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21 Abstract

22 Progesterone (P4) is a well-known steroid hormone that plays a key role in oocyte growth 23 and the maintenance of pregnancy in mammals, including cattle. Heat stress (HS) has an 24 adverse effect on P4 synthesis through an imbalance in the cellular redox status. We have 25 recently revealed that a standardized extract of Asparagus officinalis stem (EAS) increases 26 P4 through non-HS induction of heat shock protein 70 (HSP70) and a synergistic increase of 27 HSP70 by enhancing the intracellular redox balance, which was adversely affected by HS in 28 bovine granulosa cells (GCs). Bovine GCs collected from bovine ovarian follicles were 29 cultured at 38.5°C and 41°C for 12 h with or without 5 mg/mL EAS. After treatment, cells and 30 culture suppernatant were collected for the analysis. Enzyme-linked immunosorbent assay 31 (ELISA) was performed to detect in P4 level. Quantitative reverse-transcription polymerase 32 chain reaction (RT-qPCR) was used to detect expression of steroidogenesis related genes. 33 Fluorescence staining was used to detect mitochondrial activity and lipid droplet. 34 P4 level was increased by EAS treatment in association with increase in steroidogenic acute 35 regulatory protein (STAR), 3β-hydroxysteroid dehydrogenase (3β-HSD), mitochondrial 36 membrane activity and lipid droplet both under non-HS and HS conditions. Notably, 37 synergistic effect of EAS with HS co-treatment was observed to show a greater increase in 38 P4 synthesis when comparison with EAS treatment under non-HS condition. Furthermore, 39 inhibition of HSP70 significantly reduced EAS-induced P4 synthesis, mitochondrial activity

40 and synthesis of lipid droplets. These results suggest that P4 synthesis by EAS is mediated

41	by the steroidogenesis pathway via HSP70-regulated activation of STAR and 3β -HSD,
42	together with improved mitochondrial activity and lipid metabolism in bovine GCs. Moreover,
43	effect of EAS has a synergistic effect of with HSP70-regulated steroidogenesis pathway.
44	
45	Keywords: Bovine granulosa cells, EAS, HSP70, Progesterone, synergistic effect

47 **1. Introduction**

Progesterone (P4) is an important steroid hormone that plays a pivotal role in the establishment of uterine receptivity, oocyte maturation, and the maintenance of pregnancy [1,2]. In mammals, granulosa cells (GCs) play a fundamental role in the maturation and acquisition of developmental competence in bovine oocytes [3]. GCs differentiate into luteal cells that are responsible for P4 production [4]. Therefore, GCs are good models for inducing steroidogenesis under heat stress (HS) conditions to evaluate the importance of maintaining reproductive functions.

55 P4 synthesis in GCs is regulated by cholesterol transport into the mitochondria by 56 steroidogenesis enzymes, such as STAR, cytochrome P450 family 11 subfamily a member 57 1 (CYP11A1), and 3β-HSD [5,6]. Importantly, mitochondria play a central role in 58 steroidogenesis and are abundant in GCs [6,7]. Moreover, P4 synthesis in bovine GCs is 59 associated with lipid metabolism [8]. In addition, cholesterol from lipid droplets is one of the 60 available sources for P4 biosynthesis [7]. One of the main transcription factors that control 61 lipid metabolism by regulating enzymes required for cholesterol, fatty acid, triacylglycerol, 62 and phospholipid synthesis is SREBP-1 [9]. Thus, P4 synthesis in bovine GCs is regulated 63 by steroidogenesis, mitochondrial function, and lipid metabolism. However, a critical issue 64 that affects dairy farming is the reduction of reproductive performance in high-yielding dairy 65 cows [10]. The reproductive performance of dairy cows is decreased by increased body temperature [11,12] which decreases ovarian function, follicular growth, and corpus luteum
 function in summer.

High ambient temperature produces reactive oxygen species (ROS) with reduced P4 production in bovine GCs [13]. In addition, a negative effect of ROS on P4 levels has been reported in human follicular fluid [14]. Follicular fluid contains soluble factors secreted from GCs that maintain the intrafollicular redox status [15,16]. ROS induces redox status imbalance, which leads to cellular damage, alteration of cellular function, and physiological processes [17,18]. Thus, HS has an adverse impact on P4 synthesis owing to HS-induced toxic factors that disturb the redox imbalance in GCs.

Under stress conditions, cellular conditions are restored by the induction of heat shock proteins (HSPs), which act as molecular chaperones by folding and unfolding proteins while maintaining homeostasis of the cells [19]. Therefore, strengthening chaperone systems by inducing HSPs is one of the keys to regulating their functions in various biological pathways in ovarian functions, including P4 synthesis. However, as described above, HS is accompanied by the induction of ROS with harmful effects.

