Identification of crucial genes and metabolites regulating the eggshell brownness in chicken

Jing Yang¹, Zhiqiong Mao², Xiqiong Wang¹, Jingjie Zhuang¹, Sijia Gong¹, Zhouyang Gao¹, Guiyun Xu¹, Ning Yang¹ and Congjiao Sun¹

Abstract

Background: Protoporphyrin IX (Pp IX) is the primary pigment for brown eggshells. However, the regulatory mechanisms directing Pp IX synthesis, transport, and genetic regulation during eggshell calcification in chickens remain obscure. In this study, we investigated the mechanism of brown eggshell formation at different times following oviposition, using White Leghorn hens (WS group), Rhode Island Red light brown eggshell line hens (LBS group) and Rhode Island Red dark brown eggshell line hens (DBS group).

Results: At 4, 16 and 22 h following oviposition, Pp IX concentrations in LBS and DBS groups were significantly higher in shell glands than in liver (P < 0.05). Pp IX concentrations in shell glands of LBS and DBS groups at 16 and 22 h following oviposition were significantly higher than WS group (P < 0.05). In comparative transcriptome analysis, δ-aminolevulinate synthase 1 (ALAS1), solute carrier family 25 member 38 (SLC25A38), ATP binding cassette subfamily G member 2 (ABCG2) and feline leukemia virus subgroup C cellular receptor 1 (FLVCR1), which were associated with Pp IX synthesis, were identified as differentially expressed genes (DEGs). RT-qPCR results showed that the expression level of ALAS1 in shell glands was significantly higher in DBS group than in WS group at 16 and 22 h following oviposition (P < 0.05). In addition, four single nucleotide polymorphisms (SNPs) in ALAS1 gene that were significantly associated with eggshell brownness were identified. By identifying the differential metabolites in LBS and DBS groups, we found 11-hydroxy-E4-neuroprostane in shell glands and 15-dehydro-prostaglandin E1(1-) and prostaglandin G2 2-glyceryl ester in uterine fluid were related to eggshell pigment secretion.

Conclusions: In this study, the regulatory mechanisms of eggshell brownness were studied comprehensively by different eggshell color and time following oviposition. Results show that Pp IX is synthesized de novo and stored in shell gland, and ALAS1 is a key gene regulating Pp IX synthesis in the shell gland. We found three transporters in Pp IX pathway and three metabolites in shell glands and uterine fluid that may influence eggshell browning.

Keywords: Protoporphyrin IX, Brown eggshell, δ-aminolevulinate synthase 1, Shell gland, Oviposition

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Numerous studies have shown that eggshell brownness is regulated by synthases and transporters in the Pp IX/heme synthesis pathway. *ALAS1* is the target gene for the regulation of Pp IX concentration in eggshells [19, 23]. *ABCG2* has been shown to regulate ALA-mediated Pp IX levels by exporting Pp IX from the mitochondria to the cytoplasm [24]. Molecular experiments also show that coproporphyrinogen oxidase (*CPOX*) is expressed at higher levels in brown eggshell hens, but *FECH* is expressed at higher levels in white and pink eggshell hens [25]. Additionally, an iTRAQ-based quantitative proteomic analysis by Li et al. [26] revealed that proteins which transport Pp IX and iron into the mitochondrial matrix where heme is synthesized were upregulated and proteins that promote extracellular transport of Pp IX were downregulated in the shell glands of hens laying light brown eggs compared to hens laying dark brown eggs.

Existing studies on the mechanism of eggshell brownness, however, have only considered eggshell brownness from the perspective of the brownness intensity or time following oviposition, without comprehensive consideration of these two factors. Here, we held that combining these two perspectives to reveal the intrinsic driving impact of eggshell brownness was indeed necessary and of great value. Therefore, in the current study, hens with white, light brown and dark brown eggshells were utilized to explore the synthesis sites of Pp IX, and regulatory mechanism of brown eggshell formation at different stages of calcification. These findings may help shed light on the mechanism underlying brown pigmentation in eggshell.

**Results**

**Influence of eggshell brownness selection on egg quality**

To investigate the impact of eggshell brownness on egg quality, we measured egg weight and eggshell-related traits of light brown eggshell (LBS) line and dark brown eggshell (DBS) line, which have been bidirectionally selected for six generations according to their eggshell brownness. The results demonstrated that the DBS line had significantly darker eggshells than the LBS line (\(P < 0.05\), Table 1). The egg weight, eggshell strength, and eggshell weight of the LBS line eggs were significantly higher than those of the DBS line eggs (\(P < 0.05\), Table 1), however, there was no significant difference in eggshell thickness and eggshell proportion.

**Pp IX Concentrations in liver and shell glands**

Eggshell coloring occurs in accordance with eggshell calcification. At 4h following oviposition (4H), egg has not yet entered the shell gland, hence eggshell calcification does not begin. At 16h following oviposition (16H), the
egg undergoes a rapid calcification process in the shell gland, but pigment deposition does not initiate at this time. At 22 h following oviposition (22H), eggshell calcification reaches its terminal stage, and pigment deposition is extensive within shell gland (Additional file 1: Fig. S1).

