The role of dynamin in absorbing lipids into endodermal epithelial cells of yolk sac membranes during embryonic development in Japanese quail

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ABSTRACT Endodermal epithelial cells (EECs) within the yolk sac membrane (YSM) of avian embryos are responsible for the absorption and utilization of lipids. The lipids in the yolk are mostly composed of very low density lipoprotein (VLDL), uptake mainly depends on clathrin-mediated endocytosis (CME). The CME relies on vesicle formation through the regulation of dynamin (DNM). However, it is still unclear whether DNM's participate in avian embryonic development. We examined mRNA expression levels of several genes involved in lipid transportation and utilization in YSM during Japanese quail embryonic development using qPCR. The mRNA levels of DNM1 and DNM3 were elevated at incubation d 8 and 10 before the increase of SOAT1, CIDEA, CIDEC, and APOB mRNA's. The elevated gene expression suggested the increased demand for DNM activity might be prior to cholesteryl ester production, lipid storage, and VLDL transport. Hinted by the result, we further investigated the role of DNM's in the embryonic development of Japanese quail. A DNM inhibitor, dynasore, was injected into fertilized eggs at incubation d 3. At incubation d 10, the dynasore-injected embryo showed increased embryonic lethality compared to control groups. Thus, the activity of DNM's was essential for the embryonic development of Japanese quail. The activities of DNM's were also verified by the absorptions of fluorescent VLDL (DiI-vVLDL) in EECs. Fluorescent signals in EECs were decreased significantly after treatment with dynasore. Finally, EECs were pretreated with S-Nitroso-L-glu-tathione (GSNO), a DNM activator, for 30 min; this increased the uptake of DiI-vVLDL. In conclusion, DNM's serve a critical role in mediating lipid absorption in YSM. The activity of DNM's was an integral part of development in Japanese quail. Our results suggest enhancing lipid transportation through an increase of DNM activity may improve avian embryonic development.

Key words: Development, dynamin, endodermal epithelial cell, lipid, yolk sac

INTRODUCTION

Yolk serves as a main source of nutrients for embryonic growth in avian eggs (van der Wagt et al., 2020). Lipids in the yolk provide about 90% of the energy demand for the avian embryo (Speake et al., 1998; van der Wagt et al., 2020). The yolk lipids, primarily consisting of triacylglycerol, phospholipids, and cholesterol, are derived from livers lipids packaged in very low density lipoproteins (VLDLs). The VLDLs are then transported into growing oocytes (Burley et al., 1993; Schneider, 2016). During embryonic development, the yolk sac membrane (YSM) extends from the abdomen to embrace and ingest the yolk progressively (Starck, 2020; Wong and Uni, 2021). The YSM play important roles in the uptake lipids from yolk and the release lipids into embryonic circulation (Schneider, 2016). The endodermal epithelial cells (EECs), within the YSM, in the area vasculosa are considered to be the most important cells for nutrient transportation in YSM (Lambson, 1970; Sheng and Foley, 2012; Yadgary et al., 2014). Facing the yolk, EECs are responsible for absorption of yolk lipids which are sequentially transported to the embryonic circulation (Schneider, 2016). Several proteins involved in the lipid utilization by YSM in avian embryos have been demonstrated. The lipoprotein receptors expressed on the surface of YSM's and EEC's, such as LR8, Cubulin, and LRP2, have been suggested to contribute to the import of VLDLs (Hermann et al., 2000; Bauer et al., 2013). In YSM, free cholesterol absorbed from yolk is esterified by acyl-CoA:cholesterol acyltransferase to form cholesteryl ester (CE), abundant in VLDL derived from yolk sac (Noble and Cocchi, 1990; Speake et al.,
incubated eggs were automatically turned 6 times daily in the egg incubator.

**Dissemination and Primary Culture of EECs From Japanese Quail Embryos**

We followed the procedure from previous studies to isolate the primary EECs from Japanese quail embryos (Bauer et al., 2013; Lin et al., 2016; Wang et al., 2017). In brief, YSM were separated from the sinus terminals of mesoderm in Japanese quail embryos at incubation d 5 to obtain the endoderm. After digestion with collagenase type IV (17104019, Gibco, MA) at 37°C for 15 min, the endoderm of YSM from 6 embryos in the same group were collected and pooled as a single sample. The pooled samples were seeded in 24-well plates and cultured in DMEM/F12 (pH 7.4, 12400–024, Gibco) with 10% new born calf serum (16010–159, Gibco) and 1% Pen-Strep Am Env Hemo. solution (03-033-1B, Biological Industries, CT). After culture for 2 d, EEC explants were formed and used for further experiment.

**Gene Expression Was Determined by Real-Time PCR**

Total RNA was extracted from Japanese quail embryonic YSM and EECs using the GENEzol Reagent (#GZR100, Geneaid, New Taipei, Taiwan). Two µg of total RNA of each sample was utilized to produce cDNA using a high capacity cDNA reverse-transcription kit (#4368814, Applied Biosystem, MA). The synthesized cDNA samples were used to analyze the mRNA levels using the SensiFAST SYBR Hi-ROX reagent (#BIO-92005, Bioline, London, UK) via the Step One Plus Real-Time PCR System (Thermo Fisher Scientific, Waltham, MA). The real-time PCR program consisted of an initial denaturation for 7 min at 95°C, followed by 40 repeated cycles of denaturation for 10 s at 95°C, annealing for 30 seconds at 60°C, extension for 1 minute at 72°C, and a final extension for 10 min at 72°C. Relative gene expression was normalized by β-actin in the same sample and calculated using the following formula: 2^{−(Ct
target genes- Ct β-actin)} (Pfaffl, 2001). Primer sequences were shown in Table 1.

