

Figure 1

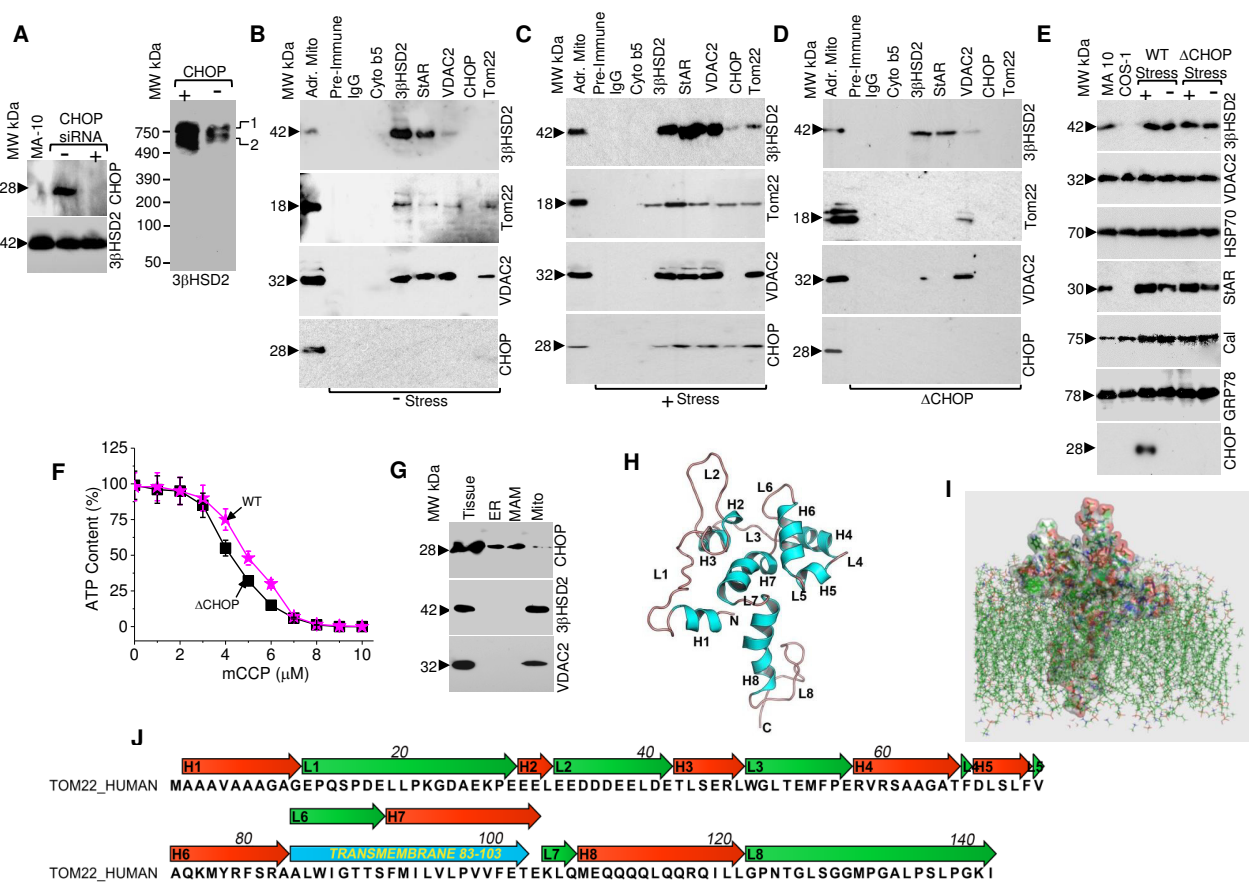


Figure 2

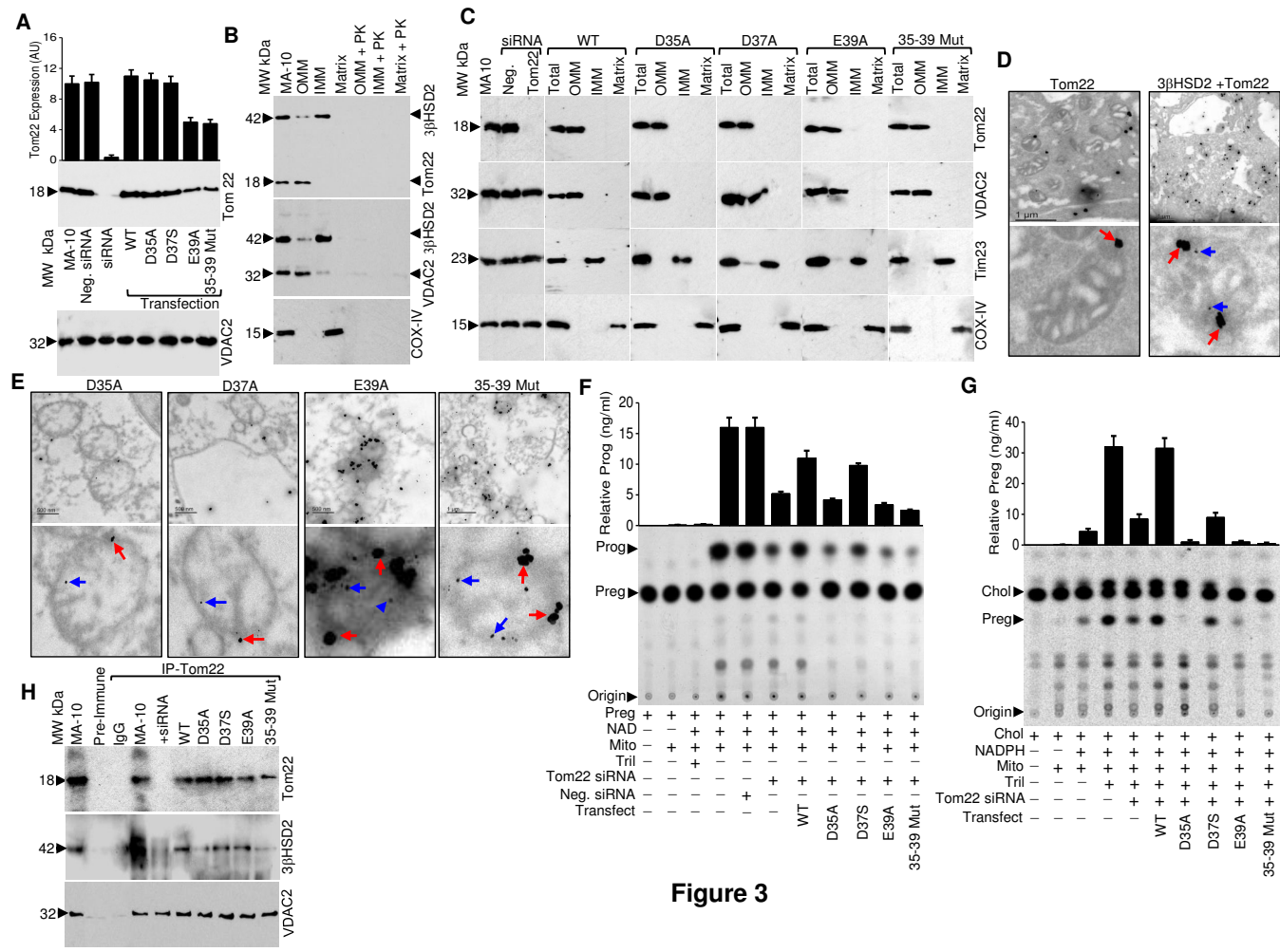


Figure 3

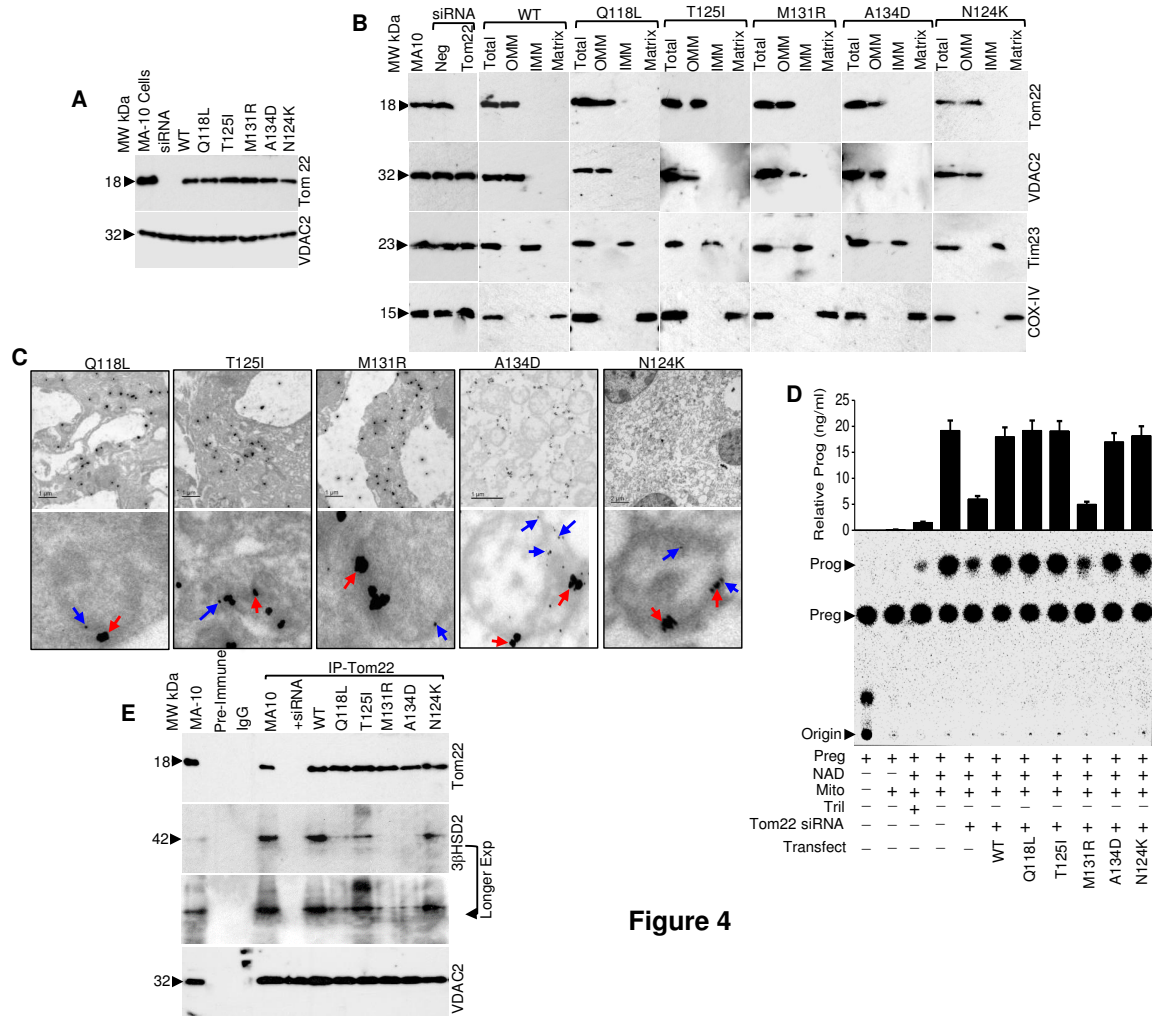


Figure 4

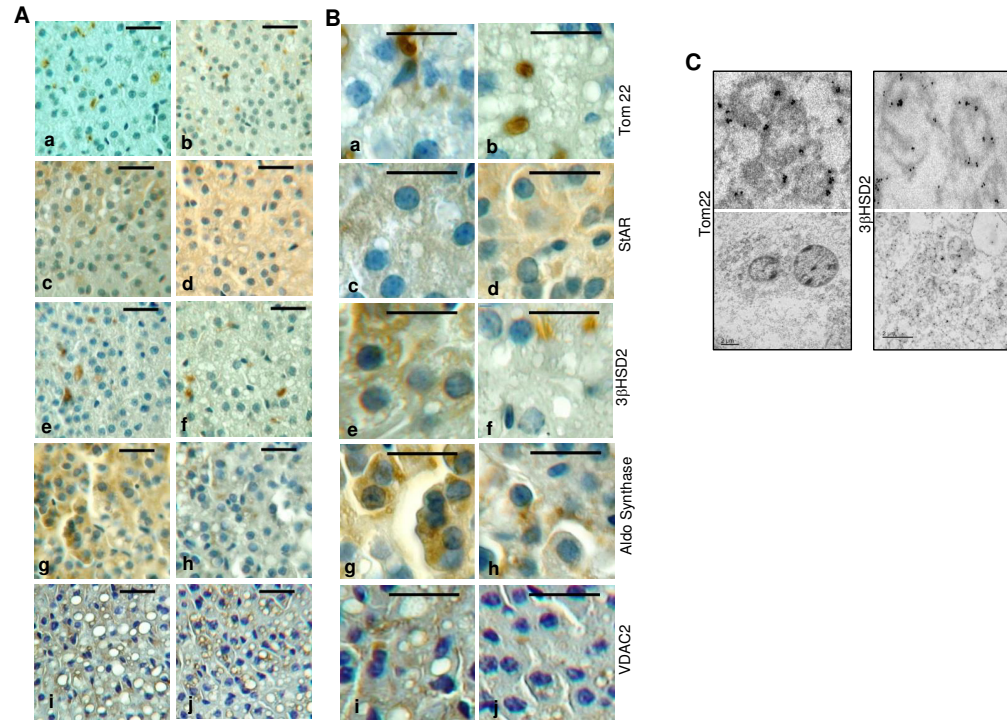


Figure 5

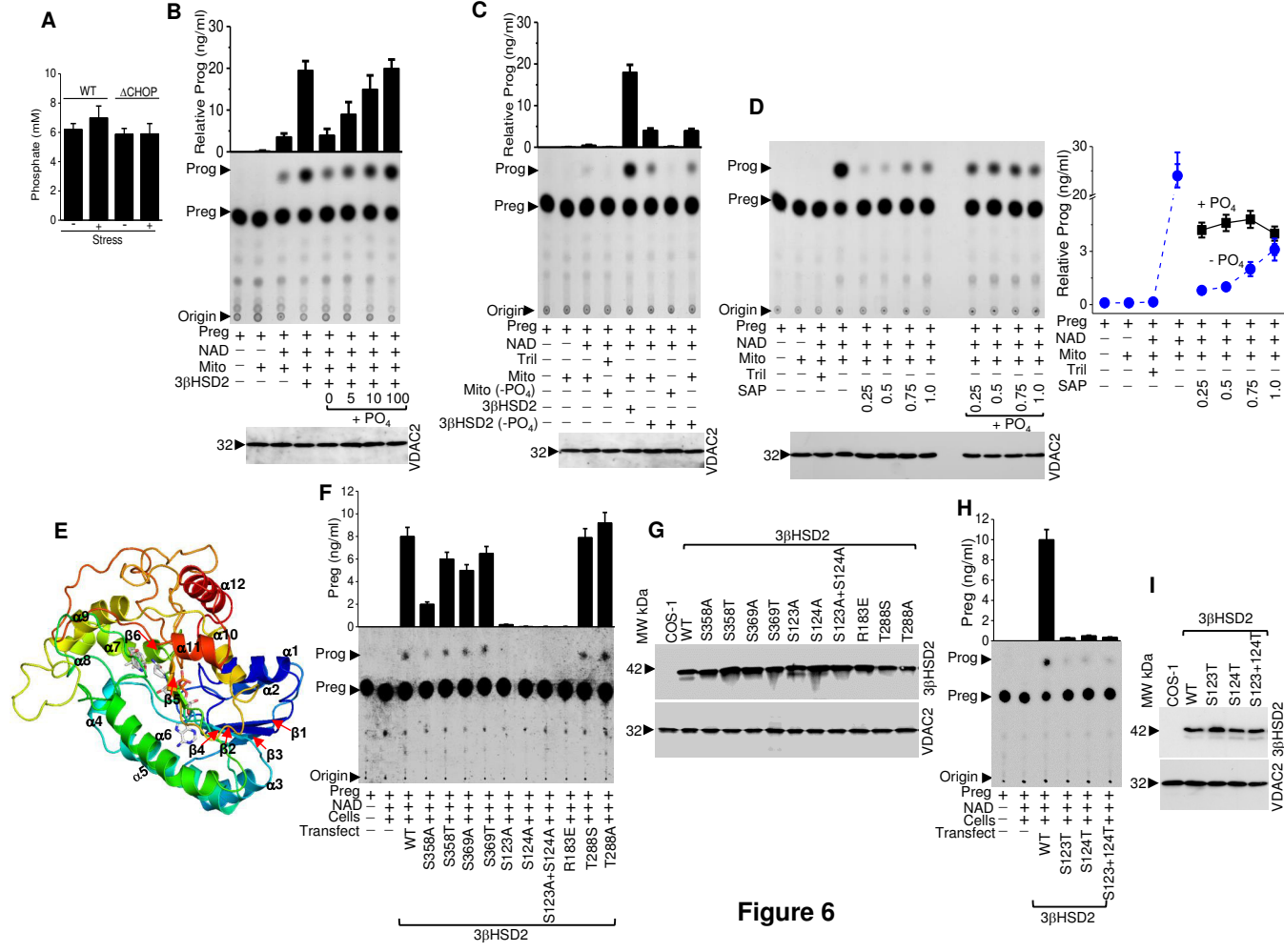


Figure 6

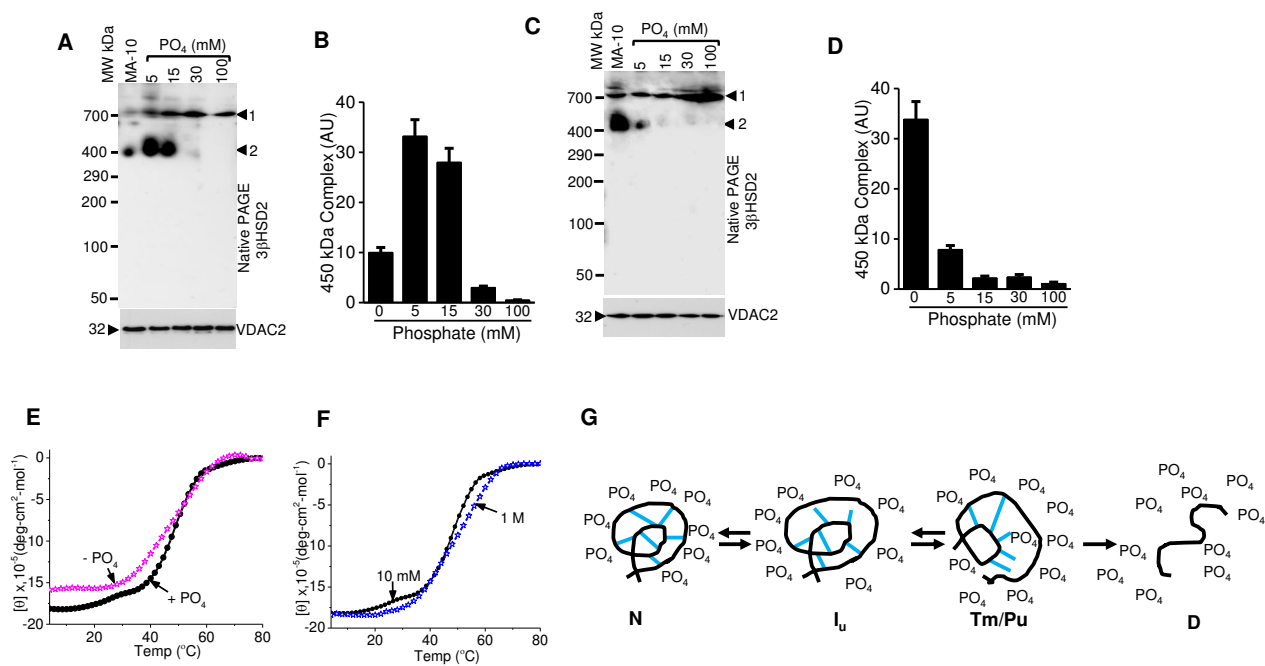


Figure 7

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Endoplasmic reticulum stress enhances mitochondrial metabolic activity in mammalian adrenals and gonads

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Running Title: Stress facilitates mitochondrial metabolism

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Keywords: 3 β -hydroxysteroid dehydrogenase type 2 (3 β HSD2), Translocase of Outer Mitochondrial Membrane 22 (Tom22), Steroidogenic Acute Regulatory protein (StAR), Pregnenolone, Progesterone, CHOP (C/EBP homology protein), Unfolded Protein Response (UPR), ER stress, Mitochondria.

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27 **ABSTRACT**

28 The acute response to stress consists of a series of physiological programs to promote survival by generating
29 glucocorticoids and activating stress-response genes that increase the synthesis of many chaperone proteins
30 specific to individual organelles. In the endoplasmic reticulum (ER), short-term stress triggers activation of the
31 unfolded protein response (UPR) module that either leads to neutralization of the initial stress or adaption to it;
32 chronic stress favors cell death. UPR induces expression of the transcription factor, C/EBP homology protein
33 (CHOP), and its deletion protects against the lethal consequences of prolonged UPR. Here, we show that stress-
34 induced CHOP expression coincides with increased metabolic activity. During stress, the ER and mitochondria
35 comes close to each other, resulting in the formation of a complex consisting of the mitochondrial translocase,
36 translocase of outer mitochondrial membrane 22 (Tom22), steroidogenic acute regulatory protein (StAR) and
37 3 β -hydroxysteroid dehydrogenase type 2 (3 β HSD2) via its intermembrane space (IMS)-exposed charged
38 unstructured loop region. Stress increased the circulation of phosphates, which elevated pregnenolone synthesis
39 by 2-fold by increasing the stability of 3 β HSD2 and its association with the mitochondrial-associated ER
40 membrane (MAM) and mitochondrial proteins. In summary, cytoplasmic CHOP plays a central role in
41 coordinating the interaction of MAM proteins with the OMM translocase, Tom22, to activate metabolic activity
42 in the IMS by enhanced phosphate circulation.

43

44 INTRODUCTION

45 The acute response to stress consists of a relatively stereotyped series of physiological programs to promote