To determine the synthesis and storage site of Pp IX, we measured Pp IX concentrations in liver and shell glands at 4H, 16H and 22H. The results showed that the concentration of Pp IX in the shell gland was significantly higher than that in the liver at all three time points in both the LBS and DBS group ($P<0.01$ or $P<0.05$, Fig. 1A and B). To clarify the synthesis pattern, white leghorn chickens that lay eggs with white eggshells (WS) were used as a control group. Figure 1C shows that the concentration of Pp IX in shell glands of the WS group was very low at all three time points (23.80–0.35 nmol/g). Nevertheless, the concentration of Pp IX in the shell glands of LBS and DBS groups reached their highest level at 16H ($P<0.05$), especially in the DBS group (84.92 nmol/g). In the liver, there was no significant difference in the concentration of Pp IX among WS, LBS, and DBS groups at any of the three time points ($P>0.05$, Fig. 1D).

Identification of differentially expressed genes from RNA-sequencing data

The elevated concentrations of Pp IX in shell glands indicates that shell gland plays a crucial role in Pp IX production and eggshell coloring. Therefore, RNA-sequencing (RNA-seq) was performed on 54 shell gland samples...
from WS, LBS and DBS groups at three time points to identify key genes determining eggshell brownness. After quality control, the number of clean reads produced per sample ranged from 39,733,000 to 55,987,008. At 16H and 22H, the concentration of Pp IX in the shell glands of the DBS groups was much higher than that at 4H; therefore, the DEGs at 16H or 22H versus 4H in DBS group deserve our attention (Fig. 2A and B), and we believe that the regulation of Pp IX formation by gene expression was positive, so the upregulated genes in DBS group shell glands at 16H or 22H compared with 4H were the focus of our analysis. However, the process of Pp IX synthesis, storage, and secretion has a large time overlap with eggshell calcification. In this case, with the WS group that produces white shelled eggs as a control, differentially expressed genes only related to eggshell calcification and not brownness can be identified and separated from those influencing brownness. Using this strategy, 284 upregulated genes at 16H (Gene List 1, Fig. 3A, Additional file 2: Table S1) and 396 upregulated genes at 22H were identified from DBS group shell glands (Gene List 2, Fig. 3B, Additional file 3: Table S2).

At 16H or 22H that belong to eggshell calcification stage, the DEGs between WS group and DBS group deserve attention (Fig. 2C and D). Similarly, we believe that the upregulated genes in shell glands in DBS group are most likely involved in regulating eggshell coloration compared with in WS group. Meanwhile, the eggshell brownness and Pp IX concentration of LBS group was between that of WS and DBS groups, therefore, we applied a criterion to ensure that the gene expression level of LBS group was between WS and DBS groups while screening the significantly expressed genes. Thus, we obtained 312 upregulated genes in the DBS group compared to the WS group at 16H (Gene List 3, Fig. 3C, Additional file 4: Table S3) and 595 upregulated genes at 22H (Gene List 4, Fig. 3D, Additional file 5: Table S4). At 4H, 16H and 22H, there were only 13, 39 and 19 DEGs between LBS and DBS groups, respectively, and these genes were not found to be associated with eggshell brownness.

Functional annotation of differentially expressed genes
To verify the biological functions of upregulated DEGs, we merged Gene Lists 1 and 3 (upregulated genes at 16H) and Gene Lists 2 and 4 (upregulated genes at 22H) for GO and KEGG analysis, respectively. For GO enrichment analysis of DEGs at 16H, signal transduction [GO:0007165], integral component of membrane [GO:0016021] and calcium ion binding [GO:0005509] were the top enriched terms for biological process, cellular component, and molecular function, respectively (Gene List 1 and 3) (Fig. 4A). For upregulated genes at 22H (Gene List 2 and 4), negative regulation of transcription from RNA polymerase II promoter [GO:0000122], integral component of membrane [GO:0016021] and transcriptional activator activity, RNA polymerase II transcription regulatory region sequence-specific binding [GO:0001228] were the corresponding top enriched terms, respectively (Fig. 4B). According to the KEGG analysis, upregulated DEGs at 16H were significantly enriched in five pathways including focal adhesion [gga04510], cytokine-cytokine receptor interaction [gga04060], glycosphingolipid biosynthesis - lacto and neolacto series [gga00601], ECM-receptor interaction [gga04512], and the MAPK signaling pathway [gga04010] (P < 0.05, Fig. 5A). Metabolic pathways [gga01000], glycerolipid metabolism [gga00561], the NOD-like receptor signaling pathway [gga04621], and the c-type lectin receptor signaling pathway [gga04625] were significantly enriched for upregulated genes at 22H (P < 0.05, Fig. 5B).