Recently, a standardized extract of *Asparagus officinalis* stem (EAS) was shown to induce HS-independent HSP70 in human promyelocytic leukemia cells [20]. In addition, we have revealed that EAS has a beneficial effect in inducing HSP70 under non-HS conditions and maintaining redox balance through ROS reduction and glutathione (GSH) synthesis in bovine GCs [21]. Moreover, EAS showed a synergistic increase in HSP70 expression and an

86	improvement in the intracellular redox status of bovine GCs, which was adversely affected
87	by HS [21]. The abundance regulation of EAS-induced HSP70 which affects redox balance
88	under both HS and non-HS conditions may compromise P4 synthesis in bovine GCs. In
89	addition, the effects of EAS on P4 biosynthesis under HS and non-HS conditions in bovine
90	GCs cells have not yet been investigated. In the present study, we investigated the effects
91	of EAS on P4 levels, steroidogenesis genes, mitochondrial function, and lipid metabolism
92	under non-HS and HS conditions.
93	
94	2. Materials and Methods
95	
96	Extract of Asparagus officinalis stem (EAS), which was extracted from asparagus (A.
97	officinalis L.), was produced following a previously described method [22] and commercially
98	available as a supplement (ETAS [®]).
99	
100	2.1 Collection and culture of bovine GCs
101	Bovine ovaries were obtained from a local abattoir and were transported within 4 hours to
102	the laboratory at 20°C. The ovaries were washed several times with a sterile saline solution.
103	Oocytes were collected from follicles (2-8 mm in diameter) without signs of atresia, using a
104	disposable 18-gauge needle attached to a 10-mL syringe. After cumulus-oocyte complexes
105	were picked up, the remaining follicular fluid with floating GCs cells was cultured in

Dulbecco's modified Eagle's medium (high glucose) (DMEM) (Wako, Osaka, Japan)
containing 5% fetal bovine serum (FBS), 0.06 g/L penicillin G potassium (Nacalai Tesque,
Kyoto, Japan), and 0.1 g/L streptomycin sulfate (Nacalai Tesque, Kyoto, Japan) at 38.5°C
under 5% CO₂ in air.

110 After overnight culture, the theca cell cluster layer was gently removed from the bottom of 111 the cell culture dishes. The cells remaining at the bottom of the dish were bovine GCs, which 112 were then washed with calcium- and magnesium-free phosphate-buffered saline (PBS) (-), 113 and cultured in 5% FBS in DMEM at 38.5°C under 5% CO2 in air. When the cells reached 114 confluence, they were washed with PBS (-) and dissociated from the substratum with PBS 115 (-) containing 0.05% trypsin and 0.53 mM Ethylenediaminetetraacetic acid (EDTA) for 2 min 116 at 38.5°C in a CO₂ incubator. After the addition of 5% FBS in DMEM to inactivate trypsin 117 activity, the separated cell suspension was centrifuged at $1,200 \times g$ for 3 min. To culture 118 these bovine GCs, 1 × 10⁵ cells/mL viable cells were seeded in each well of a 4-well dish 119 (Thermo Fisher Scientific) or 8-well slide chamber (Thermo Fisher Scientific) and cultured at 120 38.5° C under 5% CO₂ in air. After the cells reached 70% confluence, the medium was 121 replaced with 0.9 mL of DMEM in 5% FBS together with 0.1 mL of EAS solution in PBS (-), 122 and 0.1 mL of PBS (-) was added to the control group. The cells were then cultured at 38.5°C 123 or 41°C under 5% CO₂ atmosphere.

124

125 2.2 Experimental design

126 To investigate the effect of EAS on P4 production, the cells were cultured at 38.5°C for 12 127 h with 0.5, 1, 5, and 10 mg/mL of EAS. In addition to evaluating the effects of EAS on P4 128 synthesis under non-HS and HS conditions, cells were cultured at 38.5°C and 41°C for 12 h 129 with or without 5 mg/mL EAS. To confirm the role of EAS-induced HSP70 in P4 synthesis, 130 EAS-treated cells were exposed to 10 µM pifithrin-µ (2-phenylethynesulfonamide:PES) 131 (StressMarq Biosciences Inc., Victoria, Canada) under non-HS and HS conditions. After 132 culture, cells and culture medium were used for mRNA expression analysis, P4 133 measurement, and detection of lipid droplets and mitochondria.

134

135 2.3 P4 measurement

136 The concentrations of P4 in culture supernatants of bovine CGs were measured by enzyme-

137 linked immunosorbent assay (ELISA) using P4 measurement kits (ADI- 900- 011, Enzo Life

138 Sciences, USA). The intra- and inter-assay coefficients of variation for P4 were less than

139 10%. The aforementioned experiments were performed in duplicate.