46 survival. It is mediated by the hypothalamic-pituitary-adrenal (HPA) axis, resulting in hypothalamic
47 corticotropin releasing hormone (CRH) secretion that stimulates release of adrenocorticotrophic hormone
48 (ACTH) by the pituitary and subsequent adrenal glucocorticoid release. Stress signaling is a critical factor
49 regulating major morphological changes in cells that may be dependent on the activation of Ca^{2+} -dependent
50 protein kinase C (PKC) (1, 2). Mitochondrial stress may also induce retrograde signaling in mammalian cells
51 (3). In an animal model of acute stress in which mice were exposed to temperature changes, improved
52 physiological recovery, reduced mortality and hormonal changes were observed. Such stress also leads to
53 transcriptional activation of genes that harbor stress response elements within their promoters. For example,
54 heat shock elements (HSEs) are found in the promoters of genes encoding proteins representative of all
55 subcellular compartments (4), enabling cells to respond to global stress by increased synthesis of heat shock
56 proteins and other molecular chaperones (5). Cells can also respond to stress in a way that is specific to
57 individual organelles. Specifically, the endoplasmic reticulum (ER) stress response or the unfolded protein
58 response (UPR) is activated in response to mild or short-term stress triggers, inducing the expression of a wide
59 range of genes involved in the maintenance of ER function (6). In contrast, severe or long-lasting stress favors
60 activation of a proapoptotic module that will lead to cell death.

61 Abnormal protein conformation disturbs cellular homeostasis and is considered a cause of many diseases,
62 including developmental abnormalities. Signal transduction cascades are activated to restore the ER to its
63 normal physiological state. The most abundant ER chaperone, the 78-kilodalton glucose-regulated protein
64 (GRP78/BiP), is responsible for maintaining the permeability barrier of the ER during protein translocation,
65 guiding protein folding and assembly, and targeting misfolded proteins for degradation (7). In unstressed cells,
66 a fraction of ER-luminal GRP78 is bound to three different ER transmembrane proteins: (i) inositol-requiring
67 kinase/endoribonuclease 1 (IRE1), (ii) a protein kinase activated by double-stranded RNA, (PKR)-like ER
68 kinase (PERK), and (iii) activating transcription factor 6 (ATF6) (7). Binding of GRP78 to the ER-luminal
69 domains of these proteins maintains them in an inactive state. Upon ER stress and concomitant accumulation of

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70 misfolded and unprocessed proteins, GRP78 is sequestered away from PERK, IRE1, and ATF6 in order to
71 attend to the increased need for protein folding (7).