Identification of candidate genes
We screened genes for their relation to eggshell pigment from the above DEG lists, wherein ALAS1 was found in both Gene List 3 and 4. Furthermore, the expression level of ALAS1 in DBS group was nearly five times that in WS group and nearly twice that in LBS group at 4H. FLVCR1, a gene associated with heme export, was found in both Gene List 2 and 4, and its expression in the DBS group was highest at 22H among the three time points in this study. SLC25A38, encoding a protein responsible for transporting glycine from the cytoplasm to the mitochondria, was found in Gene List 2. In Gene List 3, we found ABCG2, which can transport Pp IX out of the cell, but found no significant difference in the expression level of ABCG2 at any of the three time points following oviposition in DBS group eggs. Detailed information of these four candidate genes is shown in Table 2.

Key role of ALAS1 in eggshell brownness verified at transcriptional and genetic levels
We compared the expression of ALAS1 in the shell glands of WS group and DBS groups (Fig. 6A), as well as in the shell gland and liver of the DBS group (Fig. 6B). The RT-qPCR results indicated that the expression level of ALAS1 was significantly higher in DBS group than in the WS group at 16H and 22H (P < 0.01 and P < 0.05, Fig. 6A), which was consistent with the RNA-seq results. The expression level of ALAS1 was significantly higher in the shell gland than in the liver at 16H in the DBS group (P < 0.01, Fig. 6B), but there was no significant difference at 22H (P > 0.05, Fig. 6B). ALAS1 is the rate-limiting enzyme in the Pp IX biosynthetic pathway. Thus, we compared the expression of other synthase genes in this pathway.
between WS and DBS groups based on RNA-seq data, including aminolevulinate dehydratase (ALAD), hydroxymethylbilane synthase (HMBS), uroporphyrinogen III synthase (UROS), uroporphyrinogen decarboxylase (UROD), coproporphyrinogen oxidase (CPOX), and protoporphyrinogen oxidase (PPOX). Results showed that there were no significant differences in gene expression between WS and DBS groups at 16H and 22H except for ALAS1 ($P > 0.05$, Fig. 6C and D).

To further verify the association between gene expression variations in the ALAS1 gene and eggshell brownness, the RNA-seq data was used for SNP calling and genotype extraction. Chicken ALAS1 is located at chromosome 12 and spans 7169 bp. Seven SNPs in ALAS1
gene were significantly associated with eggshell brownness ($P_{FDR} < 0.05$, Table 3). All of the seven SNPs were synonymous mutations, among which four were novel while the other three have already been reported in dbSNP database (https://www.ncbi.nlm.nih.gov/snp/).

Differential metabolites between LBS and DBS groups

Lots of genes were identified for their differential expression in shell glands of hens laying brown-shell eggs versus white-shell eggs in the current study. However, between light and dark brown eggshell groups, only a few related...
genes were found at the transcriptome level to account for their pigmented eggshell. Therefore, we utilized metabolomics approaches to investigate metabolites in the shell gland and uterine fluid that may regulate the brownness of eggshells. In the shell gland, 188 and 44 differential metabolites were identified between the LBS and DBS groups at 6H and 22H (Additional files 6 and 7: Tables S5 and S6), respectively. In the uterine fluid, the corresponding numbers of differential metabolites were 84 and 406 (Additional files 8 and 9: Tables S7 and S8), respectively. According to the results of annotation and literature searches, four metabolites were considered as candidate molecules associated with eggshell brownness (Table 4).

The concentration of Pp IX in shell glands of the DBS group was significantly higher than that in the LBS group.

**Fig. 5** KEGG annotation of upregulated genes. **A** KEGG annotation of upregulated DEGs at 16H (Gene List 1 and Gene List 3). **B** KEGG annotation of upregulated DEGs at 22H (Gene List 2 and Gene List 4).
at 6H, and we also compared Pp IX concentrations in uterine fluid at 6H and 22H in DBS group and found no significant differences. In addition, 11-hydroxy-E4-neuroprostane was upregulated in shell glands of the DBS group at 6H; 15-dehydro-prostaglandin E1(1-) and prostaglandin G2 2-glyceryl ester were upregulated in uterine fluid of the DBS group at 22H. These three metabolites belong to the class of compounds known as prostaglandins and related compounds.

**Discussion**

The subject of whether eggshell color affects eggshell quality has attracted the interest of researchers. In our study, there were no significant differences between LBS...
and DBS lines in terms of eggshell thickness and eggshell proportion, which is consistent with previous research findings [23, 26]. However, the eggshell strength of DBS line eggs was significantly weaker than that of LBS line eggs, which may be attributed to the influence of eggshell pigment on the eggshell calcification process. Eggshell pigment primarily exists in a vertical crystal layer [27, 28], and can change calcite crystal size and morphology, thus affecting eggshell quality [29]. Pigment particles stored in the shell gland epithelial cells achieve their maximum deposition rate at the end of eggshell calcification [9], which may impair the deposition of vertical crystal layers, resulting in reduced density and a looser structure, thus diminishing eggshell strength.