140

141 2.4 RNA extraction and quantitative reverse-transcription polymerase chain reaction (RT-

142 *qPCR*)

Total RNA was extracted from the cells using ISOGEN II (Nippon Gene, Toyama, Japan)
 according to the manufacturer's instructions. The RNA concentration was measured using
 spectrophotometry (NanoDrop ND-2000; Thermo Fisher Scientific). All RNA samples were

146 stored at -80°C until use. Total RNA was reverse transcribed using ReverTra Ace qPCR RT 147 Master Mix with gDNA remover (TOYOBO Life Science, Osaka, Japan) with a thermal cycler 148 (Astec GeneAtlas Type G Thermal Cycler; ASTEC, Fukuoka, Japan). All cDNA samples were 149 stored at - 30°C until use for Quantitative PCR. Specific primers (Table S1) were designed 150 using Primer-BLAST (http://www.ncbi.nlm.nih.gov/tools/primer-blast/). qPCR was performed 151 using THUNDERBIRD SYBR qPCR Mix (Toyobo Life Science) at a final primer concentration 152 of 0.5 µM. To quantify mRNA expression levels, PCR was performed under the following 153 conditions with the LightCycler Nano (Roche Diagnostics, Basel, Switzerland): 1 cycle at 154 95°C for 30 s (denaturation), 45 cycles at 95°C for 10 s (denaturation), 55°C for 15 s (primer 155 annealing), and 72°C for 30 s (extension). Relative mRNA abundance was calculated using 156 the $\Delta\Delta$ Ct method with H2AFZ as a reference gene.

157

158 2.5 Immunodetection of HSP70

HSP70 immunofluorescence analysis was performed as previously described [21]. After GCs were cultured in 8-well slide chambers, they were fixed with 4% paraformaldehyde diluted with PBS (–), permeabilized with 0.2% Triton X-100 in PBS (–), blocked with 2% (w/v) BSA (Sigma-Aldrich) in PBS (–), and incubated at 4°C overnight with rabbit anti-human HSP70 polyclonal primary antibody (SPC-103; StressMarq Biosciences Inc., Victoria, Canada). After washing three times with PBS, the cells were incubated for 30 min with the secondary antibody Alexa Fluor 488 donkey anti-rabbit IgG (A21206; Thermo Fisher 166 Scientific). The 8-well slide chambers were then mounted with mounting solution (Fluoro-

167 KEEPER Anti fade Reagent, Non-Hardening Type with DAPI, Nacalai Tesque) and observed

168 under a fluorescence microscope (Leica Camera AG, Wetzlar, Germany).

169

170 2.6 Detection of Lipid droplet and mitochondrial activity

For the detection of lipid droplets, cells were washed with PBS (–) after culture and fixed with 4% PFA in PBS (–) for 10 min at room temperature. After washing the cells three times with PBS (–) for 5 min, they were treated with 1 μ g/mL Nile Red (Wako, Osaka, Japan) for 30 min at 38.5°C in a 5% CO₂ atmosphere.

175 To detect mitochondrial activity, cells were washed with PBS (-) after culture and treated

176 with 200 nM MitoTracker™ Green FM (Thermo Fischer Scientific) for 30 min at 38.5°C in a

177 5% CO₂ atmosphere.

Cells from both detections were then washed with PBS (–), and 10 µL of mounting solution (Nacalai Tesque) was added to the cells and covered with a cover glass. Fluorescence images were obtained using a fluorescence microscope (Leica, Wetzlar, Germany). Fluorescence intensity was quantified using ImageJ software v1.52A (National Institutes of Health; http://imagej.nih.gov/ij/). Corrected total cell fluorescence (CTCF) was carried out following a previously described formula [23]:

184 1. CTCF = integrated density (total area of selected cells × mean fluorescence of
 185 background readings).

186 2. CTCF per cell = CTCF/ Ncells

187 where "integrated density" is the integrated intensity of pixels for all cells in the image, total 188 cell area is the number of pixels of all cells, background fluorescence is the average mean 189 gray value of nearby regions containing no cells, and Ncells is the number of cells that was 190 measured by counting fluorescently labeled nuclei from images. 191 Images were deconvolved using Huygens Professional software (Hilversum, Netherlands) 192 193 2.7 Statistical analysis 194 Experimental data are shown as mean ± standard error of the mean (S.E.M.). Analysis of 195 variance (ANOVA) and Tukey's test were performed using R software (version 3.5.3; 196 https://www.r-project.org/). Probabilities of less than 5% (P < 0.05) indicate statistically 197 significant differences. 198 199 3. Results

200 3.1 Effects of EAS on P4 production

201 We investigated the effects of various doses of EAS on P4 secretion in bovine GCs. As

- shown in Figure 1, we found that EAS significantly (p < 0.001) promoted P4 levels peaking
- at 5 mg/mL. Collectively, EAS showed a dose-dependent induction of P4 production.