72 The CHOP gene encoding the bZIP transcription factor, CHOP [C/EBP homology protein, also called
73 GADD 153], is unregulated by JNK2 kinase and activator protein-1 (AP-1) (8) in response to the UPR (9).
74 Studies using CHOP-null mice have established its role in ER-stress-induced apoptosis; CHOP deletion
75 protects against the lethal consequences of prolonged UPR (10). Because both cells and animals lacking CHOP
76 are protected against different physiological problems, CHOP may have a role in different cellular functions,
77 possibly impacting both viability and apoptosis (11). However, it is not clear whether CHOP is directly
78 inducing apoptosis or whether cell dysfunction and death arise as a secondary consequence of CHOP activity.
79 CHOP likely has a protective role in maintaining ER function, and CHOP expression in response to stress
80 impacts mitochondrial biogenesis through its chaperone activity (10).

81 Impaired stress response arises from either primary defects in the adrenal gland or secondary hypothalamic
82 or pituitary defects, resulting in steroidogenesis defects. Mitochondrial proteins, enzymes and translocases
83 responsible for steroid synthesis are encoded by the nucleus, synthesized in cytosol, and then targeted to
84 mitochondria (Fig. 1A) (12-15). Outer and inner mitochondrial membrane (OMM and IMM, respectively)
85 translocator protein assemblies, including TOM (Translocase, Outer Membrane) and TIM (Translocase, Inner
86 Membrane), translocate and sort proteins into mitochondria. For example, on acute stress or hormonal
87 stimulation, steroidogenic acute regulatory protein (StAR) is synthesized in the cytoplasm. It first interacts with
88 the OMM-associated voltage-dependent anion channel 2 (VDAC2) followed by VDAC1, which is necessary
89 for it to achieve an active conformation to foster cholesterol from the OMM to the IMM (16). Next, 3-beta-
90 hydroxysteroid dehydrogenase type-2 (3 β HSD2), which catalyzes pregnenolone to progesterone and DHEA to
91 androstenedione conversion, interacts with the IMM translocase, Tim50, from the C-terminal to the inter
92 membrane space (IMS) (17, 18). Given the interaction of Tim50 and Tim23 in the IMS, it was not surprisingly
93 that 3 β HSD2 also formed a transient association with Tim23 and regions of Tom22, generating a larger
94 complex (Fig. 1B) (17). Tom22, an OMM-associated protein facing the cytoplasm, acts as a receptor for protein
95 sorting and translocation of OMM proteins targeted to mitochondria (19). Knockdown of Tim50 expression
96 reduced the expression of 3 β HSD2 by 80% as well as decreased its activity (17). Moreover, knockdown of

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107 either Tim23 or Tom22 reduced the conversion of DHEA to androstenedione by 3 β HSD2 without affecting
108 P450c17 (17), supporting the notion that 3 β HSD2 requires mitochondrial translocases for steroidogenic and
109 metabolic activity. Using an *in vivo* mouse model of stress, we found that the ER and mitochondria come in
110 close proximity to each other, resulting in a complex containing mitochondrial StAR, Tom22 and 3 β HSD2
111 (Fig. 1B).

112 Given its role in stress, we hypothesized that acute stress-induced CHOP expression may increase StAR
113 protein levels and facilitate cholesterol transport (Fig. 1B), resulting in increased pregnenolone synthesis. In
114 this article, we provide the first direct evidence of how adrenal steroid production is preferentially induced by
115 CHOP expression without affecting other mitochondrial or ER proteins. Specifically, CHOP facilitates a supply
116 of phosphates into the mitochondria, resulting in increased steroid synthesis by stabilizing 3 β HSD2 protein
117 conformation and its association with neighboring proteins.

118 MATERIALS AND METHODS

119 *Cell culture, generation of CHOP knockdown cells (Δ CHOP cells) and Construction of plasmids*

120 The mouse Leydig tumor cell line (MA-10) was grown in Waymouth media containing 15% horse serum, 5%
121 fetal bovine serum and 1 \times Gentamycin. Cells were maintained at 37°C in a humidified incubator under 5%
122 CO₂. The pSilencer siRNA expression plasmid for CHOP was obtained from Open Biosystems (20). For
123 generation of stable Δ CHOP MA-10 cells, the cells were transfected with the purified plasmid using
124 Oligofectamine (Invitrogen, Carlsbad, CA, USA), and single clones were generated 48 h after transfection by
125 limiting dilution into a selection medium containing 600 mg/mL of G418 (Geneticin; Life Technology, CA).
126 Individual clones were then transferred to 24-well plates for propagation and then later transferred to 6-cm
127 plates for transfection. Individual clones were examined for CHOP expression by Western blotting.

128 Construction of the full-length Tom22 and 3 β HSD2 or different mutants cDNA expression cDNA
129 vectors was described before (16). The accuracy of all clones was confirmed by sequencing in both direction
130 through a commercial resource (MC Cloning lab, South San Francisco, CA). siRNAs for Tom22 were obtained
131 from Life Technologies. COS-1 or MA-10 cells were transfected with 30 pmol siRNA (17), respectively, using
132 oligofectamine (Life Technology, CA). Non-targeting siRNA as well as a combination of two non-targeting
133 scrambled siRNAs (Life Technology) were included as controls in all experiments. In a specific experiment, we

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124 also incubated with 100 ng/ml Trilostane (Gift from Dr. Gavin Vinson, University of London) as an inhibitor
125 for 3 β HSD2. The accuracy of the knockdown was determined by Western blotting.

126 *Animal model*

127 Male wild-type (WT) and CHOP-null mice were purchased from Jackson Laboratories (Bar Harbor
128 Maine, USA). All mice were maintained in a pathogen-free facility and placed on a standard chow diet
129 (AIN93G, Harlan Teklad Global Diets) for 4 weeks. The experimental procedure was approved by the
130 Institutional Animal Care and Use Committee (IACUC number# A1406011 on September 8, 2014). To induce
131 acute stress, animals were randomly divided into two groups: (i) control animals exposed ambient air at room
132 temperature and (ii) experimental animals exposed to 10°C for 1h in each day for 14 days. There were six
133 animals in each group and each experiment was repeated three times independently. All animals were sacrificed
134 immediately after their experimental procedure at day 14. Blood samples were collected through a heart
135 puncture at the time they were sacrificed, and the adrenal and testicular tissues were excised, rinsed in ice-cold
136 buffer, and processed for metabolic conversion immediately.

137 *Isolation, purification and fractionation of mitochondria*

138 Mitochondria were isolated from cells or mouse adrenal tissues. For mitochondrial isolation, adrenal
139 tissues were diced in mitochondrial isolation buffer (250 mM sucrose, 10 mM HEPES, 1 mM EGTA, pH 7.4);
140 MA-10 or COS-1 cells were washed with PBS two times and then isolated mitochondria following our
141 previously developed procedure (17, 21). For most of the experiments, fresh mitochondria were used
142 immediately after their isolation from tissues or cells.

143 The mitochondrial compartments were individually purified following a standard procedure with minor
144 modifications (22). In brief, the OMM fraction was extracted with 1.2% digitonin, and the remaining mixture of
145 matrix and IMM was purified through 0.5% Lubrol, where the matrix fraction remained in solution after
146 centrifugation, and the IMM fraction formed the pellet. The purity of mitochondrial fractions was confirmed by
147 assessing the levels of compartment-specific proteins.

148 *Native polyacrylamide gel electrophoresis (PAGE)*

149 To analyze complex formation, mitochondria were isolated from MA-10 cells and Δ CHOP MA-10
150 cells. The native complex was isolated by incubating the isolated mitochondria with buffer containing 1%

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151 digitonin, and samples were separated by electrophoresis through 3-16% gradient native gels. Next, the protein
152 complexes were transferred to a membrane and then probed with antibodies against 3 β HSD2 and CHOP.

153 *3 β HSD2 protein purification and Conformational analysis*

154 3 β HSD2 was expressed as a recombinant protein in the baculovirus system (23). The concentrations of
155 the unfolded proteins were determined with guanidinium hydrochloride to avoid errors in extinction
156 coefficients (24). Multiple scans were averaged to improve the signal-to-noise ratio. Appropriate buffer
157 baselines were obtained under the same experimental conditions and were subtracted from the sample spectra.
158 Far-UV (195-250 nm) circular dichroism (CD) measurements were carried out in a Jasco spectropolarimeter
159 (JASCO -815, Japan) at 20°C with a 1.0-mm path length cuvette containing 175 mg/mL (1.52×10^{-6} M) protein
160 in 10 mM NaH₂PO₄, pH 7.5. Spectra shown are the averages of three consecutive scans that were performed at
161 a scan speed of 20 nm/min and corrected by subtracting corresponding blanks. Results are presented as mean
162 residue molar ellipticity (Θ): $[\Theta]_r = \theta_{\text{obs}} / (10nlc)$, where θ_{obs} is the measured ellipticity in millidegrees, 'n' the
163 number of residues in the protein, 'l' the path length of the cell expressed in cm, and 'c' the molar
164 concentration of protein. Thermal melting (T_m) studies were performed in temperatures ranging from 4°C to
165 80°C, and the data was collected at 208 nm and 222 nm with a 0.2°C/min rate. For protein stabilization studies,
166 the T_m of 3 β HSD2 with and without shrimp alkaline phosphatase (SAP) treatment was performed with 10 mM
167 NaH₂PO₄ and 10 mM C₂H₃NaO₂.

168 *Proteolytic digestion experiments*

169 Proteolytic digestion experiments were performed at 4°C or room temperature using various
170 concentrations of proteinase K (PK; Sigma, St. Louis, MO). The limited digestion experiments were performed
171 using 12.5 μ g of total protein and different concentrations of PK for either 15 or 45 min. The reactions were
172 terminated by the addition of an equal volume of SDS sample buffer containing 2 mM PMSF and then
173 incubating in a boiling water bath. After electrophoresis, the samples were processed for Western blotting using
174 the indicated antibodies.

175 *Western blot analysis*

176 Protein (12.5 μ g) was separated by 15% SDS-PAGE and transferred to a polyvinylidene difluoride
177 (PVDF) membrane (Millipore, Billerica, MA, USA). The membrane was blocked with 3% nonfat dry milk for

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178 45 min, probed overnight with the primary antibodies, and then incubated with the peroxide-conjugated goat
179 anti-rabbit IgG or anti mouse IgG (Pierce). Signals were developed with a chemiluminescent reagent (Pierce).

180 ***Mitochondrial viability assay***

181 To check the mitochondrial activity and its membrane responsiveness, ATP assays were performed
182 using an ATP Assay System Bioluminescence Detection Kit (ENLITEN, Promega, Madison, WI, USA) with a
183 luminometer (Veritas microplate luminometer, Turner Biosystems) following the manufacturer's protocol.
184 Basic mitochondrial activity was determined in terms of ATP production, and mitochondrial activity was
185 inhibited by incubation of MA-10 cells and Δ CHOP mitochondria with various concentrations of mCCP
186 (Carbonyl cyanide m-chlorophenyl hydrazone) for 1h.

187 ***Metabolic conversion assays***

188 Isolated mitochondria from murine adrenals or steroidogenic MA-10 cells (300 μ g) were incubated in
189 phosphate buffer for metabolic conversion experiments. For the 3 H-pregnenolone to progesterone assay, 3×10^6
190 counts of 3 H-pregnenolone were used for each reaction, which was chased with 30 μ g of cold unlabeled
191 progesterone (25). The metabolic reaction was initiated by addition of NAD and incubated at 37°C with
192 continuous shaking for 4 h. In the case of 14 C-cholesterol to pregnenolone, 80,000 counts of 14 C-cholesterol
193 were used for each reaction and chased with 20 μ g of unlabeled cholesterol. For metabolic conversion of
194 cholesterol to pregnenolone, the reaction was initiated by addition of NADPH and the reaction mixture was
195 incubated at 37°C with continuous shaking for 3 h. To ensure complete conversion, we used 5-fold excess cold
196 carrier in order to reach the saturation point. The steroids were extracted with ether/acetone (9:1 v/v), and equal
197 amounts of a cold pregnenolone-progesterone mixture (for pregnenolone-progesterone assay) and cold
198 cholesterol-pregnenolone mixture (for 14 C-cholesterol to pregnenolone) in CH_2Cl_2 was added as a carrier. The
199 extracts were concentrated under a stream of nitrogen and then separated by TLC (Whatman, Sigma, St. Louis,
200 MO) using a chloroform/ethyl acetate (3:1) mobile phase.

