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### Reactive oxygen species target specific tryptophan site in the 1 mitochondrial ATP synthase

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#### ABSTRACT

The release of reactive oxygen species (ROS) as side products of aerobic metabolism in the mitochondria is an 25 unavoidable consequence. As the capacity of organisms to deal with this exposure declines with age, accumu- 26 lation of molecular damage caused by ROS has been defined as one of the central events during the ageing 27 process in biological systems as well as in numerous diseases such as Alzheimer's and Parkinson's Dementia. 28 In the filamentous fungus Podospora anserina, an ageing model with a clear defined mitochondrial etiology of 29 ageing, in addition to the mitochondrial aconitase the ATP synthase alpha subunit was defined recently as a 30 hot spot for oxidative modifications induced by ROS. In this report we show, that this reactivity is not ran- 31 domly distributed over the ATP Synthase, but is channeled to a single tryptophan residue 503. This residue 32 serves as an intra-molecular quencher for oxidative species and might also be involved in the metabolic per- 33 ception of oxidative stress or regulation of enzyme activity. A putative metal binding site in the proximity of 34 this tryptophan residue appears to be crucial for the molecular mechanism for the selective targeting of ox- 35 idative damage. 36

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### 1. Introduction

Molecular damage caused by the exposure to ROS is a major cause of numerous diseases such as Alzheimer's and Parkinson's Dementia as well as a central factor of the ageing process by the 'free radical theory of ageing' (FRTA) [1] and its refined version, the 'mitochondrial free radical theory of ageing' (MFRTA) [2]. The latter theory identifies mitochondria as the major site of ROS generation and the predominant target of damage of all kinds of biomolecules. As a result, mitochondria become dysfunctional and biological systems degenerate and die. On the other hand, however, ROS are also employed in many life-sustaining mechanisms such as in signaling pathways and defense against pathogens.

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In ageing research the conditions and effects of ROS generation, 54 signaling, scavenging and damaging, are intensively studied in vari- 55 ous systems including different model organisms. Podospora anserina 56 is one of these organisms. This filamentous fungus is characterized by 57 a short lifespan of a few weeks and is tractable to experimentation. It 58 contains a small genome [3], of which the sequence has been 59 completely determined [4]. Ageing of P. anserina has a clear mito- 60 chondrial etiology. A number of specific factors and pathways which 61 are basically involved in keeping a functional population of mitochon- 62 dria have been demonstrated to affect ageing and lifespan [3, 5-10]. 63 Although ROS generation and scavenging do clearly contribute to 64 the ageing process in P. anserina [11, 12], recently counter-intuitive 65 data were obtained when the effect of the modulation of mitochon- 66 drial superoxide dismutase (PaSOD3) was studied [13]. In contrast 67 to what is expected from the MFRTA, strains deleted for PaSod3 68 were not short-lived and strains overexpressing PaSod3 were not 69 long- but short-lived. Intriguingly, the latter strains were character- 70 ized by a reduced abundance of a mitochondrial peroxiredoxin in- 71 volved in scavenging of hydrogen peroxide, and in PaCLP protease, 72 as a part of the mitochondrial protein quality control system. Taking 73 these observations into account allowed the generation of a mathe-74 matical model that, at least partly, explains the unexpected results 75 obtained from the study (manuscript in preparation). In particular, 76 in addition to ROS metabolism mitochondrial protein quality control 77

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Abbreviations: Desferal, Deferoxamine N'-{5-[acetyl(hydroxy)amino]pentyl}-N-[5-({4-[(5-aminopentyl)(hydroxy)amino]-4-oxobutanoyl}amino)pentyl]-N-hydroxysuccinamide; FRTA, Free radical theory of ageing; Kyn, kynurenine; MFRTA, mitochondrial free radical theory of ageing; MS, mass spectrometry; NFK, N-formyl-kynurenine; ROS, reactive oxygen species; SRM, single reaction monitoring; TRP-OH, hydroxytryptophan.