205 3.2 Effects of EAS on P4 production and steroidogenic and lipid metabolism gene expression

In our previous study, we showed that EAS induces HSP70 and improves redox balance in

206 in bovine GCs under non-HS and HS conditions

207

both non-HS and HS conditions [21]. In the present study, we investigated whether EAS affects P4 synthesis in both environments, together with the expression of steroidogenesisrelated genes in EAS-treated bovine GCs, under non-HS and HS conditions. As shown in Figure 2A, P4 levels were significantly (p < 0.001) increased by EAS compared with the

212 untreated control. However, P4 levels in HS-treated cells tended to be lower than that for the

213 control cells (p < 0.1) (Figure 2A). Interestingly, P4 levels were significantly increased (p < 0.1)

214 0.001) by EAS under HS conditions (Figure 2A).

215 Furthermore, STAR and 3β-HSD mRNA expression levels were similar to P4 levels in the

216 EAS and EAS+HS groups (p < 0.001) (Figure 2B, 2C). In contrast, EAS did not affect the

217 expression of CYP11A1, progesterone receptor, or estrogen receptor under either non-HS

or HS conditions (p > 0.05) (Figure 2D and Figure S1).

219 Moreover, SREBP-1 expression was significantly increased in both the EAS and EAS+HS

groups, similar to the pattern of P4 levels and expression of steroidogenic genes (p < 0.05)

221 (Figure 2E).

222 These results confirmed that EAS-induced P4 production is affected by the enhancement of

223 STAR, 3β-HSD, and SREBP-1, and it has the greatest effect on P4 synthesis in bovine GCs

under HS conditions. Notably, HS treatment had a synergistic effect with EAS on the increase
in *STAR*, *3β-HSD*, and *SREBP-1*.

226

3.3 Effects of EAS on lipid metabolism and mitochondrial activity in bovine GCs under nonHS and HS conditions

To clarify the effects of EAS on P4 levels, steroidogenesis, and lipid metabolism gene expression, we examined lipid synthesis and mitochondrial function in EAS-treated cells under non-HS and HS conditions. As shown in Figure 3A, EAS increased lipid droplets and mitochondrial activity under non-HS and HS conditions. Remarkably, the EAS-induced HSP70 protein was expressed in both the nucleus and cytoplasm, but HS-induced HSP70 was localized mainly in the nucleus (Figure 3A).

235 Next, we deconvoluted the acquired lipid metabolism and mitochondrial activity images. 236 Deconvolution of the images efficiently removes background noise pixels in lipid droplets and 237 mitochondrial images of African green monkey kidney fibroblast cells [24]. After 238 deconvolution, the lipid droplets and mitochondrial structure appeared to be assembled from 239 multiple particles (Figure 3A). Notably, the localization of lipid droplets and mitochondrial 240 function were mainly observed in the cytosol around the nucleus (Figure 3A). The results 241 show that EAS increased lipid droplets and mitochondrial particles under non-HS and HS 242 conditions.

Similar to P4 synthesis, the fluorescence levels of lipid droplets and mitochondria in the EAS+HS group were significantly (p < 0.001) higher than those in the other treatment groups (Figure 3B, C). HS significantly (p < 0.05) increased the number of lipid droplets but significantly (p < 0.01) reduced mitochondrial activity (Figure 3B, C). These results show that EAS and its synergistic effect with HS increased lipid metabolism and mitochondrial activity, and these results may be related to EAS-induced HSP70.

249

250 3.4 Inhibition of HSP70 attenuates P4 synthesis, lipid metabolism, and mitochondrial activity

251 in EAS-treated bovine GCs under non-HS and HS conditions

252 We hypothesized that EAS-induced HSP70 may be involved in the effects of EAS on P4 253 synthesis, lipid droplet formation, and mitochondrial function in bovine GCs under non-HS 254 and HS conditions. To test this hypothesis, we evaluated the effects of HSP70 inhibition on 255 P4 synthesis, steroidogenic gene expression, lipid synthesis, and mitochondrial activity in 256 EAS-treated cells under both non-HS and HS conditions. As expected, PES significantly 257 decreased EAS-induced P4 synthesis related to reduced P4, STAR, 3β-HSD, and SREBP-258 1 levels under both non-HS and HS conditions (p < 0.05) (Figure 4A, B, C, E). However, an 259 HSP70 inhibitor did not affect CYP11A1 expression (p > 0.05) (Figure 4D). 260 In addition, HSP70 blockage inhibited the EAS-mediated upregulation of fluorescent levels 261 of lipid droplets and mitochondria, whereas HSP70 expression was unchanged (Figure 5).

