





Cellular studies of the two main isoforms of human **D-aspartate** oxidase

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Human D-aspartate oxidase (hDASPO) is a FAD-dependent enzyme responsible for the degradation of p-aspartate (p-Asp). In the mammalian central nervous system, D-Asp behaves as a classical neurotransmitter, it is thought to be involved in neural development, brain morphology and behavior, and appears to be involved in several pathological states, such as schizophrenia and Alzheimer's disease. Apparently, the human DDO gene produces alternative transcripts encoding for three putative hDASPO isoforms, constituted by 341 (the 'canonical' form), 369, and 282 amino acids. Despite the increasing interest in hDASPO and its physiological role, little is known about these different isoforms. Here, the additional N-terminal peptide present in the hDASPO 369 isoform only has been identified in hippocampus of Alzheimer's disease female patients, while peptides corresponding to the remaining part of the protein were present in samples from male and female healthy controls and Alzheimer's disease patients. The hDASPO 369 isoform was largely expressed in E. coli as insoluble protein, hampering with its biochemical characterization. Furthermore, we generated U87 human glioblastoma cell clones stably expressing hDASPO 341 and, for the first time, hDASPO_369 isoforms; the latter protein showed a lower expression compared with the canonical isoform. Both protein isoforms are active (showing similar kinetic properties), localize to the peroxisomes, are very stable (a half-life of approximately 100 h has been estimated), and are primarily degraded through the ubiquitin-proteasome system. These studies shed light on the properties of hDASPO isoforms with the final aim to clarify the mechanisms controlling brain levels of the neuromodulator D-Asp.

Introduction

The FAD-dependent flavoenzyme D-aspartate oxidase (DASPO or DDO, EC 1.4.3.1) was first identified in the '50s in rabbit kidney and liver [1] and now it is known to be widely present in eukaryotes, ranging from fungi to humans [2]. In mammals, the enzyme is primarily involved in the

catabolism of acidic D-amino acids and the best substrate is D-aspartate (D-Asp). DASPO catalyzes their oxidative deamination into the corresponding α-ketoacids, along with the production of hydrogen peroxide and ammonia [3]. Neutral and basic D-AAs are similarly deaminated by the homologous

Abbreviations

CHX, cycloheximide; CQ, chloroquine; DAAO, p-amino acid oxidase; p-Asp, p-aspartate; DASPO, p-aspartate oxidase; p-Ser, p-serine; hDASPO, human p-aspartate oxidase; MG132, benzyloxycarbonyl-L-leucyl-L-leucyl-L-leucinal; NMDAR, N-methyl-p-aspartate receptors; UPS, ubiquitin-proteasome system.

flavoenzyme, D-amino acid oxidase (DAAO, EC 1. 4.3.3) [4].

In mammals, free p-Asp has been reported to play different roles: In the endocrine system, it modulates steroidogenesis and the synthesis and release of several hormones [5–8], while in the central nervous system, it stimulates mGlu5 and presynaptic AMPA receptors [9,10], as well as N-methyl-D-aspartate receptors (NMDAR), acting as agonist [11]. In the brain, D-Asp is abundant during embryonic and perinatal phases and drastically decreases later on [7,12]. This peculiar temporal distribution pattern is due to the concomitant onset of DASPO expression and activity, mainly observed in neurons [12,13]. D-Asp has been proposed as a signaling molecule involved in neural development, brain morphology, and behavior, and the postulated role of DASPO in strictly regulating its levels has been strengthened [11,14,15].

In this regard, several studies performed in animal models demonstrated that the persistent deregulation of D-Asp levels causes age-dependent effects: an improvement of spatial memory and cognitive abilities in young individuals is followed by a rapid deterioration of learning and memory, leading to precocious brain aging [16,17]. A protective role of DASPO has been proposed: the enzyme would prevent NMDAR hyperstimulation through the strict regulation of postnatal brain levels of D-Asp.

Despite the increasing interest in hDASPO properties and physiological role, little is known about processes involved in the regulation of its activity at the cellular level. Notably, the UniProtKB database reports three different isoforms of hDASPO (identifier Q99489) encoded by alternative transcripts of the human DDO gene: (a) Isoform 1 (hDASPO_341, 341 amino acids), referred to as the 'canonical isoform', is homologous to the single protein form in rodents; (b) isoform 2 (hDASPO_282), apparently originated by alternative splicing of the transcript, whose sequence is identical to the hDASPO_341 one but lacks of 59 residues in the central region (residues 95-153 in the canonical isoform); (c) isoform 3 (hDASPO 369), which appears highly conserved in primates, is characterized by the presence of 28 additional N-terminal residues, probably due to the recognition of an upstream alternative start codon (Fig. S1). The shorter protein isoforms were produced in E. coli [18,19], while the longer one has never been expressed. Notably, the recombinant deleted hDASPO 282 form was largely produced as inclusion bodies [19]. The three-dimensional structure of human DASPO 341 (hDASPO, PDB entry code PDB: 6RKF), and an extensive characterization of its biochemical properties, has been recently published [18].

In order to look deep inside into the role of the two main hDASPO isoforms, here we investigated by a proteomic approach the presence of hDASPO in hippocampus of male and female healthy subjects and Alzheimer's disease (AD) patients: the hDASPO_369 isoform was identified exclusively in the hippocampus of female AD patients. The functional properties, the degradation kinetics, and the mechanisms involved in protein turnover were investigated by ectopically expressing hDASPO_341 and hDASPO_369 isoforms in the U87 human glioblastoma cell line. This study demonstrated that both hDASPO isoforms are active (thus, able to control p-Asp cellular level), are highly stable, and mainly degraded through the ubiquitin-proteasome system.

Results

Identification of hDASPO isoforms in human hippocampus by nLC-MS/MS

Hippocampus is a brain area relevant for the formation and consolidation of memory, and it is specifically vulnerable to damage at early stages of AD. We investigated the presence of hDASPO in hippocampus samples from male and female healthy controls and AD patients by using a nLC-MS/MS approach. The results, briefly summarized in Table 1 and Fig. S1, clearly show that the N-terminal peptide (exclusively present in the hDASPO_369 isoform) was identified in female AD samples only (1_MRPARHWETR_10_peptide) while peptides covering the regions common to both 341 and 369 isoforms were identified in all samples.

Expression of hDASPO_369 in E. coli cells

The synthetic gene encoding the His-tagged hDASPO_369 was subcloned into the pET11a expression vector, and the protein was expressed in *E. coli* BL21(DE3) LOBSTR cells using the procedure reported for hDASPO_341 [18]. Differently from the latter, the hDASPO_369 isoform was largely expressed as inclusion bodies. The use of the ArcticExpress (DE3) *E. coli* strain (expressing the 'cold adapted' chaperons Cpn10 and Cpn60) and of conditions favoring heterologous proteins folding, that is incubation at 13 °C for 24 h after adding the inducer, yielded a very low amount of soluble recombinant protein (not shown). The purification of hDASPO_369 from the crude extract by HiTrap chelating chromatography

Table 1. List of the hDASPO peptides identified by mass spectrometry in hippocampus samples from male and female healthy controls and Alzheimer's disease (AD) patients. The N-terminal peptide present in the hDASPO_369 isoform is marked in bold. The numbering refers to the hDASPO_369 isoform.