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systems need to be included in networks of pathways involved in the 78 79 control of ageing. Such systems may be triggered by and acting on specific posttranslational modifications. One of such modifications 80 81 has also been identified in P. anserina mitochondria as the irreversible oxidation of tryptophan residues and the formation of N-formyl-82 kynurenine (NFK) [14, 15]. NFK is frequently observed in proteins 83 by mass spectrometry along with the other oxidation intermediates 84 85 displayed in Fig. 1, particularly in tissues with high metabolic rates 86 and in proteins with long half lives [16]. Prominent is its presence 87 in human and bovine heart mitochondrial proteins [15, 17], rat skeletal muscle proteins [18], bovine  $\alpha$ -crystalline [19], as well as CP43 88 [20] and LHCII [21] in photosynthetic membranes. 89

Here we show that oxidative modifications are not—as intuitively
accepted—randomly distributed to all tryptophan residues over the
entire sequence of the alpha subunit of the mitochondrial ATPase,
but selectively targeted to one specific tryptophan.

#### 94 2. Materials and methods

#### 95 2.1. P. anserina cultivation and isolation of crude mitochondria fraction

P. anserina is a filamentous ascomycete which is normally growing 96 97 on solid medium. To obtain sufficient material for mitochondria isolation, after germination of monokaryotic ascospores on solid cornmeal 98 medium with 60 mM ammonium acetate, small pieces of mycelium 99 are first grown 2-3 days on solid PASM medium [22] under perma-100 nent light. To prevent transfer of solid medium into the subsequently 101 102 used liquid medium, the solid medium was overlaid with a cellophane sheet. The mycelium from this solid medium was then trans-103 ferred to liquid CM medium [23] and incubated for 2 days shaking 104 under permanent light. Different age stages were obtained by pre-105106growth to the respective age on solid medium and subsequent culture 107 in liquid medium. For instance, to isolate mitochondria from 6 days old cultures, ascospores were germinated for two days on solid corn-108 meal medium with ammonium acetate. Subsequently, pieces of my-109 celium from the growth front were transferred to solid medium 110 with cellophane. After two days of growth, the mycelium was 111 scratched from the plates and used to inoculate liquid medium. 112 After additional 2 days of growth the mycelium was harvested, giving 113 a total age of an individual of 6 days. Senescent cultures were isolated 114 by growing the fungus on solid medium until it reaches senescence 115116 (= stop of growth, alteration of pigmentation). The solid medium with cellophane was now inoculated with mycelium 2-3 cm 117 (4-5 days) behind the growth front. To prevent rejuvenation, cultiva-118 tion of senescent mycelium in liquid medium was performed without 119 shaking in Fernbach flasks (permanent light, 5–7 days). The isolation 120 121 of mitochondria from these different mycelial samples was modified according to Gredilla [12]: After high-speed centrifugation 122(15,000 g, 20 min) the mitochondria were directly resuspended in 123 isolation buffer without BSA, additional wash steps were omitted. In 124 total crude mitochondrial extracts of three independent wild-type 125



**Fig. 1.** Chemical structure of hydoxytryptophan (Trp-OH), N-formlykynurenine (NFK), and kynurenine (Kyn)-products and intermediates of the tryptophan oxidation [17]. Alternative structures with hydroxyl substitution at the benzene ring are plausible for Trp-OH and cannot be distinguished based on the mass shift. The mass shifts observed by MS relative to Trp are noted beneath the formulas.

isolates of the age stages 6 days and senescent (14–15 days) were  $^{126}$  prepared. In control experiments 100  $\mu M$  Desferal was added to all  $^{127}$  buffers used during isolation.  $^{128}$ 