In this study, Pp IX concentration in the shell glands of the DBS group was significantly higher than that in liver at three time points following oviposition. At 16H and 22H following oviposition, the Pp IX concentration in the shell gland differed significantly across WS, LBS and DBS groups, but not in liver. This confirms that Pp IX is stored in the shell gland before being released into uterine fluid, but does not necessarily imply that Pp IX is synthesized in the shell gland. However, the significantly higher expression of ALAS1 in the shell gland compared to the liver of the DBS group, as well as its higher expression in shell glands of the DBS group compared to that in the WS group, suggests that Pp IX is synthesized de novo in the shell gland from glycine and succinyl coenzyme A. As an alternative hypothesis, some scholars posit that Pp IX precursor in the shell gland comes from lysed red blood cells [21, 22]. They suggest that damaged and aging red blood cells are phagocytosed by macrophages to release heme, which is then decomposed by FECH into Pp IX. However, in this study, there was no significant difference in the expression level of FECH among the three hen groups, indicating that Pp IX in the shell gland is not produced by heme decomposition. This further supports the hypothesis that Pp IX is synthesized de novo in the shell gland.

Although the expression of ALAS1 in the shell gland differed significantly between WS and DBS groups, we did not find expression differences in other synthase genes in the Pp IX biosynthesis pathway, which may be due to substrate concentrations or transporters responsible for transporting ALA out of the mitochondria. It remains unclear how these genes are regulated. ATP binding cassette subfamily B member 10 (ABCB10) is implicated in early mitochondrial steps of heme synthesis, most likely via facilitating ALA export out of the mitochondria [30]. In this study, there was no significant difference in ABCB10 expression among the three groups at any of the three post-oviposition time points. In addition, research has demonstrated that SLC25A38 can promote ALA production by delivering glycine into the mitochondria or by exchanging glycine through the inner mitochondrial membrane of red blood cells [31].

### Table 3 SNPs identified in chicken ALAS1 DNA sequences

| Location     | SNPs | Variation type     | Region | NCBI number | Genotype<sup>a</sup> |
|--------------|------|--------------------|--------|-------------|-----------------------|
|              |      |                    |        |             | WS        | LBS      | DBS      |
| 12:3346427   | A → G | Synonymous variant | Exon2  | rs734271895 | 18:00    | 69:3     | 0:6:11   |
| 12:3346436   | G → A | Synonymous variant | Exon2  |             | 0:0:18   | 11:6:1   | 17:0:0   |
| 12:3346451   | C → T | Synonymous variant | Exon2  |             | 18:00    | 68:4     | 0:6:11   |
| 12:3347938   | G → A | Synonymous variant | Exon4  | rs317202842 | 1:3:14   | 18:00    | 17:0:0   |
| 12:3349200   | T → C | Splice region variant, Synonymous variant | Exon6 | rs316944705 | 1:4:13   | 18:00    | 17:0:0   |
| 12:3349584   | T → C | Synonymous variant | Exon7  |             | 0:0:18   | 10:6:2   | 17:0:0   |
| 12:3349587   | G → A | Synonymous variant | Exon7  |             | 1:4:13   | 18:00    | 17:0:0   |

<sup>a</sup>The number ratio of different genotypes in WS, LBS, and DBS groups, number of wild homozygous: number of mutant heterozygous: number of mutant homozygous

### Table 4 Description of metabolite molecules potentially contributing to eggshell brownness

| Metabolites                          | Comparison group | VIP | Fold change (DBS/LBS) | p value |
|--------------------------------------|------------------|-----|-----------------------|---------|
| Protoporphyrin IX                    | shell gland, 6H  | 1.66| 1.64                  | 0.0089  |
| 11-hydroxy-E4-neuroprostane          | shell gland, 6H  | 1.76| 1.94                  | 0.0091  |
| 15-dehydro-prostaglandin E1(1-)      | uterine fluid, 22H | 1.83| 2.02                  | 0.0155  |
| Prostaglandin G2 2-glyceryl Ester    | uterine fluid, 22H | 1.78| 2.02                  | 0.0334  |
results show that the expression of *SLC25A38* in the DBS group was significantly higher than that in the WS group at 22H, suggesting that the synthesis of Pp IX may still be taking place in the shell gland epithelial cells during rapid deposition of Pp IX. It remains to be determined whether *SLC25A38* in the epithelial cells of the shell gland play the same role as in red blood cells.