Sequence	Control		AD	
	Male	Female	Male	Female
1-MRPARHWETR-10	_	_	_	X
₁₄₇₋ MTEAELKKFPQYVFGQAFTTLK ₋₁₆₈	Χ	_	_	X
₁₅₅₋ FPQYVFGQAFTTLK ₋₁₆₈	_	_	Χ	_
₁₈₄₋ GSGGWTLTR ₋₁₉₂	Χ	Χ	_	Χ
₁₉₄₋ IEDLWELHPSFDIVVNCSGLGSR ₋₂₁₆	_	Χ	_	_
₂₆₆₋ QKGDWNLSPDAENSR ₋₂₈₀	Χ	_	_	_
₂₆₆₋ QKGDWNLSPDAENSREILSR ₋₂₈₅	Χ	_	Χ	Χ
₂₆₈₋ GDWNLSPDAENSREILSRCCALEPSLHGACNIR ₃₀₀	_	_	_	Χ
₂₆₈₋ GDWNLSPDAENSREILSR ₋₂₈₀	-	X	-	_
₃₁₄₋ LQTELLAR ₋₃₂₁	Χ	-	-	_

yielded $\leq 0.06~\text{mg}\cdot\text{L}^{-1}$ of fermentation broth, an amount inadequate for performing kinetic and spectral studies.

Solubilization and refolding of hDASPO_369 from inclusion bodies was attempted using different procedures and conditions (Table S1): Once more we failed in obtaining a significant amount of the soluble protein isoform. In line with protein expression results, bioinformatic evaluation of protein solubility by four different methods predicted a significantly lower solubility for the longer isoform compared with the canonical hDASPO 341 isoform (Table S2).

Expression of hDASPO isoforms in U87 cells

Expression plasmids were generated by subcloning the sequences encoding the hDASPO 341 hDASPO 369 isoforms in the pcDNA3 vector. The corresponding DNA fragments were produced by PCR amplification of hDASPO cDNA using different 5'-primers designed to anneal to the alternative ATG sites and to insert, beside two unique restriction sites, a 5'sequence coding for 3 copies of the FLAG epitope (N-terminal 3XFLAG). The generated pcDNA3 3XFLAG-hDASPO 341 pcDNA3 3XFLAGand hDASPO_369 expression constructs were used to transfect human glioblastoma U87 cells, and cell clones stably expressing the long or the short hDASPO isoform were selected.

Western blot analysis on selected cell clones confirmed the expression of both hDASPO_341 and hDASPO_369 isoforms (Fig. 1A), although the longer one at a fourfold lower level (Fig. 1B). The western blot analysis of U87 3XFLAG-hDASPO_341 cell lysates with the anti-hDASPO antibody recognized a band at the expected molecular mass (\approx 42.0 kDa, corresponding to 3XFLAG-hDASPO_341) and a band

at \approx 37.0 kDa. On the other hand, the same analysis performed on U87 3XFLAG-hDASPO 369 cell lysates revealed the presence of three bands: In addition to the 37 kDa band, signals at ≈ 40.5 and 44.0 kDa were also apparent (Fig. 1A). In both cell clones, the band at the lowest molecular mass, ≈ 37.0 kDa, should correspond to an aspecific signal since it was also present in control samples (U87 untransfected cells). Based on the molecular mass, the bands at ≈ 40.5 and 44 kDa correspond to untagged hDASPO 341 and 3XFLAGhDASPO 369 isoforms, respectively: Both bands were recognized by the anti-hDASPO antibody, while only the latter one was detected by the anti-FLAG antibody (not shown). Therefore, U87 cells transfected with the pcDNA3 3XFLAG-hDASPO 369 construct expressed both protein isoforms at comparable levels (Fig. 1B).

The identity of the expressed protein isoforms in the U87 3XFLAG-hDASPO 369 cells was determined using immunoprecipitation (IP) experiments. In detail, two consecutive IP rounds were carried out: the first one using the whole cell extract and the anti-FLAG M2 affinity resin; the second using the obtained post-IP sample and the anti-hDASPO antibody cross-linked to Dynabeads Protein G. Western blot analyses of the immunoprecipitated samples confirmed that both 3XFLAG-hDASPO 369 and the untagged hDASPO 341 isoforms were present in the cell extract (Fig. 1C,D). The untagged hDASPO 341 originates by the translation of the hDASPO 369 encoding transcript at an alternative, downstream starting codon.

hDASPO isoforms are active

A sensitive fluorimetric assay using the Amplex Ultra-Red Reagent and based on the detection of hydrogen peroxide produced by hDASPO in the presence of a saturating concentration of the substrate p-Asp

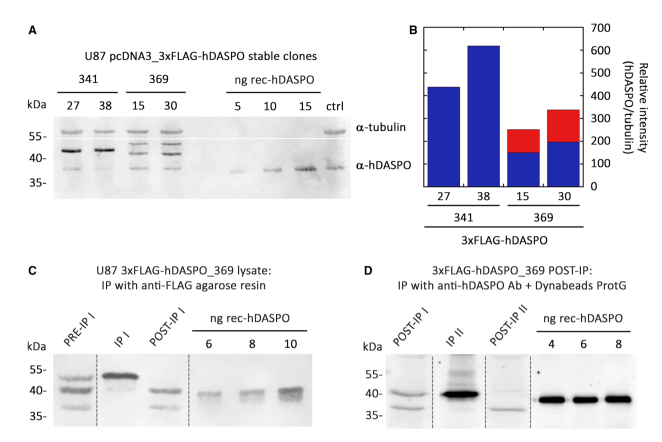


Fig. 1. Expression of hDASPO isoforms in U87 glioblastoma cells. (A, B) Relative expression levels of the protein isoforms in U87 cell clones stably transfected with pcDNA3 3XFLAG-hDASPO_341 (clones 27 and 38) or pcDNA3 3XFLAG-hDASPO_369 (clones 15 and 30) detected by western blot analysis using anti-hDASPO antibody (A) and corresponding densitometric analysis (panel B; blue bars: hDASPO_341; red bars: hDASPO_369). Amounts of sample corresponding to 5×10^4 cells were loaded. Ctrl represents the untrasfected cells sample, and recombinant hDASPO (38.6 kDa) was used as a positive control. Western blot analyses were repeated twice, and the reported densitometric values represent the average. (C, D) IP studies aimed to confirm the identity of the hDASPO forms detected in lysates of U87 cells transfected with pcDNA3 3XFLAG-hDASPO_369. The presence of hDASPO was revealed by western blot analysis using anti-hDASPO antibody: (C) 3XFLAG-hDASPO_369 (44 kDa) was immunoprecipitated from the whole cell lysate using an anti-FLAG agarose resin, and (D) the ensuing post-IP sample (showing bands at 37 and 40.5 kDa) was further immunoprecipitated using anti-hDASPO antibodies conjugated to Dynabeads Protein G. Bands corresponding to 3XFLAG-hDASPO_369 (44 kDa) and hDASPO_341 (40.5 kDa) were observed in the IP I and IP II samples, respectively. Dashed lines represent noncontinuous lanes. Immunoprecipitation experiments were repeated twice.