262 Consistently, PES suppressed lipid synthesis and mitochondrial activity in EAS-treated

bovine GCs under non-HS and HS conditions (p < 0.05) (Figure 6). Taken together, these data suggest that EAS-induced P4 synthesis is regulated by EAS-induced HSP70.

265

4. Discussion

In the present study, we confirmed that EAS induces HSP70-mediated P4 synthesis through steroidogenic genes associated with the induction of lipid metabolism and mitochondrial activity in bovine GCs under both non-HS and HS conditions. Notably, EAS treatment under HS treatment synergistically increased P4 synthesis with increased expression of *STAR*, 3β -*HSD*, and *SREBP-1*, lipid metabolism, and mitochondrial activity.

272 The present results show that P4 levels in bovine GCs were highest when treated with 5 273 mg/mL EAS. This result was similar to that of our previous study in which bovine GCs were 274 treated with or without 5 mg/mL EAS [21]. EAS exerts beneficial effects during stress and 275 improves sleep quality in humans [25]. Other studies have revealed beneficial effects of EAS 276 on brain cell function [26]. Recent evidence has suggested that the central nervous system 277 can synthesize steroid hormones from cholesterol [27]. P4 synthesis is initiated by 278 cholesterol import into the mitochondria by STAR, after which CYP11A1 transforms 279 cholesterol to pregnenolone, which is converted into P4 by 3β -HSD [28]. In GCs, P4 binds to 280 the progesterone receptor and facilitates its activity in gene transcription and cellular 281 responses [29]. Under in vitro conditions, bovine GCs are good models for P4 studies 282 because of luteinization of this cell line, which involves the switch from the production of 283 estrogens to P4 [30]. In the present study, EAS strongly increased P4 levels and the 284 expression of STAR and 3\beta-HSD, but did not affect the expression of CYP11A1, 285 progesterone receptor, and estrogen receptor. These results suggest that EAS regulates P4 286 synthesis through STAR and 3β -HSD pathways in the mitochondria of bovine GCs. In 287 addition, EAS enhanced P4 levels but did not affect P4 related signal pathway of GCs through 288 the progesterone receptor in an autocrine or paracrine manner. Furthermore, HS can 289 synergistically promote this pathway with EAS treatment, resulting in promotion of P4 290 synthesis. Importantly, mitochondria play a central role in steroidogenesis enzymatic activity 291 [6,7]. Surprisingly, in our results, EAS increased mitochondrial activity in response to HS, 292 despite the fact that HS itself decreased mitochondrial activity.

293 Generally, HS in the summer season is known to cause a decrease in ovarian function, with 294 reduced P4 levels and corpus luteum size [31–33]. In mammalian cells, lipid droplets are the 295 cholesterol preservation sites for steroid hormones [34]. In the fat tissue of Japanese black 296 cattle, SREBP-1 affects lipid metabolism [35]. Our present results show that lipid droplets 297 and SREBP-1 were induced by both EAS and HS at the same level, and the highest 298 expression levels were observed after co-treatment with EAS and HS. Collectively, EAS-299 induced P4 synthesis through enhancement of STAR, 3β-HSD enzyme activity, 300 mitochondrial activity, and lipid metabolism in bovine GCs. In addition, significant effect of 301 EAS and HS co-treatment on P4 synthesis was also found in the present study.

In our previous study, EAS was shown to enhance redox balance through the reduction of
 ROS levels under ordinary non-HS conditions, and, surprisingly, EAS had a significant effect
 on HSP70 induction and ROS reduction in bovine GCs even under HS conditions compared
 with the increased ROS and decreased GSH under HS conditions without EAS [21].

306 In a previous study, STAR, a key enzyme in P4 synthesis, was shown to be inhibited by ROS 307 in rat Leydig cells [36]. ROS also blocks P4 synthesis through CYP11A1 and 3β-HSD in 308 human granulosa luteal cells [37]. In bovine GCs, HS-induced ROS reduce STAR and 309 CYP11A1 gene expression [13]. Hydrogen peroxide, the most popular ROS, decreases P4 310 levels, STAR, and 3β -HSD gene expression in bovine GCs [38]. In addition, high 311 concentrations of ROS negatively affect mitochondria in mouse GCs cells [39]. Therefore, 312 redox balance might also be important in regulating steroidogenesis-related genes and 313 mitochondrial activity in bovine GCs.