### 2.2. 2-D Blue-native/tricine-SDS-PAGE and colorless-native PAGE 129

Isolated crude mitochondrial fractions were resuspended in a me- 130 dium containing 50 mM imidazol, pH 7.0, 50 mM NaCl, 5 mM  $\epsilon$ - 131 aminocaproic acid, and 10% glycerol. The membranes were solubi- 132 lized on ice for 30 min with the non-ionic detergent digitonin at a 133 final concentration of 1% (4 g digitonin/g protein). Insoluble material 134 was removed by centrifugation (21,000 g, 4 °C, 10 min). The superna- 135 tant was loaded directly onto blue-native gradient gels. Blue-native 136 PAGE was performed using the Hoefer SE 600 system 137  $(18 \times 16 \times 0.15 \text{ cm}^3, 10 \text{ lanes})$  as described previously [24, 25]. Stack- 138 ing gels with a total acrylamide concentration of 3.5% and separating 139 gels with linear acrylamide gradients 5–13% were prepared. 250 µg 140 solubilized protein (as determined before solubilization) was applied 141 per lane. Bovine heart mitochondria prepared from tissue and stored 142 at -80 °C (4 g digitonin/g protein, 70 µg of protein before solubiliza- 143 tion) served as molecular mass standard. 144

After electrophoresis, gels were stained with Coomassie R-250 or 145 lanes of the BN-gel were cut out and incubated in a solution of 1% 146 (w/v) SDS and 1% (w/v)  $\beta$ -mercaptoethanol at 20 °C for 30 min. Subsequently, lanes were analyzed by Tricine-SDS-PAGE in second dimension on a gel with two stacking gels, one native and one 149 denaturing, with a total acrylamide concentration of 10% and a separating gel with 16.5% [26].

After electrophoresis, gels were silver stained [27, 28]. Protein 153 bands of interest were excised inside a laminar flow hood. Silver 154 stained spots were destained [29], and in-gel digestion was performed with trypsin [30]. After extraction of the peptides with 50% 156 (v/v) acetonitrile / 0.5% (v/v) formic acid the solvent was removed 157 completely by lyophilization. Previous to LC-ESI-MS/MS analysis 158 peptides were resuspended in 2% acetonitrile with 0.1% formic acid. 159

### 2.4. LC–MS analysis

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LC–MS analysis was performed as described previously [31] with 161 slight modifications. For reverse phase chromatography, a gradient 162 of solvent A (95% water, 5% acetonitrile, 0.1% formic acid) and solvent 163 B (10% water, 85% acetonitrile, 5% isopropanol, 0.1% formic acid) was 164 used. For MS analysis, a Thermo LTQ Orbitrap mass spectrometer was 165 operated in a duty cycle consisting of one 400–2000 *m/z* FT-MS and 166 four MS/MS LTQ scans. For detection of low quantities of tryptophan 167 oxidation for all three tryptophan sites in the ATP synthase alpha subunit an inclusion list was defined for the doubly and triply charged 169 ions of the peptides ILQWEADFLSHLK, YSIVVAATASEAAPLQYLAPFT-GASIGEWFR, and WNSGNDETK and their respective variants with oxidized tryptophan residues (Table 1).

#### 2.5. Data analysis

Analysis of the LC–MS/MS data was performed using the Sequest 174 algorithm [32] implemented in the Bioworks 3.3.1 software (Thermo 175 scientific) for peptide identification versus a database [4] consisting of 176 all *P. anserina* proteins with a tolerance of 10 ppm for the precursor 177 mass accuracy and 1 u for the fragment mass accuracy. For detection 178 of oxidized peptides for tryptophan residues modifications of 179 15.9949, 31.9898, and 3.9949 u were defined as parameters during 180 the search for hydroxy-tryptophan, *N*-formyl kynurenin and kynure-181 nin, respectively. False discovery rates were estimated by the number 182

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#### t1.1 Table 1

Peptide masses defined in the inclusion list for LC-MS analysis.