(16.7 mm) [20] was used to assess the enzymatic activity of the two protein isoforms. 3XFLAGhDASPO 369 and 3XFLAG-hDASPO 341 were immunoprecipitated from the corresponding cell lysates using an anti-FLAG M2 affinity resin, and the activity assays were performed on the purified protein form. The specific activity of the two hDASPO isoforms, calculated by normalization for the amount of the immunoprecipitated protein in each well, as assessed by western blot analysis, was very similar $(34.4 \pm 7.6 \text{ and } 38.7 \pm 6.9 \text{ U} \cdot \text{mg}^{-1} \text{ for the 3XFLAG}$ hDASPO 369 and 3XFLAG-hDASPO 341 isoforms, respectively) and slightly lower than the value determined for the recombinant purified hDASPO $(55.1 \pm 3.9 \text{ U} \cdot \text{mg}^{-1})$. Immunoprecipitated

were also used to evaluate the apparent kinetic properties on D-Asp: The two isoforms show similar $k_{\rm cat}$ and $K_{\rm m}$ values (i.e., 27.8 ± 0.6 and $27.4 \pm 1.3~{\rm s}^{-1}$ and 0.44 ± 0.04 and $0.29 \pm 0.07~{\rm mm}$ for hDASPO_369 and hDASPO_341, respectively). This method gave for the recombinant hDASPO a $k_{\rm cat}$ of $76.4 \pm 7.1~{\rm s}^{-1}$ and a $K_{\rm m}$ of $2.04 \pm 0.60~{\rm mm}$ which are comparable to the previously reported values ($81.3 \pm 1.5~{\rm s}^{-1}$ and $1.05 \pm 0.06~{\rm mm}$) [18].

hDASPO expression controls cellular levels of D-aspartate

HPLC analysis of amino acid enantiomers showed that the ectopical expression of the 3XFLAG-hDASPO_341

isoform deeply affects D-Asp cellular content: Basal D-Asp levels $(0.044 \pm 0.006 \text{ nmol} \cdot \text{mg}^{-1} \text{ proteins in con-}$ trol cells transfected with the pcDNA3 empty vector) were fully depleted in 3XFLAG-hDASPO 341 expressing cells (Fig. 2A), whereas the L-enantiomer content was unaffected (5.97 \pm 1.25 and 6.67 \pm 1.14 nmol·mg⁻¹ proteins in transfected and control cells, respectively). Accordingly, a dramatic decrease in the D-Asp/total Asp ratio in the transfected cells was also evident (Fig. 2A, right panel). On the other hand, serine (Ser) enantiomer levels appeared only marginally altered by 3XFLAG-hDASPO 341 expression: L-Ser slightly increased (12.7 ± 5.2) content $8.2 \pm 0.9 \; \mathrm{nmol \cdot mg^{-1}}$ proteins in transfected cells compared with controls, respectively) and the D-Ser/total Ser ratio remained unchanged (Fig. 2B).

hDASPO isoforms are peroxisomal enzymes

The peroxisomal targeting of hDASPO should be determined by the noncanonical C-terminal PTS1

sequence (-SNL). The subcellular localization of hDASPO 369 and hDASPO 341 isoforms was verified by immunostaining and confocal analysis performed on the stable clones expressing the two protein isoforms and on U87 untransfected cells as a control. The ectopically expressed 3XFLAG-hDASPO 369 and 3XFLAG-hDASPO 341 were specifically detected using either the anti-FLAG or the anti-hDASPO antibody (no immunorecognition was observed in control cells, Fig. 3). Both protein isoforms showed a punctate distribution within the cells (Fig. 3B,C,E,F) consistent with their targeting to the peroxisomal compartment, as demonstrated by the large overlapping of FLAG immunofluorescence signals to PMP70 ones (red and green channels, respectively; Fig. 3B,C, merge panels). Conversely, no signal overlapping was apparent when the protein isoforms were costained with the antimitochondria antibody (green and red channels, respectively; Fig. 3E,F, merge panels). In this case, we used the anti-hDASPO antibody, which allowed to detect both the 3XFLAG-hDASPO 369 and the untagged

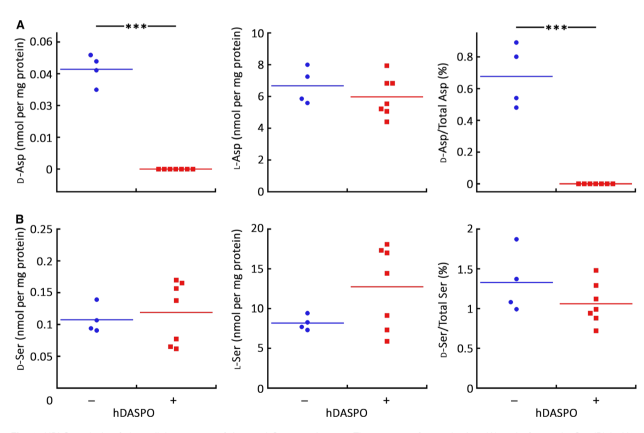


Fig. 2. HPLC analysis of the cellular content of Asp and Ser enantiomers. The amount of D- and L-Asp (A) and of D- and L-Ser (B) in U87 cells stably transfected with pcDNA3_3xFLAG-hDASPO_341 and in control cells transfected with the empty vector (red and blue values, respectively) was measured and normalized for the total protein content in each sample. The D-enantiomer/total stereoisomer ratio is also reported (right panels). ***P < 0.0001. Data were analyzed by unpaired parametric t-test. Graphs report single data point (dots) as well as mean values (n = 3; lines).

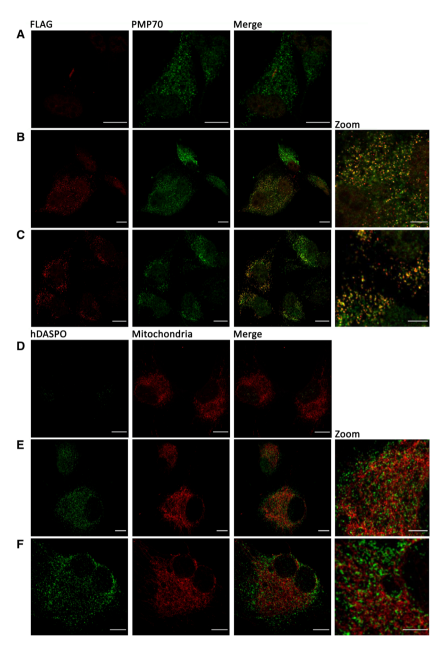


Fig. 3. Confocal analysis of U87 stable clones expressing 3XFLAG-hDASPO protein isoforms and control cells. U87 untransfected cells, as well as 3XFLAG-hDASPO_369 (B, E) and 3XFLAG-hDASPO_341 (C, F) expressing cells, were double-stained with the anti-FLAG and the anti-PMP70 (A–C) or the anti-hDASPO and the anti-mitochondria antibodies (D–F). (A, D) Control cells showed no (aspecific) signals with either the red (A) or the green (D) channel, respectively. (B, C) 3XFLAG-hDASPO_369 (B) and 3XFLAG-hDASPO_341 (C) signals (red channel) largely overlapped with PMP70 ones (green channel), strongly indicating a peroxisomal localization for both protein isoforms. (E, F) No signal overlapping with the signal for mitochondria was instead observed for both the 3XFLAG-hDASPO_369 (E) and _341 (F) hDASPO isoforms (compare the signal distribution in the merge panels). Scale bars: 10 μm. The reported images are representative of 15 acquired images for each of the stable clones in the different staining conditions.

hDASPO_341 isoforms expressed by U87 3XFLAG-hDASPO_369 cells: the same punctate distribution pattern was evident. Notably, both protein isoforms were not detected in the cytosol, at least at significant levels.