In addition, lipid droplets are markers of cellular stress, and they play an important role in maintaining redox status under stress conditions [40]. Under stress conditions, lipid droplets sequester toxic lipids and delay the release of lipids, and this lipid droplet biosynthesis maintains redox balance [40]. Thus, EAS-treated GCs in this study, while maintaining redox balance, possibly caused the stabilization of lipid droplets as a source of cholesterol compared to non-EAS-treated cells. Collectively, the enhancement of P4 synthesis by EAS treatment while maintaining redox status depends on the ROS levels in bovine GCs.

321 In the present study, HS induced lipid droplet formation and reduced mitochondrial activity in 322 bovine GCs. Our previous study suggested that ROS and ROS-induced DNA damage are 323 detected mainly in the nuclei of bovine GCs under HS conditions [21]. In bovine somatic cell 324 nuclear transfer embryos, mitochondrial and DNA damage are induced by an increase in 325 ROS [41]. Thus, impairment of DNA damage in the nucleus reduces mitochondrial function 326 in HS bovine GCs and affects gene expression. Previous studies have shown that lipid 327 droplets have dynamic functions in cellular metabolism [42]. In mice, mitochondrial damage 328 is caused by hypoxia-induced formation of lipid droplets [43]. Thus, the high levels of lipid 329 droplets in bovine GCs induced by HS can be explained by mitochondrial dysfunction. 330 Collectively, in our results, HS-induced ROS damage in the nucleus impaired mitochondrial 331 function, resulting in the accumulation of lipid droplets in bovine GCs. EAS and EAS plus HS 332 treatment also induced lipid droplet formation, and these results are related with expression 333 of HSP70 that will be discussed in more details below. In our previous study, overall data 334 confirmed that the beneficial effects of EAS-induced HSP70 are mediated by regulating redox 335 balance and protecting cells from HS-related harmful effects in bovine GCs [21]. 336 To show the relationship between HSP70, the steroidogenesis pathway, and mitochondrial 337 function, we inhibited HSP70 activity. Inhibition of HSP70 by PES did not affect the

expression of HSP70 protein; however, HSP70 inhibition clearly reduced mitochondrial activity and lipid droplet synthesis. Previous studies have shown that PES disrupts the cochaperone and substrate proteins of HSP70 without affecting HSP70 expression; thus, PES 341 inhibits the function of HSP70 protein in multiple cell signaling pathways [44]. In the present 342 study, PES strongly inhibited P4 synthesis, expression of STAR, 3β -HSD, and SREBP-1, 343 mitochondrial activity, and lipid droplet synthesis in bovine GCs, even in the presence of EAS, 344 under both non-HS and HS conditions. Therefore, these data strongly support the hypothesis 345 that an increase in EAS-induced HSP70 promotes P4 synthesis. In addition, HSP70 346 localization was detected in both the nucleus and cytoplasm of EAS-treated cells, particularly 347 under HS conditions. In mammalian cells, HSP70 has chaperone functions in both the 348 nucleus and cytoplasm [45,46].

349 In our previous study, EAS-induced HSP70 reduces DNA damage by reducing ROS, which 350 are mainly generated in the nucleus in bovine GCs [21]. Modulation of redox balance-351 regulated mitochondrial activity and lipid droplets reduces ROS damage in the nuclei of 352 bovine GCs. In addition, the transcription of steroidogenesis enzymes mainly occurs in the 353 nucleus [47]. Therefore, EAS-induced HSP70 balances the redox status to improve 354 steroidogenesis enzyme activity, mitochondrial function, and lipid synthesis in bovine GCs. 355 On the other hand, synthetic chemical chaperones increase 3β -HSD metabolic activity in the 356 inner mitochondrial membrane by promoting the folding speed of 3β-HSD from the native to 357 the active state [48]. In rat adipocyte lipid droplets, HSP70 stimulation is involved in stabilizing 358 the droplet monolayer, protein folding, and mobilization of nascent proteins to the lipid 359 droplets [49]. The size of lipid droplets and fat accumulation are increased by HSP70 through 360 the enhancement of SREBP-1 in mouse liver cells [50]. Therefore, EAS-induced HSP70

361 enhances the accumulation of lipid droplets and 3β-HSD activity in the mitochondria in the 362 cytoplasm with the chaperone function of bovine GCs. In the present study, promotion effect 363 of EAS on P4 synthesis via steroidogenesis pathway, mitochondrial functions, lipid synthesis 364 with synergistic augmentation with HS. These results suggest the EAS is a potential tool for 365 improving the reproductive functions by regulating of intracellular redox and molecular 366 chaperone. Recent study has revealed that the HSP inducer in the EAS was purified and 367 identified as a mixture of Asparagus-Derived Proline-Containing 3-Alkyldiketopiperazines 368 (Asparaprolines) [20]. However, evaluation of such components on the detailed cellular 369 functions is not clarified. Further study will clarify the beneficial effect on cellular functions 370 and regulation of reproductive performance.