201 ***Gas chromatography-mass spectrometry (GC-MS)***

202 The spots extracted from TLC plates were subjected to GC-MS analysis (Agilent 7890 GC and 5975C
203 mass spectrometer). The column was an Agilent HP-5 with dimensions of 30 $\mu\text{m} \times 0.25 \mu\text{m}$ inner diameter
204 with a 0.25 μm film thickness. Samples were dissolved in 50 μL of CH_2Cl_2 , and 1 μL was injected onto the

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column using a pulsed splitless injection. Helium was used as the carrier gas at a flow rate of 1 mL/min. The temperature program is as follows: ramp at 10°C/min from 70° to 310°C and hold for 6 min. Spectra were collected in full scan with 70 Ev ionization mode over the mass range of m/z 50 to 500 to facilitate comparison of MS spectra.

Co-immunoprecipitation (co-IP) analysis

Specific antibodies were pre-incubated with protein A-Sepharose CL 4B (0.5 µg/µL, Amersham Biosciences, Sweden) in 100 µL of 1× co-IP buffer (1% Triton X-100, 200 mM NaCl and 0.5% sodium deoxycholate) and mixed on an end-over-end rotator for 2 h at 4°C. To remove unbound antibody, the beads were washed with 1× co-IP buffer five times and then incubated again with rabbit IgG control antibody (Sigma) for 1 h to block unbound beads. After another series of washes to remove unbound antibody, the freshly isolated mitochondrial pellet (25 mg for each sample) was resuspended with ice cold lysis buffer (20 mM Tris HCl, pH 8.0, 137 mM NaCl, 10% glycerol, 1% Triton X-100, 2 mM EDTA) and incubated at 4°C for 15 min. Insoluble material was removed by ultracentrifugation (30 min at 100,000 ×g). The supernatants were incubated overnight at 4°C with end-over-end shaking in the presence of antibodies prebound to protein A Sepharose beads. After five to six washes with 1× co-IP buffer and two washes with 10 mM HEPES (pH 7.4), the protein A-Sepharose pellets were resuspended and vortexed with 100 mM Glycine (pH 3.0) for 10 sec. The sample pH was changed to pH 7.4 by adding a pre-titrated volume of 1.0 M Tris (pH 9.5), and the beads were separated from the soluble material by centrifugation at 2000 ×g for 2 min. The supernatants (immune complexes) were analyzed by Western blotting.

The dephosphorylation studies were performed with 10 mM Tris (pH 7.4), where purified mitochondria were washed with 10 mM Tris (pH 7.4) prior to the metabolic conversion assay. 3βHSD2 and mitochondria were phosphate deprived individually by incubating each with 10 mM Tris (pH 7.4) for 1 h at 37°C after which activity was determined. For rephosphorylation studies, various concentrations of 1 mM NaH₂PO₄ (pH 7.4) was added back to phosphate-devoid samples following the removal of the electrolytes through dialysis.

Histological analysis

Adrenal glands were fixed in zinc formalin, processed, and embedded in paraffin for routine light microscopy. Sections of 5-10 µm were obtained from each specimen, floated onto glass slides, stained with

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hematoxylin and eosin (H&E), and examined by microscopy. For immunohistochemistry analysis, additional slides from each specimen were deparaffinized, rehydrated, and stained with either polyclonal antisera (dilution 1:100) or the monoclonal antibodies of the following dilutions: neat, 1:20, 1:50, and 1:100. Binding of the primary antiserum was detected by using the appropriate biotinylated secondary immunoglobulin followed by an avidin-peroxidase conjugate with diaminobenzidine as the chromogen. Negative control sections were treated in identical fashion except for the substitution of nonimmune goat serum for primary antisera or antibody. Sections were counterstained with hematoxylin.

Measurement of serum corticosterone

After completion of stress experiment blood was collected immediately from the experimental animals in an eppendorf tube through cardiac puncture and incubated at room temperature for 30 minutes. The serum was collected by centrifugation at 3000Xg for 10 minutes. To determine corticosterone concentration the serum was further diluted at various concentrations to a final volume of 50 microliter and the corticosterone was measured by radioimmunoassay (RIA) following manufacturer's protocol (Corticosterone ³H RIA kit; MP Biomedicals, CA). The radioactivity was measured in a Scintillation counter (Beckman, CA).

Measurement of serum phosphate

To determine phosphate concentration 1, 2, 5, 10 and 50 microliter of serum was diluted to a final volume of 200 microliter and the phosphate concentration was determined by using Phosphate Assay Kit following manufacturer's protocol (Abcam, CA). The color development was detected in a luminometer (BioTek Synergy HT, CA) at 650 nm and the concentration was determined through an internal standard.

Computational modeling of 3 β HSD2

Amino acid and nucleotide sequences were retrieved from the Swiss Protein Database (26). Next we performed the modeling using the Pymol Molecular Graphics system (Version 1.3 Schrodinger, LLC) (27) as described before (28).

Electron microscopy (EM)

To analyze the subcellular localization of 3 β HSD2 and Tom22 in the mitochondria of both the steroidogenic MA-10 cells, EM experiments were performed. The cells (6×10^6 MA-10 cells) were washed twice with PBS, gently scraped in the presence of PBS and transferred to 50 mL plastic disposable Corning

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259 tubes. After centrifugation at 1470xg (Beckman Allegra 22R and rotor F630) for 10 min, the cells were fixed in
260 4% formaldehyde and 0.2% glutaraldehyde in 0.1M sodium cacodylate buffer, pH 7.4, dehydrated with a
261 graded ethanol series through 95% and embedded in LR white resin. Thin sections of 75 nm thickness were cut
262 with a diamond knife on a Leica EM UC6 ultramicrotome (Leica Microsystems, Bannockburn, IL) and
263 collected on 200 mesh nickel grids. The sections were first blocked in 0.1% BSA in PBS for 4 h at room
264 temperature in a humidified atmosphere followed by incubation with a 1:50 dilution of Tom22 antibodies in
265 0.1% BSA overnight at 4°C. The sections were washed five times with PBS and floated on drops of anti-
266 primary specific ultra small (<1.0 nm) NanogoldTM reagent (Nanoprobes, Yaphank, NY, USA) diluted 1:2000
267 in 0.1% BSA in PBS for 2-4 h at room temperature. After five washes with PBS and five with deionized H₂O,
268 the sections were incubated with HQ SilverTM (Nanoprobes) for 8 min for silver enhancement, followed by
269 washing in deionized H₂O.

270 For double immunolabeling, the same sections were first labeled with a Tom22 antibody (1:2000)
271 overnight at 4°C followed by incubation with 3βHSD2 antibody (1:2000) overnight at 4°C. Because the silver
272 enhancement of the gold particles labeling Tom22 occurred for twice as long as the gold particles labeling
273 3βHSD2, it, therefore, produced two different sizes of gold particles. After a final wash, the grids were stained
274 with 2% uranyl acetate in 70% ethanol to increase the contrast. The grids were washed five times with
275 deionized H₂O and air dried. The large gold particles were an average of 55 nm in diameter with 90% of the
276 gold particles being between 45-65 nm in diameter. The small gold particles were an average size of 15 nm
277 with 90% of the gold particles being smaller than 25 nm in diameter. The cells were observed using a JEM
278 1230 transmission electron microscope (JEOL USA Inc., Peabody, MA, USA) at 110 kV and imaged with an
279 UltraScan 4000 CCD camera and First Light Digital Camera Controller (Gatan Inc., Pleasanton, CA, USA).
280 Forty sections from each experiment were analyzed.

281 ***Computational modeling of Tom22***

282 We developed a 3D structural model of Tom22 (NCBI NP_064628.1, Uniprot Q9NS69) sequence (AA
283 1-142) using the structures of NAD-dependent aldehyde dehydrogenase from *Lactobacillus acidophilus*
284 (3ROS) and Esta from *Arthrobacter nitroguajacolicus* (3ZYT) as templates using the programs YASARA (29)
285 and WHATIF (30). First, we performed a PhiBlast search of the PDB database with the amino acid sequence of

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Tom22 to create a custom position-specific scoring matrix (PSSM) that was then used in further runs of PhiBlast searches to identify structurally similar sequences. A secondary structure prediction of the sequence was used to perform the structure-based alignment of the sequences using the PSI-Pred secondary structure algorithm (31). Aligned sequences were subsequently analyzed by YASARA and WHATIF. Some of the loops were modelled separately by scanning a library of loop databases that predicted that Tom22 should have a minimum amino acid region facing the IMS. Side chains were optimized by molecular dynamic (MD) simulations. The final model was refined by a 500 ps (MD) simulation using AMBER 2003 force field and checked with WHATCHECK (32), WHATIF (30), Verify3D (33, 34) and Ramachandran plot analysis (35, 36). Structures were depicted with Pymol (www.pymol.org) and rendered as ray-traced images with POV-RAY (www.povray.org). The MD simulations were performed using the AMBER03 force field (29). The simulation cell was filled with water, the pH was fixed to 7.4, and the AMBER03 (37) electrostatic potentials were evaluated for water molecules in the simulation cell and adjusted by addition of sodium and chloride ions. The final MD simulations were then run with AMBER03 force field at 298K, 0.9% NaCl and pH 7.4 for 500 ps to refine the models. The best models were selected for analysis and evaluation.

Figure preparation and statistical data analysis

The images were obtained from the autoradiogram or scanning through a phosphorimager and the data analysis was performed using Origin software (OriginLab Corporation, Northampton, MA) or Microsoft Excel. The mean of the data was compared between different animal groups and p-value was significantly concerned at 0.05 levels. The data analysis was performed by hypothesis testing for quantifying group comparisons. The parameters were compared between two specific groups and p-value was determined by Student's *t*-test.

RESULTS

Acute stress increases adrenal steroidogenesis in vivo

Within the lumen of the ER, protein chaperones assist in folding newly synthesized polypeptides and preventing aggregation of unfolded or misfolded proteins (38). The most abundant ER chaperone, GRP78, is responsible for maintaining the permeability barrier of the ER during protein translocation, and associates with the transmembrane ER stress sensors, IRE1-1 and ATF6 (Fig. 1B). Thus, we determined the impact of ER stress on the expression of ER- and mitochondrial-associated ER membrane (MAM)-resident proteins and

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313 identified an effect on mitochondrial metabolic activity. Specifically, ER stress induced by a change in
314 temperature for 1 h/day for 14 continuous days did not alter GRP78, Calnexin, COX-IV, VDAC2, HSP70, or
315 3 β HSD2 protein expression (Fig. 1C). However, the expression of StAR was marginally induced, and CHOP
316 protein was highly induced with exposure to stress (Fig. 1C).

317 To understand the effect of increased CHOP expression on mitochondrial activity, we next determined
318 pregnenolone and progesterone synthesis in response to stress. As shown in Fig. 1D (top left panel),
319 pregnenolone synthesis increased in response to stress from 15 ng/mL to 32 ng/mL (right panel). Because StAR
320 expression was only moderately increased with stress, the increased metabolic activity is likely due to the
321 appearance of CHOP, which may increase StAR activity by altering its folding to facilitate more cholesterol
322 transfer.

323 To understand the effect of stress on progesterone synthesis, we performed metabolic conversion assays
324 to evaluate pregnenolone to progesterone conversion. As shown in Fig. 1E, progesterone synthesis increased in
325 response to stress from 18 ng/mL to 37 ng/mL. Western blot analysis of the mitochondria applied in each
326 reaction from both Fig. 1D and 1E (bottom panels) shows that 3 β HSD2 and VDAC2 levels were similar.
327 Measurement of stress by serum corticosterone analysis showed an increase from 1.4 ng/ml to 1.68 ng/ml ($p \leq$
328 0.0003) after stress (Fig. 1F). Thus, stress increased the availability of the substrate, pregnenolone (Fig. 1D), by
329 two-fold, in the absence of large changes in protein expression. Because protein folding is a rapid process,
330 stress may have induced changes in 3 β HSD2 folding, which may be mediated by CHOP.