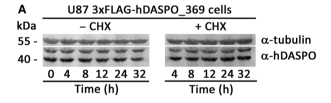
hDASPO isoforms are long-lived proteins

The cellular stability of 3XFLAG-hDASPO_369 and 3XFLAG-hDASPO_341 was investigated by treatment with cycloheximide (CHX), an inhibitor of protein

synthesis that prevents translational elongation. Western blot analysis showed only a slight decrease in both the FLAG-tagged hDASPO isoforms abundance during time: about 80% of the initially observed protein isoform was still detectable at 32 h after the treatment (Fig. 4). Thus, for both proteins, a half-life of approximately 100 h was estimated. Furthermore, the N-terminal 3XFLAG additional sequence did not appear to influence the kinetics of protein degradation, since the same half-life value was also determined for the untagged hDASPO_341 expressed in the U87 cell clones transfected with the pcDNA3_3XFLAG-hDASPO_369 construct (Fig. 4). These results indicated that hDASPO is a long-lived protein, as it was previously reported for hDAAO [21].

hDASPO isoforms are degraded by the ubiquitinproteasome system

The mechanisms involved in hDASPO degradation were investigated by treating U87 cell clones expressing 3XFLAG-hDASPO_369 or 3XFLAG-hDASPO_341 isoforms with inhibitors of autophagy or of the ubiquitin–proteasome system (blocking the preferential degradation pathway should lead to the accumulation of the protein), upon incubating the cells overnight under starvation conditions (i.e., in culture media containing 1%



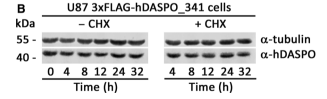


Fig. 4. Analysis of hDASPO degradation rate. U87 cells stably expressing 3XFLAG-hDASPO_369 or 3XFLAG-hDASPO_341 isoforms were treated with 100 μ g·mL⁻¹ CHX. (A) In U87 3XFLAG-hDASPO_369 CHX-treated cells, only a small decrease (~ 20% at 32 h) in the signal detected at 39 kDa (untagged hDASPO_341) and 44 kDa (3XFLAG-hDASPO_369) by anti-hDASPO antibodies was apparent compared with the control. (B) Similarly, a small change in the intensity of the 42 kDa band corresponding to 3XFLAG-hDASPO_341 was observed following CHX addition, thus indicating that both the protein isoforms are highly stable. Values are the mean \pm SD (n = 4–8), normalized to tubulin, and expressed relatively to the control without CHX (i.e., cells collected at the same time after the addition of PBS).

FBS). Starvation is a physiological stimulus inducing numerous changes within the cell. Two out of the major effects are a decreased protein synthesis and the activation of proteolytic pathways [22]; accordingly, under starvation conditions the effect of the inhibitors of protein degradation pathways should be enhanced.

Since autophagy is responsible for cytoplasmic bulk degradation and is thought to be important for the turnover of whole organelles and long-lived proteins [23,24], we investigated the effect of two autophagy inhibitors, namely chloroquine (CQ, 75 µм) and ammonium chloride (NH₄Cl, 10 mm), on the cellular levels of hDASPO variants. The chosen inhibitors act by different mechanisms: CQ inhibits autophagy mainly by impairing autophagosome fusion with lysosomes, whereas NH₄Cl acts on the proton gradient affecting lysosomal pH and thus the degradative activity of hydrolytic enzymes within this organelle [25,26]. No difference in the relative abundance of 3XFLAGhDASPO 369 and 3XFLAG-hDASPO 341 was detected in treated cells compared with controls (Fig. 5A), despite CQ and NH₄Cl treatments were prolonged up to 10 h, suggesting that they do not affect the turnover of these protein variants. This result was unexpected since hDAAO, which is also targeted to peroxisomes and shares a high degree of sequence identity and enzymatic functionality with hDASPO, is primarily degraded by the lysosome/endosome system [21].

The effect of MG132 on hDASPO cellular level was also investigated. This compound acts as a proteasomal inhibitor interacting with the 'chymotrypsin-like' component and blocking one or more peptidases within the 20S proteasome core, without affecting protein-ubiquitinating and -deubiquitinating enzymes [27]. The treatment of stably transfected cells with 25 μM MG132 led to the gradual and moderate accumulation of both 3XFLAG-hDASPO 369 and 3XFLAGhDASPO 341 proteins (Fig. 5B): a 1.6-fold increase at 6-10 h after treatment was detected for both isoforms. This resembles what was previously observed for hDAAO (i.e., a 1.5-fold increase) [21], and eventually suggests a slow synthesis of both flavoproteins under starvation conditions. hDASPO 369 hDASPO 341 behaved again exactly in the same way, indicating that the N-terminal sequence (absent in the shorter isoform) did not affect the degradation mechanism as well as the targeting to, and the recognition by, the ubiquitin-proteasome system (UPS).

Ubiquitination of hDASPO isoforms

hDAAO was demonstrated to form ubiquitin conjugates by *in vitro* and cellular studies, and it was

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Fig. 5. Effect of inhibitors of the degradation pathways on 3XFLAG-hDASPO_369 and 3XFLAG-hDASPO_341 cellular levels. (A) The cell clones were incubated under starvation conditions and treated with 75 μM CQ or PBS as a control, for up to 10 h. Western blot and quantification analysis showed no variation in the amount of 3XFLAG-hDASPO_369 (red bars) or 3XFLAG-hDASPO_341 (blue bars) in treated cells compared with controls. (B) U87 cells stably expressing 3XFLAG-hDASPO_369 or 3XFLAG-hDASPO_341 under starvation were treated with 25 μM MG132 or 0.1% DMSO for up to 10 h. Western blot analysis using anti-hDASPO antibody and quantitative analysis demonstrated that both 3XFLAG-hDASPO_369 and 3XFLAG-hDASPO_341 significantly accumulated after adding the inhibitor. For each time point, the value is expressed as the mean \pm SD (n = 4), normalized to tubulin, and expressed relative to the control (i.e., cells collected at the same time after adding PBS or DMSO, see left lanes in western blot panels).

proposed that this signal drives its targeting to the UPS during the protein degradation [21]. *In vitro* ubiquitination experiments were carried out by mixing 5 μg of purified recombinant hDASPO (i.e., hDASPO_341 isoform), 0.6 mg·mL⁻¹ ubiquitin, 5 mm ATP, 1 mm dithiothreitol, 2 mm MgCl₂, 1 μm ubiquitin aldehyde (to inhibit deubiquitinating enzymes), 25 μm MG132 (to block the proteasome degradation activity), and 4 mg·mL⁻¹ U87 cell extract (to provide the components of the UPS). Upon incubation for 60 min at 37 °C, aliquots of the reaction mixtures were analyzed

by western blot. Distinct bands at high molecular mass, corresponding in size to mono- (44.5 kDa), multi-, or polyubiquitinated (57.2–102 kDa) hDASPO species, were observed, see Fig. 6A: The same bands were absent in control samples (i.e., reaction mixtures prepared by omitting the recombinant hDASPO or ubiquitin). The assay of the enzymatic activity on the same samples (using saturating concentration of substrate and cofactor, i.e., at 15 mm D-Asp and 40 µm FAD) showed no change in hDASPO enzymatic activity (Fig. 6B).

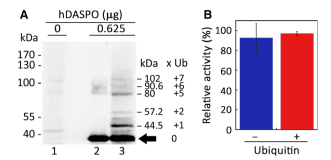


Fig. 6. In vitro ubiquitination assay of recombinant hDASPO. The purified recombinant hDASPO (5 µg) was incubated with U87 cell extracts (4 mg·mL⁻¹), ubiquitin, ATP, MG132, and ubiquitin aldehyde (in a final volume of 60 uL) and incubated at 37 °C for 60 min. To detect the formation of hDASPO-ubiquitin conjugates. an aliquot of the reaction mixture (corresponding to 0.625 ug of recombinant protein) was analyzed by western blotting using antihDASPO antibody. (A) Distinct bands, corresponding to distinct hDASPO-modified forms (mono- or polyubiquitinated), were evident in the reaction mixture only (lane 3) and absent in control samples (prepared by omitting ubiquitin, lane 2, or using the same volume of the reaction mixture without adding the recombinant protein, lane 1). The arrow indicates the unmodified protein. In vitro ubiquitination experiments and subsequent western blot analysis were repeated three times. (B) Activity assays performed on the in vitro ubiquitination and control mixtures revealed that recombinant hDASPO activity is not affected by the modification. Activity measurements are reported as relative activity after the incubation at 37 °C compared with the starting value. Data represent mean values \pm standard deviation (n = 3).