371

5.Conclusion

373 P4 synthesis by EAS is mediated by the steroidogenesis pathway via HSP70-regulated 374 activation of STAR and 3β-HSD, together with improved mitochondrial activity and lipid 375 metabolism in bovine GCs. Moreover, EAS has synergistic effect of with HSP70-regulated 376 steroidogenesis pathway. These findings provide a basis for the application of EAS to 377 improve reproductive functions in mammals, including cattle, in combination with HS 378 treatment.

379

380 Supplementary materials:

381	Table S1: Primer sequences used for qRT-PCR
382	Figure S1: Effects of EAS on progesterone receptor and estrogen receptor in bovine GCs
383	cells under non-HS and HS conditions
384	
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393	article.
394	
395	Ethics approval and consent to participate: Not Applicable . All experiments were carried
396	out by cultured cells collected from ovaries at slaughter house.
397	Competing interests: The authors declare no conflicts of interest.
398	

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591 Figure legends

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593 Figure 1. Effect of various concentrations of EAS on P4 production in bovine GCs.

- 594 Cells were subjected to various concentrations of EAS (0.5, 1, 5, and 10 mg/mL) for 12 h at 595 38.5° C (n = 5). Detection of P4 secretion was performed with ELISA in cell supernatants. 596 Data are shown as the mean ± S.E.M, a vs. b (p < 0.001), a vs. c (p < 0.05), and b vs. c (p
- 597 < 0.05).
- 598

599 Figure 2. Effect of EAS on P4 synthesis and expression of steroidogenic genes in 600 bovine GCs under non-HS and HS conditions.

- 601 Cells were treated with or without 5 mg/mL of EAS at 38.5°C (control, EAS) or 41°C (HS, HS
- 602 + EAS) for 12 h. Detection of P4 secretion was performed with ELISA in cell supernatants (A,
- 603 n = 3). The mRNA expression levels of STAR, 3 β -HSD, CYP11A1, and SREBP-1 (B–E,
- 604 respectively, n = 5) are presented relative to H2AFZ as a reference gene. Data are shown
- 605 as the mean ± S.E.M, (A) a vs. b (p < 0.001), a vs. c (p < 0.001), b vs. c (p < 0.05), (B) a vs.

608

609 Figure 3. Effect of EAS on the synthesis of lipid droplets and mitochondrial activity in

610 $\,$ bovine GCs under non-HS and HS conditions.