t1.2 t1.3	Peptide variant	Modification	[M+2H] <sup>2+</sup>	$[M + 2H]^{3+}$
t1.4	ILQWEADFLSHLK	Unmodified	800.4301	533.9558
t1.5	ILQW(OH)EADFLSHLK	Hydroxy-Trp	808.4272	539.2875
t1.6	ILQW(NFK)EADFLSHLK	NFK	816.4250	544.6191
t1.7	ILQW(Kyn)EADFLSHLK	Kyn	802.4276	535.2875
t1.8				
t1.9	YSIVVAATASEAAPLQYLAPFTGASIGEWFR	Unmodified	1643.8427	1096.2309
t1.10	YSIVVAATASEAAPLQYLAPFTGASIGEW(OH)FR	Hydroxy-Trp	1651.8402	1101.5626
t1.11	YSIVVAATASEAAPLQYLAPFTGASIGEW(NFK)FR	NFK	1659.8377	1106.8942
t1.12	YSIVVAATASEAAPLQYLAPFTGASIGEW(Kyn)FR	Kyn	1645.8402	1097.5626
t1.13				
t1.14	WNSGNDETK	Unmodified	533.7254	356.1527
t1.15	W(OH)NSGNDETK	Hydroxy-Trp	541.7229	361.4844
t1.16	W(NFK)NSGNDETK	NFK	527.7254	352.1527
t1.17	W(Kyn)NSGNDETK	Kyn	533.7254	356.1527

of spectral matches to a decoy database [33]. Acceptance criteria andfilters were set to achieve a false positive rate of 5%.

#### 185 2.6. Absolute peptide quantification

186 For quantitative analysis of the oxidative modification, single reaction monitoring (SRM) was applied on a triple quadrupole mass spec-187 trometer [34]. Custom peptides were synthesized for Trp and Kyn 188 variants of the peptide I<sup>\*</sup>LQWEADFLSHLK by Thermo Scientific using 189 isoleucine residue with stable isotopes to induce a mass shift of 190 191 7 Da to the peptides. For analysis tryptic digests were spiked with peptide standards and analyzed on a Thermo TSQ Vantage coupled 192to the reversed phase chromatography described in Section 2.4. The 193194 SRM transitions used for the analysis are listed in Supplemental Table 1. For quantification of the Trp-OH and NFK variants the Trp ref-195196 erence peptide was applied as standard, as only Trp and Kyn variants of the peptide could be obtained commercially. As judged by the sim-197ilar ionization efficiencies observed for the Trp and Kyn variant, a sys-198 tematic error of less than 20% can be expected by this approximation. 199

#### 200 3. Results

While indications for oxidative modification of tryptophan resi-201 dues in the ATP synthase in mitochondrial samples from *P. anserina* 202 and other species have been described earlier [14], in this work 203focus was put on the detailed characterization, quantification, and lo-204 calization of these oxidative modifications within the protein com-205plex. Crude mitochondria from juvenile (6 days) and senescent P. 206 anserina cultures were isolated by differential centrifugation. For the 207208investigation three independent cultures from each age stage were used. To determine the role of free iron as a factor for ROS generation, 209mitochondria were isolated from three independent P. anserina indi-210viduals in the presence and absence of the strong iron chelating agent 211 Desferal. 212

#### 213 3.1. Separation of ATP synthase complexes

Fig. 2 displays the separation of solubilized mitochondrial mem-214brane proteins by 2D-BN/SDS-PAGE. BN-PAGE in the first dimension 215separates native monomers and dimers of the ATP synthase, while 216 SDS-PAGE in the second dimension separates the protein subunits 217 constituting these protein complexes. To estimate the apparent 218 mass of the protein complexes in the blue-native gel digitonin-219 solubilized bovine heart mitochondria served as reference [14], as 220these are well characterized and are often used to estimate the 221mass of membrane protein complexes in blue-native gels [35, 36]. 222In contrast to other species, e.g., rats [37], a separated F<sub>1</sub> subcomplex 223is hardly observed in the gels from P. anserina mitochondria. The gel 224 225system resolves the alpha subunits very well into two bands originating from monomeric and dimeric ATP synthase complexes. 226 The alpha subunit contains three tryptophan residues. Apart from 227 the gamma subunit, which contains a single tryptophan, subunit 228 alpha is the only subunit of the  $F_1$  part containing tryptophan in its 229 mature protein sequence. 230