All genes in the Pp IX synthesis pathway were expressed in the WS group, however, their eggshells remain colorless. We hypothesize that a potential cause is the low expression level of *ALAS1* in the WS group eggs, resulting in low Pp IX production, which is almost entirely utilized for the synthesis of heme. Another possibility is that the Pp IX outflow route from the shell gland to uterine fluid may be restricted. *ABCG2* is a member of the ATP-binding cassette family and is essential for the export of Pp IX from the cell [24, 32, 33]. At 16H, the expression level of *ABCG2* was significantly higher in the DBS group compared to the WS group, but did not alter significantly along these three time points following oviposition, consistent with the results of previous studies [19]. So does *ABCG2* have the function of exporting Pp IX from the shell gland epithelium? There was no significant difference in the concentration of Pp IX in the uterine fluid in DBS group between 6H and 22H, and 22H is the stage when Pp IX is deposited on the eggshell surface, indicating that Pp IX has been gradually released into the uterine fluid since 6H. This means that the transporter proteins that transport Pp IX out of the shell gland epithelial cells have been expressed since 6H. Why does the Pp IX remain in the shell gland? We speculate that during the prophase and metaphase of eggshell mineralization, Pp IX transportation is in trace amounts, so it's not macroscopic. Study has shown that eggshell pigment is distributed in all layers including the shell membrane, but the amount of deposition varies greatly [34]. At 16H, the concentration of Pp IX in shell gland of DBS group was significantly higher than that at 4H, indicating that the synthesis rate of Pp IX in shell gland was greater than the release rate, and Pp IX accumulated continuously in shell gland. Intrauterine injection of prostaglandins has been shown to induce pigment secretion [35]. We speculate that when calcification is almost complete, *ABCG2* or other unknown transporters are stimulated by certain factors, such as threshold Pp IX concentration or prostaglandins, and their gene expression or protein activity is activated, resulting in rapid release of Pp IX from shell glands in large quantities and deposition on the eggshell surface. In this study, the transcript level of *ABCG2* was not significantly increased at 22H, so we speculated that the translation process or protein activity of *ABCG2* might be activated, which needs to be further studied. In addition, *FLVCR1* has been thought to encode a protein for exporting heme [36], but prior studies have shown that *FLVCR1* can export endogenous synthetic Pp IX when ALA or iron is present [37]. We discovered that the expression level of *FLVCR1* in DBS group was
significantly higher compared to WS group at 22H and was highest at 22H compared to 4H and 16H. It is noteworthy that *FLVCR1* may be able to export Pp IX synthesized in shell glands. It is the combined interactions of these genes and metabolites that leads to variation in eggshell brownness (Fig. 7).

The expression level of *ALAS1* was found to be closely related to the intensity of eggshell brownness. Seven SNPs in *ALAS1* gene are synonymous mutations, which do not change the sequence of amino acids. The replacement of synonymous codons, however, may change the synthesis rate of polypeptide chain and affect protein folding and conformation, thus affecting the function of protein [38, 39]. Therefore, SNPs of the *ALAS1* gene may regulate the expression of the *ALAS1* gene, thereby affecting eggshell brownness.

GO and KEGG functional enrichment analysis also provided us with information regarding eggshell deposition and pigmentation. At 16H, signal transduction pathways were highly enriched; at 22H, signal transduction was also prevalent and regulation of transcriptional processes was active, indicating that the expression of genes associated with eggshell and eggshell color deposition in the shell gland epithelial cells was stably regulated and maintained, possibly via the membrane. There may be a connection between lipid metabolism and eggshell deposition and pigmentation, which warrants further study.

Conclusions

The content of Pp IX in the shell gland varies with eggshell color. Eggshell pigment Pp IX is synthesized de novo and stored in the shell gland. *ALAS1* plays a key role in the Pp IX pathway and can influence eggshell brownness by influencing Pp IX production in the shell gland. Transporters such as *SLC25A38*, *ABCG2* and *FLVCR1* and metabolites such as 11-hydroxy-E4-neuroprostane, 15-Dehydroprostaglandin E1(1-) and Prostaglandin G2 2-glyceryl ester may regulate eggshell brownness. According to the results of our study, we think that ALAS1 gene should be knocked down in brown-shell laying hens next to verify its critical role in eggshell brownness, and proteomic studies that can reflect the translation level are also essential. This study lays a foundation for studying the formation mechanisms of brown eggshell color. These results may contribute to further research on the mechanism of eggshell browning.