The ubiquitination state of hDASPO isoforms at the cellular level was investigated by IP experiments on the U87 cell clones stably overexpressing the hDASPO isoforms and transiently transfected with a FLAGtagged ubiquitin expression construct. The highest expression level of FLAG-ubiquitin was observed 48 h after transfection with the pCMV-FLAG-ubiquitin vector (data not shown). Accordingly, cells were treated 48 h after transfection with 25 µm MG132 and collected 6 h after treatment: cell lysates were then immunoprecipitated with the anti-FLAG M2 affinity resin. Western blot analysis indicated that both hDASPO isoforms have been modified by ubiquitin: an intense anti-hDASPO signal was detected at high molecular mass in the immunoprecipitated protein samples from both 3xFLAG-hDASPO_369 3XFLAG-hDASPO_341-expressing cells (Fig. 7, lanes 1 and 4, respectively). The detected signal is specific for the ubiquitin-conjugated protein isoforms since it is absent in U87 control cells transiently transfected with the FLAG-ubiquitin expression vector and treated with MG132 (Fig. 7, lane 7). Notably, an antihDASPO immunorecognition signal was also evident

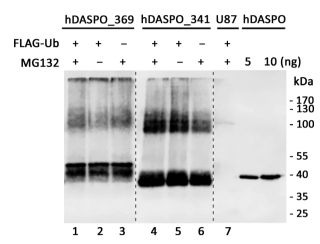


Fig. 7. Ubiquitination analysis in U87 cells stably expressing hDASPO isoforms and transiently transfected with FLAG-ubiquitin. Forty-eight hours after transfection with pCMV-FLAG-ubiquitin, U87 control cells (lane 7) and cells stably expressing the 3XFLAG-hDASPO_369 (lanes 1–3) or 3XFLAG-hDASPO_341 (lanes 4–6) isoforms were treated with 25 μm MG132 for 6 h, and then, proteins were immunoprecipitated under denaturing conditions using anti-FLAG M2 Affinity resin. Cells transfected but not treated with the proteasome inhibitor or not transfected and added with MG132 were processed as controls. Dashed lines represent noncontinuous lanes. The presence of ubiquitinated-hDASPO was revealed by western blot analysis using anti-hDASPO antibody. The experiment was repeated twice.

in U87 cell clones expressing 3xFLAG-hDASPO_369 or 3XFLAG-hDASPO_341 not treated with the proteasome inhibitor or not overexpressing FLAG-ubiquitin and subjected to MG132 treatment (Fig. 7, lanes 2–3 and 5–6). Altogether, both hDASPO isoforms are polyubiquitinated at the cellular level.

Discussion

In the brain of mammals, D-Ser is critical for neurotransmission since it modulates the activation state of NMDAR by acting as the main endogenous co-agonist of this receptor. For this reason, its synthesis, transport, and degradation processes have been characterized in depth [28–30]. A second p-amino acid, p-Asp, is also involved in neurotransmission, due to its relatively high affinity for the glutamate-binding site of NMDAR. D-Asp shows a peculiar temporal occurrence in the brain, with its levels peaking during early embryonic development and drastically falling after birth [6,7,15]: The regulation of D-Asp levels is mainly exerted by the catabolic activity of hDASPO. This peroxisomal flavoenzyme, together with hDAAO (which is responsible for the selective degradation of D-Ser) [31], belongs to the amino acid oxidase family of flavoproteins. Despite the two orthologue flavoproteins share a high sequence identity and a similar overall tertiary structure, hDASPO and hDAAO profoundly differ in their quaternary structure, cofactor binding affinity, kinetic properties, and mechanisms [18]. These findings indicate that they should regulate the brain levels of D-Asp and D-Ser differently.

Interestingly, three different isoforms of hDASPO can be identified in the UniProtKB database: hDASPO 341 (37.5 kDa, the 'canonical' protein isoform), hDASPO_282 (30 kDa), and hDASPO_369 (41 kDa). We focused on the hDASPO 341 and hDASPO 369 isoforms. For the first time, we demonstrated that the latter longer isoform is expressed in the hippocampus of female AD patients (Table 1). This is an intriguing observation that pushed the investigation of the biochemical properties of the two hDASPO isoforms. Unluckily, the recombinant hDASPO 369 cannot be characterized since it is largely produced in E. coli cells as an insoluble protein. Both protein isoforms were expressed in U87 cells after transfection: The 3XFLAG-hDASPO 369 isoform was expressed at a significantly lower level (Fig. 1). Notably, the cells transfected with the pcDNA3 3XFLAG-hDASPO 369 construct produced both the long and the untagged short protein isoforms at similar levels (Fig. 1B), suggesting that the translation of hDASPO 369 encoding transcript can occur from the alternative starting codon at a similar frequency. Notably, both hDASPO 369 and hDASPO_341 isoforms were efficiently targeted to the peroxisomes and showed similar kinetic properties on D-Asp. Furthermore, the ectopically expressed hDASPO 341 isoform fully depleted D-Asp cellular pool while not significantly affected the levels of D-Ser and the corresponding L-enantiomers (Fig. 2).

Similarly to the homologous hDAAO protein [21], both hDASPO isoforms were highly stable proteins and likely characterized by a slow cellular turnover: Their estimated half-life was $\approx 100 \text{ h}$ (Fig. 4), even higher than that for catalase (≈ 30 h) [32]. This value is in line with the peroxisomal localization of hDASPO isoforms, since most data indicate a peroxisomal halflife of around 2 days [33]: peroxisomes are thought to be mainly degraded by autophagy, with a process mediated by p62 and LC3-II named pexophagy [33] that can be upregulated in response to different stresses, such as nutrient starvation. Accordingly, ectopically expressed hDAAO accumulated in U87 cells in this condition upon blocking the lysosome/endosome pathway [21]. On the contrary, hDASPO 341 and hDASPO_369 isoforms were degraded by the UPS.

Actually, *in vitro* and cellular studies revealed that hDASPO isoforms can be ubiquitinated (Figs 6A and

7), a modification that did not affect the enzymatic activity (Fig. 6B). This observation suggests that (poly)ubiquitin conjugation represents the cellular signal that triggers the targeting of hDASPO to the UPS and thus that the modified flavoenzyme should be retrotraslocated to the cytoplasm for degradation. It is known that excessive peroxisomal matrix proteins may be exported to the cytosol where they are degraded by cytosolic proteases or the proteasome [34]. In this context, it is noteworthy that peroxisomal membrane receptors involved in the import apparatus (e.g., Pex5p, Pex7p, and Pex20p) can also be degraded by the UPS [34] and that the ubiquitination pattern controls their targeting: When the export is impaired, they are polyubiquitinated and extracted from the peroxisomal membrane for degradation by the UPS, a process called RADAR (Receptor Accumulation and Degradation in the Absence of Recycling) [35]. Since the RADAR quality control pathway is most likely conserved in mammals, it is tempting to speculate that this mechanism is involved in the regulation of hDASPO cellular levels.

We can conclude that the additional N-terminal sequence in the long hDASPO isoform does not affect the enzyme functionality, subcellular localization, half-life, and degradation pathway. The main difference is a lower solubility for the longer isoform and its selective expression in the hippocampus of AD female patients. In order to understand how the cellular levels of D-Asp are controlled, further studies will focus on the effects due to post-translational modifications and ligand interactions on hDASPO isoforms under physiological and pathological conditions.

Materials and methods

Expression vectors

The cDNA encoding hDASPO was purchased as a full ORF clone from the PlasmID Repository at Harvard Medical School (HsCD00335325 pCMV-SPORT6) and amplified with primers, reported in Table S3, specifically designed to obtain the two different isoforms with three copies of the FLAG (DYKDDDK) epitope fused to the N-terminal end. These cDNAs were then subcloned into pcDNA3 vector, and the resulting constructs were confirmed by automated sequencing. The pCMV-FLAG-ubiquitin vector for the expression of N-terminal FLAG-tagged human ubiquitin was kindly provided by professor Herman Wolosker (Technion—Israel Institute of Technology, Haifa, Israel).

