castellano et al., 2017; Weaver et al., 2016), where a certain ambiguity between cause and relation of serotonin and calcium homeostasis is described. Serotonin is mainly responsible for increasing calcium pumps in the mammary gland (Hernandez et al., 2012) and secretion into milk (Laporta et al., 2013). Infusion of serotonin acutely decreased free calcium concentrations (Erjavec et al., 2016) because of bone loss and the development of type 2 diabetes (Erjavec et al., 2016). It is discussed that a possible answer to this problem might be the explanation of a time-dependent change in metabolism, where an acute change in serotonin (such as feeding serotonin to cows for days) differs from a long-term change in metabolism (such as rats with long term inhibitory treatment). In Caenorhabditis elegans, calcium imaging studies could show that serotonin acts directly on the vulval muscles to increase the frequency of spontaneous calcium transients, thus increasing egg-laying (Shyn et al., 2003).

Current data reveals a link of FAHD-1 depletion in Caenorhabditis elegans to a significant change in the nematode’s serotonin signaling pathway. However, more elaborate experiments on serotonin signaling and calcium homeostasis in Caenorhabditis elegans are warranted to reveal a possible link to FAHD-1 depletion.

4. Discussion and outlook

4.1. Multiple physiological functions of FAHD proteins in eukaryotes

Predicted protein interaction partners of FAHD1 reflect its reported localization (Pircher et al., 2011; Wasmuth and Lima, 2017; Uhlen et al., 2010), and suggest a putative role of FAHD proteins in the pathways of fatty acid oxidation, oxidative phosphorylation, mitochondrial RNA metabolism and the ubiquitin/proteasome system. As available data from high-throughput proteomics analysis (Huttlin et al., 2015) suggest, the most probable interaction partners of FAHD1 are carnitine palmitoyltransferase 2 (CPT2), clustered mitochondria homolog (CLUH), NADH dependent ubiquinone oxidoreductase subunit S6 (NDUF6), polyribonucleotide nucleotidyltransferase 1 (PNPT1), and putative ubiquitin protein ligase 3 (UBR3). NDUFS6 is an accessory subunit of the mitochondrial membrane respiratory chain complex I. A putative interaction with FAHD1 may complement our recently hypothesized model of senescence (Etemad et al., 2019) due to the inactivation of genes required for mitochondrial function (such as SIRT3 (Hallows et al., 2011) and FAHD1 (Etemad et al., 2019)), thus explaining how in some cellular models the inactivation of either ETC complex I (by metformin) or ETC complex II (by FAHD1 knockdown) has the potential to increase p21 gene expression in the absence of AMPK (Etemad et al., 2019). In agreement with results obtained from a high-throughput proteomics study (Dittenhafer-Reed et al., 2015), we recently provided circumstantial evidence for a SIRT3 deacetylation site (Dittenhafer-Reed et al., 2015) in mouse FAHD1 (Weiss et al., 2020), which further supports this model.

4.2. A new role for FAHD1 in calcium homeostasis?

FAHD proteins are members of the FAH superfamily of metabolic enzymes, the physiological role of which is only partially explored. In the case of FAHD1, existing evidence suggests that it is a mitochondrial protein which can catalyze hydrolysis of acylpyruvates and the decarboxylation of oxaloacetate. However, several features of FAHD1 activity remain largely unexplored, in particular due to the fact that experiments with FAHD1/2 depleted cells and animals still lack considerable mechanistic detail. The main purpose of this review is to stimulate discussions in this understudied field of research, and to critically review the research agenda how to unmask molecular mechanisms of action for these proteins.

We have proposed a model of how FAHD1 catalytic activity as oxaloacetate decarboxylase in mitochondria may describe FAHD1 as a regulator of TCA cycle flux and as a possible regulator of mitochondrial function and senescence (Etemad et al., 2019). We now propose a complementary model of how the actual presence of FAHD1 protein (or lack thereof), independent of its catalytic function, may influence intracellular calcium levels. It is well reported that FAHD1 expression in human organs correlates with the regulation of calcium metabolism in...
the human body, and experimental results described in this work are in line with the hypothesis that FAHD1 may be a calcium binding protein. Calcium binding proteins are present in various cellular compartments and serve to mediate effects of increased calcium concentration on biological responses. On the other hand, it is conceivable that calcium binding proteins serve as buffering systems to fine-tune the concentration of intracellular calcium. Our unpublished observation that increasing levels of calcium inactivate FAHD1 catalytic activity in vitro is in line with the model of how calcium levels modulate the TCA cycle flux (Etemad et al., 2019) (Fig. 4). The model predicts coordinated but inverse regulation of FAHD1 and the canonical TCA cycle enzymes IDH and OGD, respectively, suggesting a regulatory mechanism by which increasing calcium levels in the mitochondrial matrix booster flux through the TCA cycle.

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CRediT authorship contribution statement

Alexander K.H. Weiss: Conceptualization, Project administration, Investigation, Data curation, Validation, Methodology, Resources. Eva Albertini: Investigation, Data curation, Validation. Max Holzknecht: Methodology. Elia Cappuccio: Methodology. Ilaria Dorigatti: Methodology. Elisabeth Damisch: Methodology. Hubert Gstach: Investigation, Data curation, Validation, Methodology, Resources. Pidder Jansen-Dürr: Conceptualization, Project administration, Investigation, Data curation, Validation, Resources.

Declaration of Competing Interest

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

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Appendix A. Supplementary data

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