611	Cells were treated with or without 5 mg/mL of EAS at 38.5°C (control, EAS) or at 41°C (HS,
612	HS + EAS) for 12 h. Fluorescence staining and enlarged image of mitochondria, lipid droplets,
613	and HSP70 are shown (A). Scale bar = 50 μ m; original magnification = 63×. The fluorescence
614	intensity of mitochondria and lipid droplets was analyzed using CTCF (B and C, respectively,
615	n = 5). Data are shown as the mean \pm S.E.M, (B) a vs. b (p < 0.05), a vs. c (p < 0.001), b vs.
616	c (p < 0.001), (C) a vs. b (p < 0.01), a vs. c (p < 0.01), a vs. d (p < 0.001), b vs. c (p < 0.001),
617	b vs. d (p < 0.001), c vs. d (p < 0.001).
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619	Figure 4. Effect of HSP70 inhibition on EAS-induced P4 synthesis and expression of
620	steroidogenic genes in bovine GCs under non-HS and HS conditions.
621	Cells were treated with or without 10 μ M PES to 38.5°C (control, EAS, EAS + PES) or at
622	41°C (HS, HS + EAS, HS + EAS + PES) for 12 h. Detection of P4 secretion was performed
623	
	with ELISA in cell supernatants (A, n = 3). The mRNA expression levels of STAR, 3β -HSD,
624	with ELISA in cell supernatants (A, n = 3). The mRNA expression levels of STAR, 3β -HSD, CYP11A1, and SREBP-1 (B–E, respectively, n = 5) are presented relative to H2AFZ as a
624 625	with ELISA in cell supernatants (A, n = 3). The mRNA expression levels of STAR, 3 β -HSD, CYP11A1, and SREBP-1 (B–E, respectively, n = 5) are presented relative to H2AFZ as a reference gene. Data are shown as the mean ± S.E.M, (A) a vs. b (p<0.05), a vs. c
624 625 626	with ELISA in cell supernatants (A, n = 3). The mRNA expression levels of STAR, 3 β -HSD, CYP11A1, and SREBP-1 (B–E, respectively, n = 5) are presented relative to H2AFZ as a reference gene. Data are shown as the mean ± S.E.M, (A) a vs. b (p<0.05), a vs. c (p<0.01), and b vs. c (p<0.01), (B) a vs. b (p<0.001), a vs. c (p<0.001), and b vs. c
624 625 626 627	with ELISA in cell supernatants (A, n = 3). The mRNA expression levels of STAR, 3β-HSD, CYP11A1, and SREBP-1 (B–E, respectively, n = 5) are presented relative to H2AFZ as a reference gene. Data are shown as the mean \pm S.E.M, (A) a vs. b (p<0.05), a vs. c (p<0.01), and b vs. c (p<0.01), (B) a vs. b (p<0.001), a vs. c (p<0.001), and b vs. c (p<0.05); (C) a vs. b (p<0.05); a vs. c (p<0.001), b vs. c (p<0.001), and (E) a vs. b (p<0.05); a vs. c (p<0.001), b vs. c (p<0.001), and (E) a vs. b (p<0.05); a vs. c (p<0.001), b vs. c (p<0.001), and (E) a vs. b (p<0.05); a vs. c (p<0.001), b vs. c (p<0.001), and (E) a vs. b (p<0.05); a vs. c (p<0.001), b vs. c (p<0.001), and (E) a vs. b (p<0.05); a vs. c (p<0.001), b vs. c (p<0.001), and (E) a vs. b (p<0.05); a vs. b (p<0.001), b vs. c (p<0.001), and (E) a vs. b (p<0.05); a vs. b (p<0.05); a vs. c (p<0.001), b vs. c (p<0.001), and (E) a vs. b (p<0.05); a vs. b (p<0.05); a vs. c (p<0.001), b vs. c (p<0.001), and (E) a vs. b (p<0.05); a vs. b (p<0.05); a vs. c (p<0.001), b vs. c (p<0.001), and (E) a vs. b (p<0.05); a vs. b (p<0.05); a vs. c (p<0.001), b vs. c (p<0.001), a vs. b (p<0.05); a vs. b (p<0.05); a vs. c (p<0.001), b vs. c (p<0.001), a vs. b (p<0.05);
 624 625 626 627 628 	with ELISA in cell supernatants (A, n = 3). The mRNA expression levels of STAR, 3 β -HSD, CYP11A1, and SREBP-1 (B–E, respectively, n = 5) are presented relative to H2AFZ as a reference gene. Data are shown as the mean ± S.E.M, (A) a vs. b (p<0.05), a vs. c (p<0.001), and b vs. c (p<0.01), (B) a vs. b (p<0.001), a vs. c (p<0.001), and b vs. c (p<0.05); (C) a vs. b (p<0.05); a vs. c (p<0.001), b vs. c (p<0.001), b vs. c (p<0.001), and (E) a vs. b (p<0.001), a vs. c (p<0.001), a vs. c (p<0.001), b vs. c (p<0.001).

630 Figure 5. HSP70 inhibitor reversed EAS-induced lipid metabolism and mitochondrial

631 activity in bovine GCs under non-HS and HS conditions.

- Cells were treated with or without 10 μ M PES at 38.5°C (control, EAS, EAS + PES) or at 41°C (HS, HS + EAS, HS + EAS + PES) for 12 h. Fluorescence staining images of mitochondria, lipid droplets, and HSP70 are shown. Scale bar = 50 μ m; original magnification = 63×.
- 636

637 Figure 6. Fluorescence intensity of lipid droplets and mitochondrial activity in bovine

638 GCs treated with EAS under non-HS and HS conditions.

639 The fluorescence intensity of lipid droplets and mitochondrial activity in cells treated with EAS

- 640 under non-HS and HS conditions was analyzed by CTCF (A and B, respectively, n = 5). Data
- are shown as the mean \pm S.E.M, (A) a vs. b (p < 0.05), a vs. c (p < 0.001), and b vs. c
- 642 (p < 0.05), (B) a vs. b (p < 0.05), a vs. c (p < 0.05), a vs. d (p < 0.001), b vs. c (p < 0.001), b
- 643 vs. d (p < 0.05), c vs. d (p < 0.001).
- 644

Highlights

- EAS increases progesterone synthesis by increasing of steroidogenesis-related genes
- EAS increases lipid synthesis and mitochondrial activity
- Synnergistic effect of EAS and heat shock on progesterone synthesis
- HSP70 inhibition decreases EAS-stimulated progesterone synthesis pathway

Graphical abstract



Figure 1



Figure 2



Figure 3



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Figure 4







Figure 5