For *E. coli* expression, the synthetic cDNA coding for hDASPO_369 (produced by GeneArt®) was designed to

contain an N-terminal 6xHis encoding sequence and was optimized in the codon usage [18]. It was amplified by PCR using the oligos reported in Table S3 and subcloned in the pET11a expression vector (Novagen, Merck KGaA, Darmstadt, Germany) using the inserted restriction sites.

E. coli expression systems

Heterologous expression of hDASPO_369 in *E. coli* cells was performed in BL21(DE3) LOBSTR (Kerafast Inc., Boston, MA, USA) and ArcticExpress (DE3) (Agilent Technologies, Santa Clara, CA, USA) cells. In BL21(DE3) LOBSTR cells, hDASPO_369 expression was induced as previously reported for the canonical DASPO_341 isoform [18], while the conditions indicated by the supplier were adopted for protein expression in ArcticExpress (DE3) *E. coli* cells. Briefly, cells grown at 30 °C under shaking (OD $_{600\text{nm}} = 0.7$) were added of either 1 mM or 0.1 mM IPTG and harvested after a 24 h of incubation at 13 °C under shaking. The recombinant protein expression levels were evaluated by SDS/PAGE; both the soluble and insoluble fractions of cell lysates were analyzed.

Attempts to purify the soluble His-tagged hDASPO_369 from the crude extracts were carried out using a HiTrap chelating column (GE Healthcare, Chicago, IL, USA) as reported in Ref. [18].

Refolding and purification of insoluble hDASPO_369

After cell lysis and centrifugation, the insoluble fraction of the E. coli cell extracts was recovered and washed twice in 20 mm Tris/HCl pH 8.0, 2 m urea, 0.5 m NaCl, 2% (v/v) Triton X-100. Different refolding procedures were used: (a) the column-refolding of inclusion bodies solubilized in 20 mм Tris/HCl pH 8.0, 6 м guanidine hydrochloride, 0.5 M NaCl, 5 mm imidazole, and 1 mm 2-mercaptoethanol, loaded on a HiTrap Chelating HP column, refolded using a linear gradient from 6 to 0 M of urea and from 0 to 40 µM FAD and then eluted with 20 mm Tris/HCl pH 8.0, 0.5 m NaCl, at increasing concentration of imidazole (100, 250 and 500 mm); (b) the solubilization of inclusion bodies as above, followed by dialysis to remove the chaotropic agent; (c) the solubilization of inclusion bodies in 50 mm Tris/HCl pH 8.0, 2 m urea, and 6 m n-propanol followed by a refolding step by dialysis to remove urea and n-propanol. At the end of the procedures, samples were centrifuged (at 39 000 g for 30 min at 4 °C) and the soluble and insoluble fractions were analyzed by SDS/PAGE.

Cell culture and transfection

The U87 human glioblastoma cells (ATCC) were maintained in Dulbecco's modified Eagle medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 1 mm

sodium pyruvate, 2 mm L-glutamine, 2.5 µg·mL⁻¹ amphotericin B, and 1% penicillin/streptomycin (all from Euroclone, pero, Italy) at 37 °C in a 5% CO2 incubator and transfected using the FuGENE HD transfection reagent (6 μL; Roche, basel Switzerland) and 2 μg of pcDNA3 3X-FLAG-hDASPO 341 or pcDNA3 3XFLAG-hDASPO 369 constructs. Stable clones were selected adding 0.4 mg⋅mL⁻¹ G418 to the growth medium. To perform ubiquitination studies, U87 3XFLAG-hDASPO_341 or U87 3XFLAGhDASPO 369 cells were transiently transfected with 2 µg of the pCMV-FLAG-ubiquitin vector. For immunolocalization studies, 10⁴ U87 3XFLAG-hDASPO 341, 3XFLAGhDASPO 369 cells, and U87 control cells were seeded onto previously gelatinized coverslips (diameter 12 mm, Thermo Scientific, Walthman, MA, USA). At 24 h after seeding, cells were extensively washed with PBS (10 mm dibasic sodium phosphate, 2 mm monobasic potassium phosphate, 137 mm NaCl, 2.7 mm KCl, pH 7.4) and fixed with 4% pformaldehyde for 10 min at room temperature.

To evaluate the stability of hDASPO_341 and _369 isoforms and assess their rate of degradation, U87 3XFLAG-hDASPO_341 and U87 3XFLAG-hDASPO_369 cells were seeded into 6-well plates (2.7 × 10⁵ cells per well) and treated up to 32 h with cycloheximide (CHX, 100 μg·mL⁻¹; Sigma), a potent inhibitor of protein synthesis, and collected at different times for western blot analysis [21]. Changes in hDASPO isoforms cellular levels were determined by densitometric analysis upon acquisition with an Odyssey Fc Imaging System (LI-COR Biosciences, Lincoln, NE, USA). Data were fit to a single exponential decay equation to estimate the half-life.

Processes involved in hDASPO degradation were investigated by treating the cell clones expressing the two protein isoforms with specific inhibitors of autophagy/lysosomal pathway or the ubiquitin–proteasomal system (UPS) [21]. After seeding, the cells were grown overnight under starvation conditions (DMEM supplemented with 1% FBS), and then, the lysosomal hydrolase inhibitors chloroquine (CQ, 75 μμ; Sigma) or NH₄Cl (10 mμ; Sigma, St. Louis, MO, USA) or the proteasomal inhibitor benzyloxycarbonyl-Lleucyl-L-leucyl-L-leucinal (MG132, 25 μμ; Sigma) was added. At different time points (up to 10 h), cells were collected by trypsinization and the hDASPO protein levels were analyzed by western blot. Starvation conditions have been used to maximize the effect of inhibition of the degradation pathways, as previously reported [21,36].

Immunoblot

For western blot analysis, stably transfected or control cells were resuspended in sample buffer (12.5 mm Tris/HCl, pH 6.7, 3% SDS, 5% glycerol, and 62.5 mm dithiothreitol) to have 5000 cells μ L⁻¹ and 10 μ L of each sample was subjected to SDS/PAGE. Proteins were then transferred to polyvinylidene difluoride membranes (Immobilon-P,

Millipore, Burlington, MA, USA) and saturation of aspecific sites was performed by incubation (2 h at room temperature or overnight at 4 °C) in a blocking solution containing 4% dried milk in Tris-buffered saline (TBS; 10 mm Tris/HCl pH 8.0, 0.5 m NaCl) with the addition of 0.1% Tween-20 (TBST). Membranes were then incubated with primary antibodies at room temperature for 1.5 h, extensively washed in TBST, and then incubated for 1 h at room temperature with specific peroxidase-conjugated immunoglobulins. The immunoreactivity signals were detected by enhanced chemiluminescence (WESTAR ETA C Ultra 2.0 reagents; Cyanagen, Bologna, Italy) using the Odyssey Fc Imaging System (LI-COR Biosciences).

The primary antibodies used were as follows: rabbit polyclonal anti-hDASPO (1:1000, Davids Biotechnologie, Regensburg, Germany); rabbit polyclonal anti-FLAG (1:500, Sigma); mouse monoclonal anti-β-tubulin (1:2000, Thermo Fisher Scientific, Waltham, MA, USA).

The different immunorecognition signals in cell lysates were measured using the Image Studio Lite Software (LI-COR Biosciences). The intensity values of the bands detected by the anti-hDASPO antibody were normalized to the values of the anti- β -tubulin ones. After normalization, the ratio of the intensity signals corresponding to treated and control cells, collected at the same incubation time, was calculated.