#### 3.2. Identification of tryptophan oxidation site

For localization and structural characterization of tryptophan oxi-232 dation sites in the primary structure of the alpha subunit, bands cor-233 responding to the dimeric and monomeric ATP synthase in the blue-234 native gel (Fig. 2:  $V_2$ ,  $V_1$ ) were digested with trypsin and the resulting tryptic peptides were applied to LC–MS/MS analysis. On average about 6500 MS/MS spectra could be assigned to peptides of ATP synthase subunits, yielding excellent sequence coverage of 30–80% for all proteins of the  $F_1$  part. Although all subunits of the membrane integral  $F_0$  part were also detected, the number of distinct peptides observed was significantly lower due to the reduced number of tryptic cleavage sites in these proteins [38]. For alpha and beta subunits sequence coverage of 50% and 72% respectively, was obtained with 243



**Fig. 2.** Monomers and dimers of ATP synthase from *P. anserina* mitochondria are efficiently resolved in 2D-BN/SDS-gels. In contrast to mitochondrial samples from other species, significant amounts of  $F_1$  subcomplex are not observed in the gels.

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Fig. 4. Relative proportion of different variants of the peptide ILQWEADFLSHLK quantified by the use of isotope-labeled reference peptides.

more than 2500 spectra assigned to peptides of each subunit. Fig. 3A 244 displays a basepeak chromatogram of the analysis. All three Trp sites 245 in the alpha subunit (Table 1) were sequenced by LC-MS/MS spectra 246 from the corresponding tryptic peptides. Trp oxidation, however, could 247 only be observed for  $Trp^{503}$  in the peptide ILQWEADFLSHLK. Species of 248 this peptide with three intermediates of tryptophan oxidation-hydroxyl- 249 tryptophan (Trp–OH), N-formlykynurenine (NFK), and kynurenine 250 (Kyn) (Fig. 1)—were identified with p-values of  $2 \cdot 10^{-6}$  and better. 251 The assignment of these intermediates is unambiguous due to the 252 high quality of the obtained MS/MS spectra (Fig. 3B-E). While the oxi- 253 dation of the peptide could be demonstrated merely by the mass shift 254 of the precursor and an incomplete ion series in the fragment spectra, 255 the presence of  $y_9$  and  $y_{10}$  ions unambiguously proves the oxidation 256 of the Trp residue. The lower retention times as compared to the 257 unmodified peptide (Trp: 68 min, Trp-OH: 51 min, NFK: 63 min, 258 Kyn: 45 min) depict the reduced hydrophobicities of the oxidized 259 Trp species. 260

Two other peptides of the alpha subunit containing Trp residues261(Table 1) were observed, however, any corresponding peptides with262oxidized Trp species could not be detected even when an inclusion263list (Table 1) was applied for the LC-MS analysis.264

#### 3.3. Quantitation of tryptophan oxidation

For a quantitative analysis of the different oxidation intermediates 266 of Trp<sup>503</sup> observed in the LC–MS analysis isotope-labeled reference 267 peptides were applied [34]. Fig. 4 depicts the relative proportion of 268 the different oxidation intermediates of the peptide ILQ- 269 WEADFLSHLK. Under all investigated conditions the NFK form was 270 the most abundant oxidized species. 271

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While the location of the oxidized tryptophan was well defined in 272 all analyzed replicates, and also the amount of oxidized Trp was reproducible in technical replicates, as judged by peak areas of the respective peptides, a significant variation in the amount of oxidized 275 Trp was observed between different biological replicates. In respect 276 to the biological variation (for juvenile and senescent cultures), as 277 well as monomeric and dimeric ATP synthase complexes, no statistically significant difference in the degree of oxidation was detected 279 in the analysis in juvenile versus senescent cultures nor in monomeric versus dimeric ATP synthase. The degree of oxidation was, however, significantly reduced by more than 30%, when the strong iron 282

**Fig. 3.** A) Basepeak chromotogram of a tryptic digest from ATP synthase isolated from BN-gels analysed by LC–MS/MS. The four variants of the peptide ILQWEADFLSHLK are eluted with retention times of 68 min (Trp), 51 min (Trp-OH), 63 min (NFK), and 45 min (Kyn). Oxidized species from the two other tryptophan residues in the ATP synthase alpha subunit are not observed in the analysis. B–E) MS/MS spectra of the four variants of the peptide. The fragment ions  $y_9$  and  $y_{10}$  representing the transition from EADFLSHLK to  $W^*$ EADFLSHLK allow the differentiation of oxidation intermediates.