Materials and methods

Animal management and sample collection

A chicken population that was derived from the Rhode Island Red line were bidirectionally selected for six generations according to their eggshell brownness and resulted in two experimental lines with light brown eggshells (LBS) and dark brown eggshells (DBS). These two lines were raised in individual cages and allowed free access to food and water. The temperature of the room was controlled at 22 °C, and the daily lighting plan was 16 L:8D. At 54 weeks of age, 150 laying hens were randomly selected from LBS and DBS lines respectively, and egg quality was measured on the same day. Eighteen White Leghorn hens, 35 LBS line hens, and 35 DB line hens were selected at 57 weeks of age, with regular laying time and uniform eggshell color, and were assigned to the white eggshell (WS) group, light brown eggshell (LBS) group, and dark brown eggshell (DBS) groups, respectively. A total of 54 hens (18 hens per group) were euthanized — six hens per group were euthanized at 4-, 16- and 22-hour timepoints following oviposition. The location of the next ovulated egg in the oviduct at different time points following oviposition is shown in Additional file 10: Fig. S2. Liver and mucosal layer of shell gland samples (108 total) were collected and stored at −80 °C for subsequent ultra-high performance liquid chromatography - tandem mass spectrometry (UHPLC-MS/MS) analysis. Shell glands and Uterine fluid were collected as follows: at 6 h following oviposition, the oviduct was cut off at the vaginal site, and the shell gland was carefully cut longitudinally though the vaginal opening to expose its mucosal layer. The egg with a very thin shell was removed, and the uterine fluid on the mucosa surface of the shell gland was repeatedly aspirated with a syringe. At 22 h following oviposition, we cut off the oviduct at the vaginal site, extruded the egg in the shell gland through the vaginal site, and extruded the shell gland from the top down to extrude the uterine fluid through the vaginal opening. After collecting the uterine fluid, we cut out the entire shell gland and rinsed it with normal saline, then gently scraped the mucosal layer of the shell gland with the handle of a small medicine spoon.

Eggshell quality determination

Egg weight, L, a, and b values, eggshell strength, eggshell thickness, eggshell weight, and eggshell proportion were determined for 150 eggs of LBS line and DBS line respectively. Egg weight and eggshell weight were measured using an electronic balance. The values of L, a, and b were determined at the center of sharp end,
equator, and blunt end by a reflectometer (Minolta CM600). An egg force reader (EMG-0502, Robotma-
tion, Japan) was used to measure eggshell strength. Eggshell thickness was measured at the sharp end,
equator, and blunt end of eggs using a micrometer screw gauge (Mitutoyo293-100, Kawasaki, Japan). Egg-
shell proportion was the ratio of the weight of the eggshell to the weight of the egg.

**Pp IX concentration determination**
The extraction solution was prepared as described pre-
viously [18], which consisted of hydrochloric acid, ace-
tonitrile, and water. The shell gland and the liver were
ground to powders in liquid nitrogen. Prepared shell
gland and liver powders (both 0.25g) were vortexed with
1.5 mL extraction solution in a centrifuge tube for 15 sec-
onds. Dissolved samples were centrifuged for 10 min at
17,800 x g at 25°C, and 0.3 mL supernatant was trans-
ferred to a microtiter plate for analysis using a microplate
reader (SpectraMax i3x, Meigu Molecular Instrument
Co., Ltd., Austria) at the wavelength of 412 nm. Pp IX
standard curve was established to calculate the Pp IX
concentration of the measured samples.

**RNA extraction, library construction, and transcriptome
sequencing**
Total RNA from 54 shell gland samples was isolated
using TRIzol reagent [40] (Invitrogen, CA, USA) follow-
ing the manufacturer’s protocol. RNA Nano 6000 Assay
Kit of the Bioanalyzer 2100 system (Agilent Technolo-
gies, CA, USA) was used to accurately detect RNA integ-
rity [41]. The initial amount RNA for library construction
was 1 μg. The mRNA with polyA tail was enriched by poly-T oligo-attached magnetic beads [42]. The obtained
mRNA was then randomly fragmented with divalent ca-
tions in Fragmentation Buffer at high temperature. The
first strand of cDNA was synthesized in the M-MuLV
Reverse Transcriptase system using mRNA fragments as
templates and random oligonucleotides as primers, then
the RNA strand was degraded with RNase H, and the
second strand of cDNA was synthesized from dNTPs in
the DNA polymerase I system. Subsequently, the end of
double-stranded cDNA was repaired. After adenyla-
tion at 3’ ends of DNA fragments, Adaptor with hairpin
loop structure were ligated to prepare for hybridization.
AMPure XP system (Beckman Coulter, Beverly, USA)
was used to screen 370-420 bp cDNA fragments, then
PCR was performed. At last, PCR products were purified
by AMPure XP system, and the library quality was evalu-
ated by Agilent Bioanalyzer 2100 system [43]. TruSeq PE
Cluster Kit V3-CBOT-HS (Illumina) was used to cluster
the index-coded samples on a cBot Cluster Generation
System. Following cluster generation, library preparation
was sequenced on Illumina Novaseq platform to generate
150bp paired-end reads.

**Expression quantification and differentially expressed
gene analysis**
Raw data of fastq format were firstly processed through
Perl scripts which removed reads with adapter, reads with
plo-y N and low-quality reads from raw data to obtain the
clean data. Clean reads were aligned to Genome Reference
Consortium Chicken Build 6a (GRCg6a) assembly
provided by Ensembl (https://asia.ensembl.org/index.
html) using HISAT (ver 2-2.1.1). The mapped reads of
each sample were assembled and quantified by StringTie
(ver 2.1.4).