Immunostaining and confocal analysis

In order to analyze the subcellular localization of the hDASPO isoforms, p-formaldehyde-fixed U87 3XFLAGhDASPO 341, 3XFLAG-hDASPO 369, and U87 control cells were permeabilized and the unspecific binding sites were blocked by incubation in PBS supplemented with 0.2% Triton X-100 and 4% horse serum. DASPO isoforms were subsequently stained using rabbit polyclonal anti-hDASPO (1:500, Davids Biotechnologie) and mouse monoclonal anti-FLAG antibody (1:500, Invitrogen, Carlsbad, CA, USA); peroxisomes and mitochondria were stained by rabbit polyclonal anti-PMP70 (peroxisomal membrane protein 70, 1:500, Sigma) and mouse monoclonal antimitochondria antibody (1:500, Millipore). Cells were incubated with primary antibodies overnight at 4 °C and, after extensive washing in PBS supplemented with 1% horse serum, with antirabbit Alexa 488 and anti-mouse Alexa 546 antibodies (1:1000, Molecular Probes, Invitrogen) diluted in PBS, 0.1% Triton X-100, and 1.5% horse serum.

Immunostained coverslips were imaged using an inverted laser scanning confocal microscope (TCS SP5, Leica Microsystems, Wetzlar, Germany), equipped with a 63.0×1.25 NA plan apochromatic oil immersion objective. Confocal image stacks (five sections with optimized thickness) were acquired using the LEICA TCS software (Leica Microsystems) with a sequential mode to avoid interference between each channel due to spectral overlap and without saturating any pixel.

Immunoprecipitation

determine the specific activity of 3XFLAGhDASPO 341 and 3XFLAG-hDASPO 369, the enzymes were purified from cell extracts by immunoprecipitation (IP) using anti-FLAG M2 Affinity resin (Sigma). Briefly, U87 cell clones stably expressing the two isoforms were suspended in ice-cold lysis buffer (50 mm sodium phosphate pH 8.0, 0.7 μg·mL⁻¹ pepstatin, 1 μg·mL⁻¹ leupeptin, 5 μм FAD, 0.1% ethanol, 1 mg·mL⁻¹ DNAse) and sonicated. The cell lysates were then centrifuged at 13 000 g for 30 min at 4 °C; the protein concentration of the supernatant (Pre-IP sample) was quantified using the Bradford reagent (Sigma). A volume of sample corresponding to 0.5 mg (or 2 mg for U87 3XFLAG-hDASPO 369 cells) of total protein was subjected to IP using 40 µL of anti-FLAG M2 Affinity resin and incubated overnight at 4 °C under constant rotation. The sample was then centrifuged at 8000 g for 1 min, and the supernatant (Post-IP sample) was stored for western blot analysis. The pelleted resin and the bound flagged protein were resuspended in 50 µL of lysis buffer for the subsequent activity measurements (IP sample).

To confirm the identity of the protein forms expressed in transfected with the pcDNA3 3XFLAGhDASPO 369 vector, after a first IP using the whole cell extract and the anti-FLAG M2 Affinity resin, a second IP was performed using the anti-hDASPO antibody. In details, after the removal of flagged proteins, the obtained post-IP sample was incubated with 50 µL of Dynabeads Protein G (Invitrogen) previously cross-linked to 10 µg of rabbit anti-hDASPO antibody (Davids Biotechnologie) using dimethyl pimelimidate (DMP, Thermo Fisher Scientific), and the procedure previously reported [21]. After an overnight incubation at 4 °C with rotation, the supernatant (post-IP sample) was collected by separating the beads on the magnet; beads were extensively washed with lysis buffer, suspended in 50 µL of nonreducing SDS/PAGE sample buffer (IP sample), and boiled. SDS/PAGE and western blot analyses were performed using an amount of pre-IP and post-IP samples corresponding to 20 µg of total protein and 10 μL of the IP samples.

DASPO activity assay and kinetic measurements

hDASPO activity was assayed on total cell lysates and on hDASPO isoforms purified by IP using the Amplex Ultra-Red Assay Kit (Invitrogen) based on the detection of H_2O_2 by the peroxidase-mediated oxidation of the fluorogenic Amplex UltraRed Dye [20]. Cells were suspended in ice-cold 50 mm sodium phosphate buffer, pH 8.0, containing 0.7 μ g·mL⁻¹ pepstatin, 1 μ g·mL⁻¹ leupeptin, 5 μ m FAD, 0.1% ethanol, and 1 μ g·mL⁻¹ DNAse, sonicated, and centrifuged at 13 000 μ g for 30 min at 4 °C. The protein concentration in the supernatant was quantified using the Bradford assay (Sigma). The IP samples were obtained as

reported above. A volume of cell extract corresponding to 2.5 and 5 µg of total protein content or to 2 and 3 µL of the resuspended IP samples was aliquoted in black 96-well plates and added to lysis buffer to a final volume of 100 μL. Then, 50 μL of the activity assay solution (55 μм Amplex UltraRed, 0.15 U·mL⁻¹ horseradish peroxidase, 7.5 mm NaN₃, 50 mm D-Asp, and 15 µm FAD in 50 mm sodium phosphate buffer, pH 8.0) was added to each well. The fluorescence measurements (excitation at 535 nm and emission at 590 nm) were recorded at 25 °C every 5 min for a total of 30 min, and values were corrected for control wells (without the samples) to subtract background signal. As a positive control, different amounts of the recombinant purified enzyme (0.0025-0.1 milliunit range) were assayed in parallel. The assay specificity was assessed using lysates of control U87 cells stably transfected with the empty pcDNA3 vector, as a negative control, and adding 2 mm 5aminonicotinic acid (an inhibitor of hDASPO, $K_i = 3.8 \mu M$) [37] to hDASPO expressing cells.

The concentration of H_2O_2 in the wells was determined using a calibration curve (0.25–7.5 μ M H_2O_2). hDASPO-specific activity (μ mol·min⁻¹·mg⁻¹) was then calculated by dividing the nanomoles of H_2O_2 produced in a minute by the milligrams of enzyme present in the samples, determined by western blot and densitometric analysis, using a calibration curve prepared by analyzing known amounts of the recombinant purified hDASPO.

The apparent kinetic parameters of hDASPO isoforms on p-Asp were determined using the same assay, after the purification of the isoforms by IP, and were calculated according to a Michaelis-Menten equation using the initial velocity values determined at increasing substrate concentrations (0.4–16 mm range). As a positive control, a fixed amount of the recombinant hDASPO (corresponding to 0.07 mU) was assayed.

In vitro ubiquitination

Recombinant hDASPO was expressed in E. coli BL21 (DE3) LOBSTR host cells (Novagen, Merck KGaA) and purified as reported in Ref. [18]. In vitro ubiquitination experiments were carried out using 2 and 5 µg of hDASPO, incubated with 0.6 mg·mL⁻¹ ubiquitin, 1 μM ubiquitin aldehyde, 5 mm ATP, 25 um MG132, 1 mm dithiothreitol, 2 mм MgCl₂, and 4 mg·mL⁻¹ U87 cell extract in 20 mм Tris/HCl (pH 8.0), in a total volume of 60 µL, at 37 °C for 60 min. Controls were performed by omitting ubiquitin or hDASPO in the mixture. The ubiquitin-hDASPO conjugates were resolved by SDS/PAGE and analyzed by western blot using rabbit anti-hDASPO antibody. For preparation of U87 cell extract, the cells were collected, resuspended (10⁷ cells·mL⁻¹) in lysis buffer (20 mm Tris/ HCl pH 8.0, 1 mm dithiothreitol, 5 mm KCl, 2 mm MgCl₂, 2 μM leupeptin, 1 μM pepstatin, 50 μM phenylmethanesulfonyl fluoride, 25 µm MG132), and subjected to sonication

(three cycles of 10 s each). The cell lysate was cleared by centrifugation (see above).