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chelator Desferal was present during the isolation of mitochondria
suggesting a potential involvement of iron or another transition
metal in the oxidation process.

#### 286 4. Discussion

The ATP synthase is one of the most abundant proteins in the 287 288 inner mitochondrial membrane generating most of the approximately 70 kg ATP utilized in humans every day. It is devoid of redox-active 289prosthetic groups and does not conduct any redox functions. Howev-290er, the regulation of the chloroplast ATP synthase by the redox state is 291 well known [39] and also for the mitochondrial enzyme indications 292293for a redox regulation have been shown recently [40]. Due to its spatial proximity to the mitochondrial electron transport chain, the ATP 294295synthase is a primary target of ROS.

#### 296 4.1. Selectivity of tryptophan oxidation

Using an LC-MS/MS approach the selective targeting of Trp<sup>503</sup> of 297 the ATP synthase alpha subunit for oxidative modification can be un-298ambiguously displayed, while no oxidation can be observed for the 299other Trp residues in the protein complex. Although oxidative stress 300 301 appears as an unstructured and random process, channeling of oxidative damage to defined locations within protein complexes and pro-302 tein subunits appears as a common characteristic of proteins frequently 303 304 exposed to ROS. This selectivity has been observed for mitochondrial proteins-like the aconitase from bovine mitochondria displaying a pre-305 dominant oxidation of Trp<sup>373</sup> [15], as well as for proteins from other sys-306 tems and organelles as the selective N-formylkynurenylation of Trp<sup>365</sup> 307 identified in subunit CP43 of the photosynthetic PSII complex under 308 high-light stress [20]. Recently, a targeting of oxidative modifications to 309 310 specific tryptophan residues was also recognized in pharmaceutical formulations [41] and a detailed analysis could attribute selectivity to a 311 metal binding site in the proximity of the Trp residue. 312

#### 313 4.2. Susceptibility of tryptophan towards oxidation

Although the oxidation of tryptophan is regularly observed in iso-314 lated proteins, in contrast to cysteine and methionine the reactivity 315 towards superoxide anion radicals and hydrogen peroxide is quite 316 low. Taylor and coworkers observed the different susceptibility of 317 tryptophan and methionine towards oxidation, when analyzing pep-318 tides from the 18 kDa subunit of complex I [17], and suggested differ-319 ent oxidation mechanisms for the two amino acids. In pharmaceutical 320 products the low reactivity of H<sub>2</sub>O<sub>2</sub> and peroxides with tryptophan 321 322 [42] is documented, while the reactivity of Trp with singlet oxygen [43] and hydroxyl radicals released by the Fenton reaction is high 323 and is only surpassed by cysteine and methionine [44]. 324

#### 4.3. Localization within the ATP synthase structure

Fig. 5A displays a structural representation of the F<sub>1</sub> portion of the 326 ATP synthase. Due to the high homology with a sequence identity of 327 80%, based on the ATP synthase structure from S. cerevisiae [45] a 328 structure of subunit alpha from P. anserina was generated by homol- 329 ogy modeling using Swiss-Model [46-48]. In the resulting structure, it 330 is evident that all three tryptophans of subunit alpha are exposed to 331 the surface and potentially accessible from the surrounding. Trp<sup>503</sup>, 332 which is the sole target for oxidative modification, is located in the 333 peripheral region of the alpha subunit at a remote distance to catalyt- 334 ic and regulatory nucleotide binding sites or inter-subunit interfaces. 335 A feature distinguishing Trp<sup>503</sup> from the other two Trp sites is the lo- 336 cation of three amino acid residues-Glu<sup>504</sup>, Asp<sup>506</sup> and His<sup>510</sup> in its 337 proximity (Fig. 5B). These residues are able to constitute a binding 338 site for transition metal ion and induce an increased local concentra- 339 tion of these ions in the proximity of Trp<sup>503</sup>. While this sequence pat- 340 tern is well conserved in the subphylum Pezizomycotina and present 341 in 55 of 59 sequences available for subunit alpha from this subdivi- 342 sion in the NCBI database, it appears to be absent in unicellular 343 fungi in the subdivision Saccharomycotina and higher eukaryotes, in 344 which Trp<sup>503</sup> is replaced by a phenylalanine in all known sequences. 345