DEGs were identified using the DEseq2 Bioconduc-
tor package based on the generalized linear model. The
screening criteria for differentially expressed genes was
set at p-value < 0.05, |Log2FC| >1. The online biological
tool DAVID (https://david.ncifcrf.gov/) was used for
gene ontology (GO) and Kyoto encyclopedia of genes and
 genomes (KEGG) [44–46] pathway enrichment analysis
of the DEGs. A value of P < 0.05 was considered to have
statistical significance.

**Quantitative real-time polymerase chain reaction analysis**
Total RNA was extracted from the shell gland and the
liver with EasteR® Super Total RNA Extraction Kit (Pro-
mega, Shanghai, China). All operations were performed
at 4 °C or on ice following the manufacturer’s instruc-
tions. Briefly, 1 μg of total RNA was used for cDNA syn-
thesis using PrimeScript™ RT reagent Kit (Takara, Japan).
RT-qPCR was performed using cDNA with CFX96 Real-
Time PCR Detection System (Bio-Rad, USA) and all sam-
ple were run in triplicate using TB Green® Premix Ex
Taq™ (Takara, Japan). The primer sequences were as fol-
ows (5’ to 3’): β-actin, forward: TATGTGCAAGGCCGG
TTTC, reverse: TGCTTTTCTGCCCATACCAA [18];
ALASI, forward: GGTGACAGGAAAGTAAAGA,
reverse: ACTGGTCATACTGGAAGGTG [18]. The RT-
quPCR procedure was 95°C for 30 s, 40cycles of 95°C for
5s and 60°C for 35s.

**SNPs calling with transcriptome data**
Clean data of the same eggshell color were com-
bined into one group and divided into WS group, LBS
group and DBS group. We mapped reads to GRCg6a
assembly with STAR (ver 2.7.9a). Repeated reads were
removed using GATK (ver 4-4.2.3.0). SNP detection
was performed using GATK’s HaplotypeCaller. Dual
alleles were extracted using GATK’s SelectVariants. Variable filtration was used for strict quality control of SNP detected, with the following criteria: 1) QD < 2.0; 2) QUAL < 30.0; 3) MQ < 40.0; 4) FS > 60.0; 5) MQRankSum < −12.5; 6) ReadPosRankSum < −8.0. Vcftools (ver 0.1.16) [47] was used to conduct quality control on SNP data sets. The quality control criteria were: 1) Sample detection rate > 90%; 2) Minimum allele frequency > 5%. Then Beagle (ver 5.2) [48] was used for genotyping filling. PLINK (ver 1.90) was used for case-control association analysis and extracted genotypes of significant SNP contained in candidate gene sequences. Finally, SNPS are annotated based on GRCh38 assembly.

As the traditional Bonferroni correction is too conservative [49], simpleM package [50] was used to calculate the number of valid independent tests In this study, the number of valid independent tests was 2200, so at the genome-wide significance level, the threshold p-value was set as 2.27E-05.

**Metabolite extraction**

Shell gland tissues (100 mg) were ground separately with liquid nitrogen. The homogenate was vortexed with precooled 80% methanol. Then the mixtures were centrifuged at 15000 × g, 4 °C for 20 min after placed on ice for 5 min. The methanol concentration in supernatant was reduced to 53% by adding LC-MS grade water. Then the samples were centrifuged at 15000 × g, 4 °C for 20 min after moved to a Eppendorf tube. In the end, the obtained supernatant was used for LC-MS/MS analysis [51].

Uterine fluids (100 μL) were individually transferred to the Eppendorf tubes and vortexed with precooled 80% methanol. The following procedure was the same as described above for the shell gland part. In the end, the obtained supernatant was used for LC-MS/MS analysis [52, 53].

**UHPLC-MS/MS analysis**

UHPLC-MS/MS analyses were performed using a Vanquish UHPLC system (ThermoFisher, Germany) coupled with an Orbitrap Q Exactive™ HF mass spectrometer (Thermo Fisher, Germany). A 17-min linear gradient with a flow rate of 0.2 mL/min was used to infuse samples into a Hypesil Goldcolumn (100 × 2.1 mm, 1.9 μm). The eluents in positive pole mode were 0.1% formic acid aqueous solution (A) and methanol (B) as well as those in negative pole mode were 5 mM ammonium acetate, pH 9.0 (A) and methanol (B). The solvent gradient was set as follows: 2% B, 1.5 min; 100% B, 10 min; 100-2% B, 10.1 min; 2% B, 12 min. The mass spectral data were obtained from Q ExactiveTM mass spectrometer carrying out in positive/negative mode, and the parameters included spray voltage of 3.5 kV, capillary temperature of 320 °C, sheath gas flow rate of 35 psi and aux gas flow rate of 10 L/min, S-lens RF level of 60, Aux gas heater temperature of 350 °C.