331 *Cytoplasmic CHOP interacts with MAM-associated StAR*

332 To determine the role of CHOP in folding IMS-resident proteins, we established stable MA-10 cells
333 expressing CHOP siRNA (Δ CHOP MA-10 cells). As shown in Fig. 2A, CHOP expression was undetectable
334 following its silencing (left panel). We next isolated the crude mitochondria, which also includes proteins from
335 the MAM (39), from the WT and Δ CHOP MA-10 cells, solubilized with digitonin, separated the protein
336 complexes via native-gradient PAGE, and stained with a 3 β HSD2 antibody (Fig. 2A, right panel). Two
337 complexes of 750 and approximately 550 kDa were identified; however, the intensity of the complexes was
338 lower in the Δ CHOP MA-10 cells, which may be due to the reduced concentration of the active 3 β HSD2 and
339 StAR in the absence of CHOP expression.

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340 To identify the proteins present within the two complexes, we performed mass spectrometry (LC MS/MS)
341 analysis. As shown in Tables 1 and 2, the proteins present in both the complexes from the WT cells, including
342 MAM- and OMM-resident proteins, were similar, suggesting that the 550 kDa complex (Table 2) is possibly a
343 breakdown of the larger 750 kDa complex (Table 1). Only Tom22 disappeared from the 3 β HSD2-containing
344 complex in the absence of CHOP (Tables 3 and 4), suggesting that it may bridge the OMM side facing the
345 cytoplasm and the IMS of the mitochondria. The absence of Tom22 in the 3 β HSD2-containing complex in
346 Δ CHOP cells suggests that its signal is either below the level of detection of the system or that it is buried
347 within the sequence, where stress facilitates the interaction of Tom22 with other mitochondrial proteins, as well
348 as the proteins present at the MAM.

349 To understand the impact of stress and, therefore, CHOP expression on 3 β HSD2 and Tom22
350 interaction, we performed co-IP analysis with adrenal mitochondrial fractions. Under unstressed conditions,
351 3 β HSD2 interacts with StAR, VDAC2, but not with Tom22 (Fig. 2B, top panel). However, Tom22 antibody
352 recognized 3 β HSD2 minimally showing a partial interaction between Tom22 and 3 β HSD2 possibly due to the
353 availability of all the Tom22 epitopes (Fig. 2B, Tom22 panel). In the absence of stress, CHOP is not expressed
354 and, thus, did not interact with 3 β HSD2 or VDAC2 (Fig. 2B, bottom Panel). However, in the presence of stress,
355 the interaction between 3 β HSD2 and StAR was increased, and interaction between 3 β HSD2 and Tom22 was
356 observed (Fig. 2C, top panel). As expected, CHOP expression was induced (Fig. 2C, bottom panel). In the
357 CHOP-null mice after stress, no interaction was noticed between StAR and Tom22 (Fig. 2D, second panel from
358 the top) or between 3 β HSD2 and Tom22 (Fig. 2D, top panel). For further confirmation about the roles of
359 CHOP and VDAC2, we analyzed protein expression in MA-10 and COS-1 cells along with adrenal lysates
360 from mice with or without stress. In CHOP knockdown mice adrenals, StAR expression was similar to the level
361 observed in WT mice (StAR lane), and the expression of mitochondrial 3 β HSD2, GRP78, and calnexin
362 remained unchanged (Fig. 2E). As expected nonsteroidogenic COS-1 cells only expressed VDAC2, GRP78,
363 calnexin and HSP70, but not StAR and 3 β HSD2 (Fig. 2E). We have also examined mitochondrial viability by
364 measuring the ATP release before and after addition of increasing amounts of mCCP to the mitochondria
365 isolated from WT and Δ CHOP MA-10 cells. As shown in Fig. 2F, no difference in mitochondrial ATP content
366 was detected between the two cell lines. Thus, CHOP may facilitate the interaction between StAR and 3 β HSD2

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367 or Tom22. Given that the majority of CHOP is found within the ER and MAM (Fig. 2G, top panel), its
368 interaction with 3 β HSD2 may be facilitated by Tom22.

369 ***Domains mediating StAR-Tom22 interaction***

370 Tom22 is an OMM-associated mitochondrial receptor that interacts with IMM-associated 3 β HSD2 via its
371 C-terminus that is exposed to the IMS (17). We hypothesized that CHOP might directly influence Tom22
372 activity or alternatively work in conjunction with StAR. Because the complete 3-dimensional (3D) structure of
373 Tom22 is unknown, we developed a 3D structural model of the Tom22 (NCBI NP_064628.1, Uniprot
374 Q9NS69) sequence (AA 1-142) using the structures of NAD-dependent aldehyde dehydrogenase from *L.*
375 *acidophilus* (3ROS) and Esta from *A. nitroguajacolicus* (3ZYT) as templates and YASARA (29). As shown in
376 Fig. 2H and J, the Tom22 structural model indicates that it should have a minimum amino acid region facing
377 the IMS. A space filling model integrated with the membrane (Fig. 2I) indicates that the L2 loop is out of the
378 membrane facing the cytoplasm. Similarly only a small, unstructured stretch of 20 amino acids from the C-
379 terminus is exposed to the IMS. Thus, it is likely that the L2 loop comprising the acidic amino acids
380 “EEDDDEELD” might be responsible for interaction with proteins from the cytoplasm, including StAR.
381 However, the complete unstructured L8 loop in combination with the H8 helix might be responsible for
382 interaction with 3 β HSD2. Furthermore, the basic amino acids within the smaller transmembrane loop L6 may
383 participate in interaction; however, its short length makes this less likely because of the cost of entropy required
384 for opening will be higher than the energetic stabilization forming complex. Taken together, the loose regions
385 of Tom22 (Fig. 2H) facing the IMS are likely critical for its interaction with 3 β HSD2, which has a flexible
386 conformation.

387 To identify specific amino acids required for Tom22-3 β HSD2 interaction we expressed Tom22 mutants, in
388 which an L2 aspartic acid (amino acids 35 and 37) or glutamic acid (amino acid 39) is changed to an alanine, in
389 Δ Tom22 MA-10 cells. As shown in Fig. 3A (top panel), Tom22 siRNA reduced Tom22 expression by >90%,
390 and expression of WT, D35A and D37S amino acids restored Tom22 expression to levels similar to untreated
391 MA-10 cells. The E39A and Δ 35-39 Tom22 restored about 50% expression. VDAC2 expression remained
392 unchanged (Fig. 3A, bottom panel). Mitochondrial compartmental fractionation (Fig. 3B) confirmed that the
393 Tom22 proteins were localized at the OMM (Fig. 3B top panel) similar to the OMM marker, VDAC2 (Fig. 3B,

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middle panel). To confirm that the mutants were integrated with the OMM in MA-10 cells after knockdown with siRNA of Tom22, we isolated mitochondrial compartments, and then immunoblotted with a Tom22 antibody. In addition to WT Tom22, the mutants, D35A, D37A, E39A and the three amino acid substituted mutant Tom22 (35-39 Mut), were present only with the OMM fraction similar to that observed for VDAC2 (Fig. 3C, top and second panels). The accuracy of mitochondrial fractionation was confirmed by staining the same fractions with the mitochondrial matrix protein, COX-IV, the OMM-associated VDAC2, the IMS resident protein 3 β HSD2 and the IMM-associated Tim23 (40).

We next confirmed the localization of WT and mutant Tom22 through immune electron microscopy staining with Tom22 and 3 β HSD2 antibodies independently and also together. WT Tom22 was present at the OMM (Fig. 3D, left bottom panels, red arrow). Co-staining for the IMS resident, 3 β HSD2, confirmed that Tom22 (red arrows) was present at the OMM while 3 β HSD2 (blue arrows) was localized inside the mitochondria (Fig 3D, right panels). The bottom panels show an enlarged view of mitochondrion presented above. Similar analyses with the Tom22 mutants revealed that 3 β HSD2 (blue arrows) was present inside the mitochondria while the Tom22 mutants (red arrows) were at the OMM (Fig. 3E). In summary, within the L2 loop region facing the cytoplasm whether it is a point mutant or substitution of four amino acids still Tom22 folding remained unchanged.

Analysis of metabolic conversion of pregnenolone to progesterone showed that it was greatly reduced upon Tom22 knockdown (from 16.5 to 5 ng/mL) (Fig. 3F). Furthermore, expression of the D35A or E39A mutants did not restore metabolic activity as compared to the D37S mutant (Fig. 3F); the activity resulting following transfection with the triple mutant, D35A, D37S and E39A (35-39 Mut), was reduced by 90%. Overexpression of WT Tom22 restored metabolic activity by >80%. Similar results were observed when we analyzed the metabolic conversion of 14 C-cholesterol to 14 C-pregnenolone (Fig. 3G). In summary, the L2 loop region of Tom22 may play a significant role in regulating metabolic activity.

3 β HSD2 is a loosely folded protein, and it is likely to interact with the loosely binding regions of Tom22. We next analyzed the interaction of the Tom22 mutants with 3 β HSD2 by co-IP analysis. As shown in Fig. 3H, the intensity of the interaction decreased with the triple mutant almost completely, and also minimal interaction

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420 with the D35A mutant was observed (Fig. 3H, middle panel). Wild-type cells do not express CHOP, which
421 possibly assists in the interaction from the cytoplasmic side upon expression. Alternatively, the change in L2
422 loop amino acids possibly altered Tom22 conformation, resulting in a misfolded protein and suggesting that
423 this region is responsible for interaction with StAR directly or through CHOP.

424 ***Tom22 L8 –H8 is essential for metabolic regulation***

425 Tom22 has many unstructured regions (41), which allows it to remain flexible while associated with the
426 membrane. The partially developed 3D-solution structure predicted that Tom22 should have a minimum amino
427 acid region facing the IMS, comprising amino acids 128 to 142 (41), that may mediate its interaction with
428 3 β HSD2 at the IMS. Although a smaller transmembrane loop with basic amino acids may participate in the
429 interaction, its short length makes this less likely. To confirm that indeed the C-terminus of Tom22 is
430 responsible for its interaction with 3 β HSD2, we expressed various Tom22 mutants from small to large amino
431 acids or from polar to nonpolar or basic amino acids, generating Q118L, T125I, M131R, A134D and N124K
432 Tom22, and expressed these mutants in Δ Tom22 MA-10 cells. So we knocked down Tom22 expression by
433 siRNA in MA-10 cells (Fig 4A). Western blot analysis confirmed the Tom22 mutant expression (Fig. 4A, top
434 panel), and mitochondrial fractionation confirmed their localization to the OMM (Fig. 4B). Electron
435 microscopy was next employed to determine Tom22 mutant localization as well as that of the IMS resident,
436 3 β HSD2 (Fig. 4C). As expected, 3 β HSD2 (blue arrows) was present inside mitochondria, and the Tom22
437 mutants (red arrows) were associated with OMM.

438 Although progesterone conversion was not restored with M131R expression, it was restored upon
439 expression of all other Tom22 mutants, Q118L, T125I, N124K, and A134D (Fig. 4D). Although amino acids
440 125-142 are within the unstructured L8 loop, only the M131R mutant did not restore complete activity,
441 suggesting that the loop region possibly has a specific conformation, which is altered when the hydrophobic
442 methionine is replaced with a charged, bulky polar amino acid. Surprisingly, the helix region Q118L mutant
443 restored activity, which may be due to the similar size of the amino acid or that the mutation did not affect the
444 folding due to lipid integration (Fig. 4D). Co-IP analysis confirmed that 3 β HSD2 interacted with the N124K
445 and T125I mutants preferentially at a higher intensity than the M131R, A134D and Q118L mutants (Fig. 4E,
446 top and middle panels), suggesting that the unstructured region of Tom22 interacts with 3 β HSD2. Arginine is a

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large amino acid with a long carbon chain. There is possibly a change in conformation from methionine to Arginine of M131R resulting in loss of activity. However, it is possible that the A134D mutant is present inside the loop and thus it is not easily accessible to 3β HSD2 exposed sites for interaction resulting no loss in activity. In summary, these results confirmed that metabolic activity is increased due to the presence of CHOP in the cytoplasm, which acted on the charged amino acids of L2 loop, and that L8 loop is responsible for interaction with 3β HSD2 at the IMS.