#### 4.4. Biological implications

As removal of damaged protein is a critical function for the main- 347 tenance of mitochondria [49] and an impairment of protein degrada- 348 tion and repair has been connected to age-associated accumulation of 349 oxidized proteins, for cellular homeostasis efficient means for the 350 perception of oxidative stress are required. While oxidized cysteine 351 and methionine can be efficiently reduced by repair mechanisms 352 [9], tryptophan oxidation products are irreversible and have the po- 353 tential to form markers visible for the mitochondrial quality control 354 system. While an effect of the oxidation of Trp<sup>503</sup> on the enzymatic 355 activity of the ATP synthase is unlikely due to the distance to inter- 356 subunit contact sites and catalytical as well as regulatory nucleotide 357 binding sites, the disturbance of its structural integrity might be suf- 358 ficient to trigger recognition by the protein quality control system. 359 The LON protease, as an important component in the degradation of 360 oxidized proteins in the mitochondrial matrix [50, 51], is induced by 361 the presence of oxidized proteins both in activity and expression 362 level [34, 39]. Although not shown yet, the quality of the mitochon- 363 drial ATP synthase may be controlled by proteases like iAAA protease 364 or mAAA protease inserted in the inner mitochondrial membrane. 365



**Fig. 5.** A) Trp residue 503 (red) is surface exposed and located in the peripheral region of the alpha subunit with a distance of more than 30 Å from both the catalytical and regulatory nucleotide binding sites. The two other Trp residues (blue) are also exposed to the surface of the protein and are also accessible from the solvent. For the generation of a structural model of the alpha subunit of *P. anserina* a Swiss modeller [46–48] was used based on the structure published for *S. cerevisiae* [45] (PDB ID: 2XOK). B) The residues Glu<sup>504</sup>, Asp<sup>506</sup> and His<sup>510</sup> located in the proximity of Trp<sup>503</sup> at a distance of 5–9 Å are highlighted in a close-up of the structure. These residues are able to constitute a metal ion binding site. No such site can be defined for the two other Trp residues in the protein. For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.

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With its reactivity towards hydroxyl radicals and singlet oxygen, 366 367 tryptophan oxidation might mediate a sensory function for strong ROS or in connection with the putative metal binding site could be in-368 369 volved in the perception of metal catalyzed protein oxidation. Due to its abundance and its location in the inner mitochondrial membrane 370 the ATP synthase might be able to perceive compounds with diffusion 371 ranges limited by short half-lives [52] and mediate an up-regulation 372 of the protein turnover before more sensible components are 373 374 affected.

375 The channeling of oxidative damage to specific subunits and do-376 mains is a common feature observed in different enzymes evolved under the evolutionary pressure imposed by oxidative stress, like 377 the D1 protein of the photosystem II as a prominent example [53] 378 379 which accounts for up to 50% of protein synthesis [54, 55] in chloroplasts under high-light conditions. This focuses the effort for mainte-380 nance of protein complexes to the degradation and replacement of a 381 single protein subunit [56] and, thereby transforms the random and 382 untargeted process releasing ROS into a unidirectional signal that 383 can be perceived by the quality control system. 384

Supplementary materials related to this article can be found on-385 line at doi:10.1016/j.bbabio.2011.11.006. 386

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