**Identification and quantification of metabolites**

The raw data were processed using The Compound Discoverer 3.1 (CD3.1, ThermoFisher) for peak alignment, peak picking and quantitation. Peak intensities were normalized to the total spectral intensity, and the molecular formula was predicted based on additive ions, molecular ion peaks and fragment ions. Whereafter, mzCloud (https://www.mzcloud.org/), mzVault and MassList database were used for peak matching. At last, accurate qualitative and relative quantitative results were obtained. OPLS-DA and t-test were used to analyze the expression of metabolites. The screening criteria for differential metabolites were VIP > 1, P < 0.05, and |Log1.5FC| > 1. KEGG pathway enrichment analysis of differential metabolites was performed using MetaboAnalyst (https://www.metaboolyst.ca/) online analysis platform. A value of P < 0.05 was considered to have statistical significance.

**Statistical analysis**

Statistical analysis was performed with R software (ver 4.1.1). The eggshell quality between LBS line and DBS line, Pp IX concentrations between liver and shell glands, the gene expression levels between WS group and DBS group as well as between the liver and shell gland were analyzed by t-test. The eggshell quality, the Pp IX concentrations among WS, LBS, and DBS groups were analyzed by analysis of variance. Differences were considered statistically significant at P < 0.05.

**Abbreviations**

Pp IX: Protoporphyrin IX; ALAS1: δ-aminolevulinate synthase 1; SLC25A38: Solute carrier family 25, member 38; ABCG2: ATP binding cassette subfamily G member 2; FLVCR1: Feline leukemia virus subgroup C cellular receptor 1; DEGs: Differentially expressed genes; SNPs: Single nucleotide polymorphisms; ALA: δ-aminolevulinic acid; FECH: Ferrochelatase; CPOX: Coproporphyrinogen oxidase; LBS: Light brown eggshell; DBS: Dark brown eggshell; H: 4 hours following oviposition; 16H: 16 hours following oviposition; 22H: 22 hours following oviposition; WS: White eggshell; RNA-seq: RNA-sequencing; ABCB10: ATP binding cassette subfamily B member 10; UHPLC-MS/MS: Ultra-high performance liquid chromatography - tandem mass spectrometry; GRCh38: Genome Reference Consortium Chicken Build 38; GO: Gene ontology; KEGG: Kyoto encyclopedia of genes and genomes; ALAD: Aminolevulinate dehydratase; HMBS: Hydroxymethylbilane synthase; UROS: Uroporphyrinogen III synthase; UROD: Uroporphyrinogen decarboxylase; CPOX: Coproporphyrinogen oxidase; PPOX: Protoporphyrinogen oxidase.
Supplementary Information

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Additional file 1: Fig. S1. Eggs in shell glands of WS, LBS, and DBS groups at 16 hours and 22 hours following oviposition.

Additional file 2: Table S1. Differentially upregulated genes in DBS group at 16 h compared with 4 h following oviposition.

Additional file 3: Table S2. Differentially upregulated genes in DBS group at 22 h compared with 4 h following oviposition.

Additional file 4: Table S3. Differentially upregulated genes in DBS group compared with WS group at 16 h following oviposition.

Additional file 5: Table S4. Differentially upregulated genes in DBS group compared with WS group at 22 h following oviposition.

Additional file 6: Table S5. Differential metabolites between LBS and DBS groups at 6 h following oviposition in shell glands.

Additional file 7: Table S6. Differential metabolites between LBS and DBS groups at 22 h following oviposition in shell glands.

Additional file 8: Table S7. Differential metabolites between LBS and DBS groups at 6 h following oviposition in uterine fluid.

Additional file 9: Table S8. Differential metabolites between LBS and DBS groups at 22 h following oviposition in uterine fluid.

Additional file 10: Fig. S2. The location of the next ovulated egg in theoviduct at different time points following oviposition.

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Not applicable.

Authors’ contributions

CS, NY, and GX conceived the experimental scheme. JY, ZM, XW, JZ, SG and ZG participated in the sample collection and experiment. JY, ZM, and XW analyzed the data. JY wrote the manuscript. CS made important revisions to the manuscript. All the authors read and approved the final manuscript.

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Availability of data and materials

The raw data are available from the BioProject database with accession number PRJNA871302 (Reviewer link: https://dataview.ncbi.nlm.nih.gov/object/PRJNA871302?reviewer=nkrvnjcf1ttvmjdtefa1b99ec).

Declarations

Ethics approval and consent to participate

All methods related to animal experiments in this study were performed in accordance with the standard guidelines and were reviewed and approved by the Animal Welfare Committee of China Agricultural University. The study was in accordance with ARRIVE guidelines.

Consent for publication

Not applicable.

Competing interests

The authors declare that they have no competing interests.

Author details

1 National Engineering Laboratory for Animal Breeding and Key Laboratory of Animal Genetics, Breeding and Reproduction, Ministry of Agriculture and Rural Affairs, China Agricultural University, Beijing 100193, China. 2 Beinongda Technology Co, Ltd, Beijing 100083, China.

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