Metabolic activity is dependent on phosphate circulation

We next sought to identify the condition in which Tom22 facilitates metabolic activity at the IMS. We hypothesized that phosphate circulation across the OMM is essential for mammalian pregnenolone synthesis albeit through an unknown mechanism possibly increasing stability in the complex (42). 3β HSD2 expression is not changed in response to stress and requires a molten globule conformation in an acidic environment for activity, facilitated by the mitochondrial proton pump (23). The presence of CHOP increased pregnenolone synthesis more than 2-fold in comparison to the absence of stress (Fig. 1D); therefore, we hypothesized that the OMM receptor, Tom22, possibly facilitates interaction with 3β HSD2 while interacting with StAR-VDAC2, resulting in increased enzymatic activity at the IMS in the presence of circulating phosphates. CHOP is expressed during stress, and facilitates interaction with 3β HSD2 through the StAR-VDAC2-Tom22 complex (Fig. 2C) and, thus, possibly plays a central role in increasing the circulation of phosphates in association with Tom22. However, no changes in adrenal 3β HSD2, Tom22, and VDAC2 expression were detected in either WT or CHOP-null mice in response to stress (Fig. 2E). Thus, we first attempted to determine a change in cellular architecture by immunohistochemistry analysis of Tom22, StAR, 3β HSD2, Aldosterone Synthase (AS), and VDAC2 expression (Fig. 5A and B); however, no remarkable change was noticed. Furthermore, EM analysis following Tom22 and 3β HSD2 staining showed that the mitochondrial architecture was not altered (Fig. 5C).

Mitochondrial metabolic activity is related to the circulation of phosphates from the OMM to the IMM, as knocking down phosphate carrier protein (PCP) reduced phosphate circulation and pregnenolone synthesis (22). Analysis of serum phosphate levels showed that it increased from 6.2 to 7.4 mM in WT mice ($p \leq 0.00029$) in response to stress; however, the CHOP-null mice had no significant change ($p = 0.153$) in phosphate concentration with stress (Fig. 6A). These results prompted us to explore the possibility that phosphates

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474 increase metabolic activity by association with neighboring proteins, forming a network from the MAM to
475 mitochondria. As shown in Fig. 6B (middle panel), addition of 5 mM phosphate to mitochondria increased the
476 progesterone synthesis from 4 ng/mL to 8.0 ng/mL, peaking at 20 ng/mL with 100 mM phosphates (Fig. 6B,
477 top panel).

478 To determine whether the increased phosphate concentration facilitated 3 β HSD2 association with the
479 translocases and increased its activity, we examined the progesterone levels produced with dephosphorylated
480 3 β HSD2 protein and mitochondria. In the absence of 3 β HSD2, no progesterone was synthesized, confirming
481 that the isolated mitochondria have minimal endogenous activity (Fig. 6C). Incubation of 3 β HSD2 with the
482 mitochondria in the absence of phosphate or with mitochondria devoid of phosphate resulted in 4.1 ng/mL
483 progesterone synthesis as compared to the 18 ng/mL of progesterone produced with normal 3 β HSD2 and
484 mitochondria (Fig. 6C). Incubation of mitochondria and dephosphorylated-3 β HSD2 with sodium phosphate
485 restored about 50% activity (Fig. 6D, top and right panels), suggesting that 3 β HSD2 requires phosphate ions for
486 progesterone synthesis. An increase in activity with SAP more than 0.5 Unit/ μ l suggests a reversible
487 phosphorylation (43). The SAP enzymatic reaction proceeds through a covalent phosphoserine intermediate to
488 produce inorganic phosphates or to transfer phosphoryl group to alcohols resulting more phosphate ions (44).
489 So, the activity did not increase beyond 50%, as the inorganic phosphates inhibits shrimp alkaline phosphatase
490 at higher concentrations (43); thus, when we incubated the reaction with external inorganic phosphates, only
491 50% of 3 β HSD2 activity was restored.

492 The predicted phosphorylation sites were determined by the NetPhos program (45). Based on the
493 computer modeling (46) and phosphorylation prediction (45), serine residues 123 and 124 were most crucial for
494 S-S bonds and S-T bonds (Fig. 6E). Three more amino acids, T288, S358 and S369, are also possibly
495 phosphorylated as these residues are exposed to the outer surface of the helical structure (Fig. 6E). The best
496 structure binding site for post-translational modification of arginine 183 is present in the groove. To determine
497 the specificity, we mutated arginine 183 to glutamic acid and changed S123, S124, T288, S358 and S369 to
498 alanine and threonine, and determined progesterone synthesis following our earlier procedure (22). As shown in
499 Fig. 6F, WT 3 β HSD2 synthesized 7.9 ng/mL of pregnenolone; the S123A and S124A mutants had no activity.
500 The S358A mutant also reduced the activity, synthesizing 2-3 ng/mL of pregnenolone. This is also possible that

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the reduction in activity is due to the change in amino acids. To our surprise, mutating the bulky arginine at position 183 to glutamine also ablated 3 β HSD2 activity similarly to the S123A or S124A mutants. Western blot analysis of the transfected cells showed similar levels of mutant 3 β HSD2 expression as compared to the WT protein (Fig. 6G, top panel); VDAC2 expression levels were also similar (Fig. 6G, bottom panel). To confirm whether the inactivity is due to conformational change and not on phosphorylation, we changed S123A to S123T and similarly S124A to S124T. Also to completely rule out any error we also developed a combined serine to threonine mutant together of the 123 and 124 residues. The activity from S123T and S124T combined mutation or individual mutation ablated activity completely, suggesting that the reduction in activity is possibly due to change in amino acids not likely due to phosphorylation (Fig. 6H). Western blot analysis of the transfected cells showed similar levels of mutant S123T and S124T 3 β HSD2 expression as compared to the WT and the amount of protein content applied in each lane was also identical as observed by staining with VDAC2 antibody (Fig. 6I). In summary, our results confirm that the reduced activity observed in the other mutants was likely not due to reduced phosphorylation but due to a change in the amino acids.

3 β HSD2 conformation is stabilized by the circulation of phosphates

To better understand the role of phosphate circulation on Tom22 and 3 β HSD2 function, we isolated mitochondria from MA-10 cells, which were devoid of phosphate. After various concentrations of phosphates were added, the complexes were isolated, solubilized with digitonin, and analyzed by native gradient PAGE and staining with a 3 β HSD2 antibody (Fig. 7A). The intensity of 450 kDa complex was increased more than 3.5 fold on addition of 5 mM phosphate to the phosphate deprived cells, but the intensity decreased dramatically with concentrations more than 15 mM (Fig. 7A and B). As a comparison, we isolated the mitochondria from MA-10 cells, then added various concentrations of phosphate and analyzed in an identical fashion as described above (Fig. 7C). The bottom panels show that the same amount of mitochondrial proteins was applied in each reaction (Fig. 7A and C). In the absence of phosphate, two complexes of 750 kDa and 450 kDa were observed (Fig. 7A). In the presence of low phosphate concentrations, the 450 kDa complex became more intense (Fig. 7C) with only the 750 kDa complex detected at high phosphate concentrations, suggesting that the addition of extra phosphate increased the stability of high molecular weight complex possibly through increased association of neighboring proteins. An analysis is presented in Figs. 7B and 7D.

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528 We next determined how the phosphorylation of 3 β HSD2 impacts its conformation by dephosphorylating
529 active 3 β HSD2 protein prepared by a baculovirus system and analyzing its conformation by CD. The
530 equilibrium CD spectra of dephosphorylated and phosphorylated 3 β HSD2 have minima at 208 and 222 nm,
531 respectively, which is typical of a α -helical conformation (data not shown). We next determined the stability
532 3 β HSD2 protein in presence and absence of phosphate by thermal denaturation to identify different states of the
533 protein conformation. As shown in Fig. 7E, 3 β HSD2 unfolds partially at 27°C, after which it stabilizes until it
534 starts unfolding at 41°C and is completely unfolded at 59°C with a T_m of 51°C. The two-step unfolding is
535 suggestive of complete denaturation (Fig. 7E), where the initial unfolding is possibly due to a cooperative
536 intramolecular association with most of the domains and was resistant to unfolding because of its molten
537 globule conformation (23). In the absence of phosphate anions, the protein displayed reduced stability, and the
538 initial unfolding is lost. The initial T_m of the 3 β HSD2 protein at 33°C is suggestive of its retaining the initial
539 change of one conformational state to the second one. This first unfolding is indicative of a pseudo-stable
540 conformation in which most of the secondary structure is preserved and may serve as a way for the protein to
541 maintain equilibrium with the native state.

542 At 33°C, the enthalpy (ΔH) of 3 β HSD2 was 1.2 kCal/mol, increasing to 4.1 kCal/mol at 50°C (Fig. 7E).
543 When the salt concentration was increased to 1.0 M, the two-step unfolding was lost due to salt-induced
544 denaturation (Fig. 7F). As seen from the model structure (Fig. 6E), the globular structure is extremely flexible.
545 Thus, it is likely that the intramolecular association is favored by the phosphate anion on the positively charged
546 amino acids, changing 3 β HSD2 organization even at lower temperatures. In presence of phosphates, an
547 equilibrium between the initial state of unfolding from N \rightarrow I_u \leftrightarrow Pu \geq IU(T_m) \rightarrow D to the unfolded state occurs
548 through a three-step process (Fig. 7G). 3 β HSD2 conformation from the initial unfolding state (I_u) to the
549 pseudo-unfolded state (Pu) is in equilibrium at 33°C. The protein undergoes an initial step of unfolding or an
550 intermediate stage of unfolding (IU), which is the thermal unfolded state at 50°C. In the next step, the protein
551 undergoes complete unfolding or denaturation into an unfolded (D) state. In summary, the results strongly
552 suggest that phosphate circulation increased 3 β HSD2 association with neighboring proteins, resulting in
553 increased stability and interaction with Tom22 and StAR.

554 **DISCUSSION**

555 Both the ER and mitochondria are dynamic organelles capable of modifying their structure and function in
556 response to changing environmental conditions. ER and mitochondria interact both physiologically and
557 functionally due to their close proximity (47). Newly synthesized proteins, including mitochondrial proteins,
558 require folding within the ER lumen prior to trafficking to the next cellular compartment as programmed by
559 their signal sequences. Specifically, nascent polypeptide chains emerge in the ER lumen, where
560 posttranslational modifications, such as N-linked glycosylation and intra- and intermolecular disulfide bond
561 formation, facilitate the folding of polypeptides to form specific tertiary and quaternary structures for proper
562 protein function (48). The quality of protein folding is precisely monitored by an ER quality control system that
563 only allows properly folded proteins to be transported to the specific organelle and directs misfolded proteins
564 for ER-associated degradation (ERAD) by the 26S proteasome or for degradation through autophagy (49, 50).
565 ER stress impairs this process and results in the accumulation of unfolded or misfolded proteins, resulting in
566 UPR and the expression of a wide range of genes and activation of proteins involved in the maintenance of ER
567 function (6), including IRE1, PERK, ATF6, and CHOP.

568 In the present study, CHOP expression did affect the expression of ER-, MAM-, or mitochondrial-resident
569 proteins in the adrenal glands. The presence of CHOP at both the ER and MAM possibly influenced MAM and
570 mitochondrial proteins. ER and mitochondria form close contacts with 20% of the mitochondrial surface in
571 direct contact with the ER (51), which facilitates communication between the organelles (39). Thus, the
572 purified MAM showed the presence of Tom22 and VDAC2, which are part of the MAM but permanently
573 present at the mitochondria facing the OMM. Since Tom22 interacts with the IMS resident, 3 β HSD2, it was
574 pulled out during the co-IP analysis.