Before and after *in vitro* modification, recombinant hDASPO activity was assayed in 100 mm disodium pyrophosphate buffer (pH 8.3), 40 µm FAD using 15 mm D-Asp as substrate, at 25 °C and air saturation, measuring oxygen consumption by the Clark electrode [18].

In vivo ubiquitination

To evaluate whether hDASPO undergoes ubiquitination in the selected cell line, U87 3XFLAG-hDASPO 341 or U87 3XFLAG-hDASPO 369 cells were transiently transfected with FLAG-ubiquitin, and, 48 h after transfection, the cells were treated for 6 h with 25 µM MG132. Cells were subsequently collected by trypsinization, resuspended in denaturing lysis buffer (20 mm Tris/HCl, pH 8.0, 150 mm NaCl, 1% SDS, and 1 mm EDTA, supplemented with protease inhibitors as above), boiled for 7 min, and sonicated. The cell lysate was then centrifuged at 13 000 g for 30 min at 4 °C, and the supernatant was diluted 1:10 in IP buffer (20 mm Tris/HCl, pH 8.0, 150 mm NaCl, 0.5% NP-40, 0.5% Triton X-100, and 1 mm EDTA, supplemented with protease inhibitors). The FLAG-tagged hDASPO isoforms and ubiquitin were co-immunoprecipitated using anti-FLAG M2 Affinity Gel (Sigma). A volume of sample corresponding to 0.5 or 1 mg of total proteins for U87 3XFLAG-hDASPO 341 and U87 3XFLAG-hDASPO 369 cells, respectively, was added to 50 µL of the anti-FLAG agarose resin, and immunoprecipitation was performed as detailed above. The resin was resuspended in 50 µL of a nonreducing SDS/PAGE sample buffer (IP sample) and boiled. 20 µL of the IP samples was subjected to SDS/ PAGE on an 8-15% polyacrylamide gel and the ubiquitinhDASPO conjugates revealed by western blot analysis using the anti-hDASPO antibody (Davids Biotechnologie).

HPLC analysis

To determine cellular D- and L-Asp levels, as well as D- and L-Ser, the cells stably expressing the 3XFLAGhDASPO 341 variant were analyzed using the procedure reported in Ref. [15] with minor modifications. Cell pellets were homogenized in 1:5 (w/v) 0.2 M TCA and sonicated (three cycles, 10 s each), and the resulting cell extracts were clarified by centrifugation at 13 000 g for 20 min. The precipitated protein pellets were stored at -80 °C for quantification, while 10 µL of the supernatants was neutralized with NaOH and subjected to precolumn derivatization with 20 µL of 74.5 mm o-phthaldialdehyde (OPA) and 30.5 mm N-acetyl L-cysteine (NAC) in 50% methanol. Diastereoisomer derivatives were resolved on a Symmetry C8 reversedphase column (5 μ m, 4.6 \times 250 mm, Waters) under isocratic conditions (0.1 M sodium acetate buffer, pH 6.2, 1% tetrahydrofuran, and 1 mL·min⁻¹ flow rate). A washing step in 0.1 M sodium acetate buffer, 3% tetrahydrofuran, and 47% acetonitrile was performed after each run. Identification of peaks was based on retention times and confirmed by: (a) adding known amounts of external standards to the samples, (b) by the selective degradation catalyzed by the M213R or wild-type RgDAAO for D-Asp and D-Ser, respectively: 10 µg of the enzymes was added to the samples, which were incubated at 30 °C for 60 min and then derivatized. The peak area for D-Asp or D-Ser corresponded to the one decreased following the enzymatic treatment. Total protein content of homogenates was determined using the Bradford assay method after resolubilization of the TCA precipitated protein pellets in 1% SDS. The total amount of D- and L-amino acids detected in cell extracts was normalized by the total protein content.

nLC-MS/MS analysis

To investigate the presence of hDASPO isoforms in hippocampal cortex from male and female healthy controls and AD patients, samples were analyzed by MS using a shotgun label-free proteomic approach. Frozen hippocampus cortex samples were obtained from London Neurodegenerative Diseases Brain Bank (London. UK). Hippocampus tissues were homogenized using a Potter homogenizer in 200 µL of extraction buffer (8 m urea, 20 mm HEPES pH 8, with proteases, and phosphatase inhibitors) at full speed for 1 min. The homogenate was sonicated 3 times for 20 ms on ice with a ultrasonic homogenizer and centrifuged at 13 000 g for 15 min to sediment unhomogenized tissue and large cellular debris. The pellet was discarded, and the protein content was determined by a Bradford assay (Sigma). Proteins were subjected to reduction with 13 mm dithioerythritol (30 min at 55 °C) and alkylation with 26 mm iodoacetamide (IAA; 30 min at room temperature). Peptide digestion was conducted using sequence-grade trypsin (Promega, Madison, WI, USA) for 16 h at 37 °C using a protein: trypsin ratio of 20:1 [38]. The proteolytic digest was desalted using Zip-Tip C18 (Millipore) before MS analysis. LC-ESI-MS/ MS analysis was performed on a Dionex UltiMate 3000 directly connected to an Orbitrap Fusion Tribrid Mass Spectrometer (Thermo Fisher Scientific) by a nanoelectrospray ion source. Peptide mixtures were enriched on 75 µm ID × 150 mm Acclaim PepMap RSLC C18 column and separated employing the following LC gradient: 4% ACN in 0.1% formic acid for 3 min, 4-28% ACN in 0.1% formic acid for 130 min, 28-40% ACN in 0.1% formic acid for 20 min, 40-95% ACN in 0.1% formic for 2 min, and 95-4% ACN in 0.1% formic acid for 3 min at a flow rate of 0.3 μL·min⁻¹. MS spectra of eluting peptides were collected over an m/z range of 375–1500 using a resolution setting of 120 000, operating in the data-dependent mode to automatically alternate between Orbitrap-MS and Orbitrap-MS/MS acquisition. HCD MS/MS spectra were collected for the 20 most abundant ions in each MS scan using a normalized collision energy of 30%, and an isolation window of 1.7 m/z. Rejection of +1 and unassigned charge states were enabled [39]. Raw label-free MS/MS files from Thermo Xcalibur software (version 4.1) [40] were analyzed using Proteome Discoverer software (version 1.4, Thermo Fisher Scientific) and searched with the Sequest algorithm against the hDASPO from UniProt 05-11-2020. Only peptides with high confidence and a high cross-correlation score (\geq 1.5) were considered. The minimum required peptide length was set to 6 amino acids with carbamidomethylation as fixed modification, Met oxidation, and Arg/Gln deamidation as variable modifications.

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Conflict of interest

The authors declare no conflict of interest.

Author contributions

VR conceived, designed, and performed the experiments under the supervision of SS. GT and EM performed MS analysis. LP conceived the work, contributed to the analysis and discussion of the results, and critically revised the manuscript. All authors have read and agreed to the published version of the manuscript.

Peer Review

The peer review history for this article is available at https://publons.com/publon/10.1111/febs.15797.

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Fig. S1. Multiple sequence alignment of hDASPO isoforms.

Table S1. Conditions used to solubilize recombinant hDASPO 369 from *E. coli* inclusion bodies.

Table S2. Bioinformatics prediction of protein solubility for hDASPO isoforms.

Table S3. Primers used for cloning human hDASPO_341 and hDASPO_369 cDNA in the pcDNA3 mammalian expression vector.