575 The expression of CHOP through stress coincided with increased mitochondrial metabolic activity.
576 Because serum phosphate concentrations in CHOP-null mice exposed to stress were reduced by approximately
577 20% as compared to WT mice exposed to stress (Fig. 6A), we hypothesized that an increase in mitochondrial
578 phosphate supply could stabilize 3 β HSD2 conformation, increasing metabolic activity that resulted in >2-fold
579 progesterone synthesis. The presence or absence of CHOP expression did not alter the expression of other ER-
580 resident or mitochondrial proteins. Because CHOP is a chaperone, it may assist in the folding of Tom22 and

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581 StAR, especially given that it was located both in the ER and MAM regions of adrenal cells. Tom22 is a MAM-
582 associated protein from the mitochondrial side; however, CHOP faces the ER side of the MAM, suggesting that
583 they only transiently interact, which may be why CHOP interaction with Tom22 was not demonstrated by co-IP
584 analysis in the absence of stress (Fig. 2B, top panel). Similarly, StAR is synthesized on acute hormonal
585 regulation or during stress, remaining at the MAM prior to loading onto the VDAC-containing region of the
586 OMM. Tom22, VDAC2, and StAR are all part of a MAM fraction. There is a partial interaction between
587 Tom22 and 3β HSD2 at the IMS, but interaction is stronger under stress condition. It is possible that stress
588 generated changes in the conformation of StAR that were mediated by the chaperonic properties of CHOP.

589 Expression of Tom22 mutants did not fully restore the metabolic activity of Δ Tom22 cells, suggesting that
590 the mutants were possibly misfolded as their accurate localization was confirmed. Although Tom22 appears
591 flexible, it may become significantly structured upon OMM integration, especially given that the mutation in
592 the H8 helix did not ablate activity but a mutation in the unstructured loop region (L8) completely ablated
593 activity. An L2 mutant also failed to interact with 3β HSD2 via co-IP analysis, suggesting that this region is
594 crucial for metabolic regulation at the IMS. Since StAR residency is shorter at the MAM and longer at the
595 OMM, stress may have facilitated a structural change in Tom22 due to the presence of flexible folding
596 associated with the lipid membrane. As a result, the mitochondrial import channel may have opened, resulting
597 in increased metabolic activity with the presence of more phosphates, while the substrate remained the same for
598 the isomerase and dehydrogenase activities at the IMS (23).

599 Stress-induced CHOP expression coincided with a 2-fold increase in pregnenolone synthesis (Fig. 1D),
600 which is possibly due to the 1.5-fold increase in expression of StAR (Fig. 1C). Through physical interaction,
601 CHOP may have facilitated a conformational change in Tom22 or in StAR-Tom22. StAR is transiently
602 associated at the MAM from the ER side, so it is more likely facilitated through StAR. Thus, CHOP may push
603 Tom22 like a piston, resulting in the L8 loop of Tom22 coming in close proximity to 3β HSD2, facilitating its
604 catalysis. This would explain how a mild increase in StAR expression by stress increased the catalytic reaction
605 by two-fold largely independent of changing protein concentration.

606 The presence of increased phosphates stabilized 3β HSD2 protein conformation by increasing its interaction
607 with neighboring proteins. The intensity of the 450 kDa 3β HSD2-containing complex increased following

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608 addition of low concentrations of phosphate with higher phosphate concentrations stabilizing the 750 kDa
609 complex alone (Fig. 7A), suggesting that the presence of phosphate increased association of Tom22 with the
610 neighboring proteins from the OMM-exposed region. The association is possibly facilitated by the stabilization
611 of 3 β HSD2 conformation as confirmed by its higher T_m (Fig. 7E and F). Thus, the circulation of phosphates
612 possibly facilitates the stabilization of 3 β HSD2 in a dynamic state and may be mediated by the presence of
613 CHOP interacting with Tom22 or StAR-Tom22.

614 The molecular identity of the mitochondrial permeability pore, which is comprised of dimers of the
615 mitochondrial ATP synthase (52), is now better understood. IMM permeability is increased in response to
616 stress, leading to the formation of the voltage-dependent nonspecific pore, PTP (53), that is permeable to water,
617 ions, and substances of low molecular mass. Chronic stress leads to mitochondrial dysfunction due to extensive
618 PTP opening and ATP hydrolysis, during which the cells cannot maintain structural and functional integrity. In
619 addition, protein conformation cannot be restored under chronic stress. In the stress model used in the present
620 study, acute stress likely restored PTP to its original conformation, increasing the phosphate concentration
621 within the mitochondria due to PTP closure.

622 In summary, short periods of stress increased the metabolic activity of adrenal cells that coincided with
623 increased CHOP expression. Either directly or through StAR, CHOP acts on the mitochondrial translocase,
624 Tom22, facilitating its interaction with 3 β HSD2 via its IMS-exposed unstructured region. Stress also facilitated
625 phosphate transport into the mitochondria, which stabilized 3 β HSD2 conformation likely via circulation of
626 phosphates and increasing its activity.

627 ACKNOWLEDGEMENTS

628 HB was supported by grants from the National Institutes of Health (HD057876) and the Anderson Cancer
629 Institute. We are thankful to Dr. Satyam Banerjee for participating during preliminary experiments and
630 performing the animal surgeries. We are also thankful to Mr. Greg Roberson for the maintenance of animals
631 and his assistance during the experimental procedures and John Knights for assistance in the
632 immunohistochemistry image capturing. MP is thankful to Mr. Subodh Thapliyal for his help in statistical
633 analysis.

634 **Conflict of Interests:** The authors declare that they have no conflict of interest.

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635 **Authors Contribution:** HB conceptualized the project, analyzed the data and wrote the manuscript; MP,
636 WEB, JK, SAP, AW, AVP, GP and RMW performed the experiments; JLT provided reagents and MP prepared
637 all the figures and analysed data. A complete manuscript was available to all of the authors prior to initial
638 submission.
639

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- 764

765 **FIGURE LEGENDS**

766 **Figure 1.** Effect of stress on adrenal steroid synthesis. (A) Schematic presentation of steroid biosynthesis.
767 StAR is synthesized upon stress and hormonal stimulation, but CHOP is expressed only on stress. After
768 reaching near the outer mitochondrial membrane, StAR facilitates fostering cholesterol from the outer to inner
769 mitochondria to initiate steroid synthesis. (B) Schematic presentation of localization of different membrane
770 associated proteins, which are part of the MAM fraction present in the cytoplasm. A possible role of the outer
771 and inner mitochondrial translocases and their interaction with steroidogenic proteins is clearly described. (C)
772 Protein expression in the adrenal glands of mice with or without exposure to stress. The adrenal glands were
773 isolated immediately following the last period of stress, and proteins were analyzed by Western blotting with
774 the indicated antibodies. Mouse Leydig MA-10 cells were serum starved and cell lysates were used as a
775 control. (D) Measurement of ^{14}C -cholesterol to ^{14}C -pregnenolone conversion by mitochondria isolated from
776 the adrenal glands of the experimental animals. The metabolic reactions were initiated with NADPH at 37°C .
777 Trilostane (Tril) was applied as an inhibitor of $3\beta\text{HSD2}$ activity. The right panel shows a quantitative estimate
778 of the amount of pregnenolone synthesized from the metabolic reactions. The bottom panels show the Western
779 blots of the mitochondrial lysates stained with VDAC2 and $3\beta\text{HSD2}$ independently. (E) ^3H -Pregneolone to
780 ^3H -progesterone conversion by adrenal mitochondria isolated from animals with and without exposure to stress.
781 The metabolic reaction was initiated with NAD and Trilostane (Tril) was applied as an inhibitor of $3\beta\text{HSD2}$
782 activity. The right panel shows a quantitative estimate of the amount of progesterone synthesized from the
783 metabolic reactions. The bottom panels show the Western blots of the mitochondrial lysates stained with
784 VDAC2 and $3\beta\text{HSD2}$ independently. (F) Analysis of the change in serum corticosterone synthesis before and
785 after stress. The serum corticosterone increased from 1.4 ng/ml to 1.68 ng/ml following stress ($p \leq 0.0003$).
786 Data in panels D-F are the mean \pm SEM of at least three independent experiments performed at three different
787 times.

788 **Figure 2.** CHOP expression impacts $3\beta\text{HSD2}$ -Tom22-VDAC2 interaction. (A) Left, siRNA mediated CHOP
789 knockdown. Right, Crude mitochondria were isolated from the wild-type MA-10 and ΔCHOP MA-10 cells,
790 solubilized in digitonin, analyzed through 4-16% native gradient PAGE and stained with $3\beta\text{HSD2}$ antibody.
791 The left bottom panel show equal $3\beta\text{HSD2}$ levels from the mitochondria applied in each reaction. (B - D) Co-

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792 immunoprecipitation of the digitonin-solubilized lysate from adrenal mitochondria of wild-type mice without
793 (B) and with stress (C), and CHOP-null mice with stress (D). In all control lanes (First lanes in B-D) lysate
794 from stressed animal adrenals were applied. The immunocomplexes were isolated using the indicated
795 antibodies and then stained with 3 β HSD2, Tom22, VDAC2, and CHOP antibodies independently. (E)
796 Expression of 3 β HSD2, VDAC2, HSP70, StAR, GRP78, calnexin (cal) and CHOP by the wild-type and
797 CHOP-null adrenals with and without stress by Western blotting. Nonsteroidogenic COS-1 cells were applied
798 to show its difference from MA-10 cells. Serum starved MA-10 cell lysate was used as control. (F)
799 Measurement of ATP release by mitochondria isolated from wild-type and Δ CHOP MA-10 cells after
800 incubation with various concentrations of mCCP. (G) Percoll density distribution analysis of the subcellular
801 localization of CHOP, 3 β HSD2 and VDCA2 after stress. (H) Modeling of Tom22 and analysis of its membrane
802 association. A computer model structure of Tom22 showing its extremely flexible nature with the maximum
803 unstructured region and the absence of sheets. (I) The space filling model of Tom22 showing its organization
804 within the OMM. (J) Helical organization of the amino acids present in the Tom22 model shown in panel H.
805 Data in panel F is the mean \pm SEM of at least three independent experiments performed at three different times.
806 **Figure 3.** Domains mediating 3 β HSD2-Tom22 interaction. (A) Expression of the Tom22 L2 loop region
807 mutants, D35A, D37S, E39A and 35-39 Mut, in Δ Tom22 cells. The scrambled siRNA was used as a negative
808 control (Neg siRNA). The top panel shows the quantitative expression level of the Tom22 mutants. Bottom,
809 VDAC2 expression remained unchanged. (B) Fractionation of the mitochondrial compartments of MA-10 cells
810 and staining with the indicated antibodies either in combination or independently. The import fragment was
811 proteolyzed by proteinase K (PK). (C) Mitochondrial fractionation of Δ Tom22 MA-10 cells following
812 expression of the indicated mutants and staining with Tom22, VDAC2, Tim23 and COX-IV antibodies
813 independently. (D) Immunoelectron microscopy of the Tom22 (red arrows) independently or with 3 β HSD2
814 (blue arrows) as indicated. (E) Immunoelectron microscopy of Δ Tom22 cells after expression of the indicated
815 mutants (red arrows) and then colocalization with 3 β HSD2 (blue arrows). The bottom panels are enlarged
816 images of representative mitochondria from the same experiment indicated above. (F) Metabolic conversion of
817 3 H-pregnenolone to 3 H-progesterone by Δ Tom22 cells expressing the indicated Tom22 mutants and initiated
818 with NAD. Trilostane (Tril) was applied as a negative control, and Neg siRNA was applied to show the

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819 specificity of the Tom22 siRNA. The top panel shows the quantitative measurement of the progesterone
820 synthesized. (G) Metabolic conversion of cholesterol to pregnenolone by Δ Tom22 cells expressing the
821 indicated Tom22 mutants. The top panel shows a quantitative estimation of the amount of pregnenolone
822 synthesized in the bottom panel. (H) Co-immunoprecipitation of the Tom22 mutants and Western staining with
823 Tom22, 3β HSD2 and VDAC2 antibodies independently. Reprobing the same membrane with VDAC2 indicates
824 equal amount of protein present in each lane. Data in panels A, F and G are the mean \pm SEM of at least three
825 independent experiments performed at three different times.

826 **Figure 4.** Specificity of IMS resident amino acids. (A) Expression of the H8 and L8 loop region mutants,
827 Q118L, N124K, T125I, M131R and A134D, in Δ Tom22 cells. The scrambled siRNA was used as a negative
828 control. Bottom, VDAC2 expression remained unchanged. (B) Mitochondrial fractionation of Δ Tom22 cells
829 after expression of the indicated mutants and analysed after staining with Tom22, VDAC2, Tim23 and COX-IV
830 antibodies independently. (C) Immunoelectron microscopy of the Δ Tom22 cells following expression of the
831 mutants (red arrows) and their colocalization with 3β HSD2 (blue arrows). The bottom panels are enlarged
832 images of representative mitochondria from the same experiment indicated above. (D) 3 H-Pregnenolone to 3 H-
833 progesterone conversion following expression of the indicated mutants and initiation with NAD. Trilostane was
834 applied as a negative control, and the Neg siRNA was applied to show the specificity of the Tom22 siRNA.
835 The top panel is the quantitative estimation of the amount of steroid synthesized. (E) Co-immunoprecipitation
836 of the indicated Tom22 mutants with Tom22, 3β HSD2 and VDAC2 antibodies independently. 3β HSD2
837 antibody staining on longer exposure is also presented just bottom of the 3β HSD2 panel and shown by a
838 curved arrow. Data in panel D is the mean \pm SEM of at least three independent experiments performed at three
839 different times.

840 **Figure 5.** Analysis of cellular architecture by immunohistochemistry. (A) Sections of wild-type (left
841 panels) and CHOP null (right panels) adrenal glands probed by immunohistochemical staining for the expression
842 of Tom22 (panels a and b); StAR (panels c and d); 3β HSD2 (panels e and f); aldosterone synthase (panels g
843 and h); VDAC2 (panels i and j). The cells contain intracytoplasmic lipid stores. Aldosterone synthase
844 immunoreactivity was localized in the Zona Glomerulosa of both specimens (panels g and h). The remainder of

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845 the panels shows findings from the Zona Fasciculata of the two specimens (hematoxylin counterstain, original
846 magnification: X400). (B) The identical preparations at X1000 magnification from the panel A. The pattern of
847 immunoreactivity is cytoplasmic for all antigens in both specimens. (C) Electron microscopy of adrenals from
848 wild-type and CHOP $-/-$ mice stained with Tom22 without and with 3β HSD2. The top panels show the enlarged
849 view of mitochondria.

850 **Figure 6.** Role of phosphates in steroid synthesis. (A) Serum phosphate levels in wild-type and CHOP $-/-$ mice
851 exposed to stress as compared with controls. The serum phosphate of the wild-type mice was increased from
852 6.2 mM to 7.4 mM ($p \leq 0.0029$) following stress but CHOP null mice had no difference ($p \leq 0.153$) before and
853 after stress. (B) ^3H -Pregneolone to ^3H -progesterone conversion by mitochondria isolated from MA-10 cells
854 incubated with 3β HSD2 and various NaH_2PO_4 concentrations, ranging from 0 to 100 mM. The top panel
855 shows the quantitative measurement of the amount of steroid synthesized. The bottom panel shows that equal
856 amounts of mitochondria were employed. (C) ^3H -Pregneolone to ^3H -progesterone conversion by MA-10 cell
857 mitochondria with 3β HSD2 preincubated with Tris (pH 7.4) for phosphate removal. In some cases,
858 mitochondria or 3β HSD2 were preincubated with Tris buffer. The quantitative estimation is presented in the
859 top. (D) Similar metabolic conversion after dephosphorylation of 3β HSD2 with varying amounts of SAP and
860 then restoring activity after incubation with 100 mM NaH_2PO_4 . Quantitative measurements are shown in the
861 right panel. The bottom panels, C and D, were stained with VDAC2 antibody showing same amount of
862 mitochondrial protein was applied in each reaction. (E) The model, regenerated by Pymol molecular graphics
863 with the solvent preset, shows the presence of six β -sheets numbered 1-6. The protein structure of 3β HSD2
864 shows 12 segments of residues (15–26, 43–56, 66–75, 88–93, 96–115, 154–174, 191–200, 202–207, 225–238,
865 245–253, 331–335, and 346–360) that form 12 α -helices, shown as $\alpha 1$ - $\alpha 12$, and six segments of residues
866 (1–7, 31–36, 57–60, 78–82, 181–183, and 264–266) that form six β -sheets, shown as $\beta 1$ - $\beta 6$. In the solvent
867 view structure of 3β HSD2, the residues for the α -helices are shown in different colors depending on their
868 location. (F, G) Determination of the activity of the 3β HSD2 mutants to define possible role of serine residues.
869 (H, I) Determination of activity (H) and expression (I) of the 3β HSD2 mutants after transfection in
870 nonsteroidogenic COS-1 cells. VDAC2 expression shows the equivalent loading in each lane. Data presented in

871 panels A-D,F and G are the mean \pm SEM of at least three independent experiments performed at three different
872 times.

873 **Figure 7.** Role of phosphate in 3 β HSD2 stabilization. (A) Native gradient PAGE of the digitonin solubilized
874 complex stained with a 3 β HSD2 antibody from the mitochondria isolated from steroidogenic cells deprived of
875 phosphate followed by external addition of 5, 15, 30 and 100mM sodium phosphate. (B) An analysis of the
876 3 β HSD2-containing 450 kDa complex from the phosphate deprived cells followed by addition of indicated
877 concentrations of phosphate. (C) Similar native gradient PAGE of the digitonin-solubilized complex from MA-
878 10 mitochondria after external addition of sodium phosphate in the absence of phosphate deprivation. (D) A
879 comparative analysis of the 450 kDa native complex intensity by external addition of phosphate. (E) Thermal
880 unfolding of the dephosphorylated (purple line) and untreated (black line) 3 β HSD2 recorded in a CD
881 spectropolarimeter at 222 nm from 4°C to 90°C. (F) Thermal unfolding of 3 β HSD2 recorded in a CD
882 spectropolarimeter at 222 nm from 4°C to 90°C with two different concentrations of sodium ions, 1.0M (blue
883 line) and 10mM (black line) NaH₂PO₄, pH 7.4. (G) Schematic presentation explaining the stability of 3 β HSD2
884 with and without phosphate. The wild-type is in a native state “N” and upon gradual thermal denaturation; it
885 forms an intermediate denaturation “Iu” generating thermal unfolding “Tm/Pu” at 33°C with a pseudo plateau
886 at 41°C. Then, 3 β HSD2 goes through a two-step unfolding process with a Tm of 47°C and final denaturation
887 “D”. In the case of the phosphate-preincubated protein, the protein undergoes a two-step unfolding with a Tm
888 of 52°C. Data in panels C and D represent the mean \pm SEM of at least three independent experiments performed
889 at three different times.

890 Table Legends

891 **Table 1 and 3.** Proteins in the 750-kDa 3 β HSD2-containing complex isolated from WT (1) and CHOP
892 knockdown (3) MA-10 cells following native gradient PAGE (4-16%) and mass spectrometry. The common
893 molecules between the 750 and 550 kDa complex of the MA-10 cells are in italics and the CHOP knockdown
894 MA-10 cells are indicated in italics with underlined.

895

896 **Table 2 and 4.** Proteins in the 550-kDa 3 β HSD2-containing complex isolated from WT (2) and CHOP
897 knockdown (4) MA-10 cells following native gradient PAGE (4-16%) and mass spectrometry.

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Table 1 (MA-10 750 kDa native complex)

Accession No.	Protein	MS/MS Score	No. of unique peptides
26984237	mitochondrial ATP-dependent protease Lon	1959	29
344268714	60 kDa heat shock protein, mitochondrial	1319	31
6755967	voltage-dependent anion-selective channel protein 3 isoform 2	589	9
19745150	NADH-cytochrome b5 reductase 3	570	11
73987072	lon protease homolog, mitochondrial isoform 1	527	10
154816168	stress-70 protein, mitochondrial	527	7
351694722	Aldehyde dehydrogenase, mitochondrial	314	9
351708365	NADH-cytochrome b5 reductase 3	256	5
726095	long-chain acyl-CoA dehydrogenase	181	3
149410696	voltage-dependent anion-selective channel protein 3-like isoform 1	179	4
31542438	cytochrome b5 type B precursor	265	4
340546142	voltage-dependent anion channel 1	136	4
9910382	mitochondrial import receptor subunit TOM22	132	3
345305898	voltage-dependent anion-selective channel protein 2-like isoform 1	126	2
47058986	mitochondrial import receptor subunit TOM40	119	2
13385998	heat shock protein 75 kDa, mitochondrial precursor	115	3
301070250	hydroxyacyl-Coenzyme A dehydrogenase	110	2
339243731	ADP/ATP translocase 1calcium binding 140k protein	78	21
16758454	phosphatidate cytidylyltransferase 2	64	1

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Table 2 (MA 10 550 kDa native complex)

Accession No.	Protein	MS/MS Score	No. of unique peptides
194044029	<i>60 kDa heat shock protein, mitochondrial</i>	1066	23
238427	Porin 31HM	561	9
297265430	protein disulfide-isomerase A6 isoform 1	368	6
6755967	<i>voltage-dependent anion-selective channel protein 3 isoform 2</i>	335	7
225706314	<i>Voltage-dependent anion-selective channel protein 2</i>	287	7
860986	protein disulfide isomerase	254	5
307199045	<i>60 kDa heat shock protein, mitochondrial</i>	252	7
340546142	<i>voltage-dependent anion channel 1</i>	144	3
109075624	membrane-associated progesterone receptor component 2	118	1
297272126	<i>fatty aldehyde dehydrogenase isoform 6</i>	110	2
12055497	Ca ²⁺ -ATPase 1	109	1
386758	GRP78 precursor	97	2
550062	GTP-binding protein	93	2
2414516	surface 4 integral membrane protein	76	2
2662291	<i>cytochrome b5</i>	76	1
553254	<i>NADH cytochrome b5 reductase</i>	71	2

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Table 3 (Δ CHOP MA-10 750 kDa native complex)

Accession No.	Protein	MS/MS Score	No. of unique peptides
51455	<u>heat shock protein 65</u>	969	19
344268714	60 kDa heat shock protein, mitochondrial	882	16
6753036	<u>aldehyde dehydrogenase, mitochondrial precursor</u>	875	16
4105605	<u>voltage dependent anion channel</u>	270	5
726095	<u>long-chain acyl-CoA dehydrogenase</u>	250	5
386758	<u>GRP78 precursor</u>	202	4
5733504	<u>voltage-dependent anion channel VDAC3</u>	131	3
205632	<u>Na,K-ATPase alpha-1 subunit</u>	114	2
247307	<u>cytochrome P450 reductase</u>	99	2
118596522	<u>acetaldehyde dehydrogenase</u>	89	2
199599757	protein disulfide isomerase-associated 3 precursor	83	2
147898791	ADH-cytochrome b5 reductase 2 precursor	71	1
4566870	malate dehydrogenase	68	1
225706314	Voltage-dependent anion-selective channel protein 2	57	3

905

906

907

Table 4 (Δ CHOP MA-10 550 kDa native complex)

Accession No.	Protein	MS/MS Score	No. of unique peptides
116089322	lon protease homolog, mitochondrial precursor	1681	25
51455	<i>heat shock protein 65</i>	1442	26
6753036	<i>aldehyde dehydrogenase, mitochondrial precursor</i>	1142	23
147900646	heat shock 70kDa protein 9	500	8
355762276	<i>78 kDa glucose-regulated protein</i>	471	9
63018	beta-actin	464	9
205632	<i>Na,K-ATPase alpha-1 subunit</i>	449	7
726095	<i>long-chain acyl-CoA dehydrogenase</i>	436	7
6679421	<i>NADPH--cytochrome P450 reductase</i>	409	7
4105605	<i>voltage dependent anion channel</i>	304	6
13385998	heat shock protein 75 kDa, mitochondrial precursor	271	4
31542438	cytochrome b5 type B precursor	228	5
5733504	<i>voltage-dependent anion channel VDAC3</i>	207	3
212653	Ca ²⁺ ATPase	192	4
158422314	molecular chaperone DnaK	153	3
124245114	<i>glucose-regulated protein 78</i>	130	4
56188	glyceraldehyde 3-phosphate-dehydrogenase	123	2
340546142	voltage-dependent anion channel 1	117	3
118596522	<i>acetaldehyde dehydrogenase</i>	83	2
348500154	isocitrate dehydrogenase [NAD] subunit alpha	73	1

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