**Dynasore Injection of Fertilized Japanese Quail Eggs Yolk**

Eggs were incubated in an incubator at 37.5°C until d 3. Then, a candling inspection was implemented to exclude unfertilized eggs and early dysplasia of embryos. The eggs at incubation d 3 were randomly divided into 4 groups as follows: 1) a control group with no treatment (n = 11); 2) a sham group with a drilled hole and puncture by a 30 G needle into the egg yolk to stimulate the dynasore (#14062, Cayman Chemical, MI) injection (n = 12); 3) a vehicle group injected with the same volume of dimethyl sulfoxide (DMSO) as the volume of injected dynasore (n = 15), and 4)
Table 1. Primer set for real-time PCR.

| Gene | Primer sequence (5’−3’) | Product size (bp) | NCBI reference number |
|------|-------------------------|------------------|-----------------------|
| SOAT1 | S5’-GAAGGCGCTATCTGGAACG-3’ | 168              | XM_015290507.1        |
| APOB | S5’-ATCTGCAGCTGACATGACA-3’ | 144              | XM_015850945.1        |
| CIDEA | S5’-CTCGGATCTGGGCTTGGG-3’ | 145              | XM_015852240.1        |
| CIDEC | S5’-CCAGGCTCATGGGACCTAC-3’ | 76               | XM_015874420.1        |
| DNM1 | S5’-TCGTCGACAGGACGAGAC-3’ | 120              | XM_015879114          |
| DNM3 | S5’-CGACTGGATGGGACCTAC-3’ | 101              | XM_015869656.1        |
| ACTB | S5’-GGGCTTCCTTCCTGAGGT-3’ | 151              | NM_205518.1           |

1SOAT1, sterol O-acyltransferase 1; APOB, apolipoprotein B; CIDEA, cell death inducing DFFA like effector A; CIDEC, cell death inducing DFFA like effector C; DNM1, dynamin 1; DNM2, dynamin 2; and ACTB, β-actin.

dynasore group injected with 35 μL of 225 mM dynasore, dissolved in DMSO, into the egg’s yolk (final dosage = 2.538 mg / egg) (n = 15). The holes on the egg shells were filled by paper tape and wax. Then the eggs were incubated until incubation d 10. At d 10, the eggs were opened and the morphology of embryos was observed. To quantify the effects of dynasore injection on the fertilized eggs, ratio of embryonic lethality was calculated using the following formula: ratio of embryonic lethality = number of lethal embryo / number of total embryo.

Isolation of Yolk VLDL and Labeling With 1,1′-Dioctadecyl-3,3,3′,3′ Tetramethylindocarbocyanine Perchlorate (DiI)

The isolation of yolk VLDL from the previous study was used in the present study following the published procedure (Bauer et al., 2013). Fertilized egg yolk at incubation d 5 was mixed with VLDL isolation buffer (20 mM Tris/HCl, 0.2 mM EDTA, 150 mM NaCl). The mixture of yolk and VLDL isolation buffer was centrifuged at 40,000 g for 24 h at 4°C. After centrifugation, the yolk VLDL (yVLDL) was obtained from the top fraction and then dissolved in VLDL isolation buffer. Protein concentration of the isolated yVLDL was determined via the Bradford assay (#23238, Thermo Fisher Scientific). The isolated yVLDL sample containing 1 mg of protein was mixed with 50 μL of Dil solution (2.5 mg/mL in DMSO, #D3911, Thermo Fisher Scientific) The mixture of yVLDL and Dil was incubated and gently shaken for 16 h at 37°C in the dark. After centrifugation at 40,000 × g for 24 h at 4°C, Dil-labeled yVLDL (Dil-yVLDL) was collected from the top fraction of the tube. The Dil-yVLDL was diluted in Dil VLDL buffer (0.02 mM EDTA, 150 mM NaCl) to 150 mg/mL and stored at 4°C.

Treatment of EECs With Dynasore, S-Nitroso-L-Glutathione, and Dil-yVLDL

EECs were seeded on cover slides. EECs were incubated in serum-free DMEM/F12 for 4 h prior to treatments. Then, the medium was replaced by serum-free DMEM/F12 containing dynasore (final concentration 80 μM) to treat EECs for various time periods. The Dil-yVLDL was added to treat EECs for 2 h. The EECs were pretreated with 100-μM S-Nitroso-L-Glutathione (GSNO) (#82240, Cayman Chemical, MI) for 30 min before exposure to Dil-yVLDL. To perform confocal microscopy, EECs were washed twice with PBS and fixed with 4% paraformaldehyde for 30 min. After removing the paraformaldehyde, the cells were washed twice with PBS. 4′,6-diamidino-2-phenylindole dihydrochloride (DAPI) (#sc-24941, Santa Cruz Biotechnology, TX) was used to stain the nucleus. After DAPI staining, the cover slide with stained cells was mounted with nail polish. Images were captured from the Leica Confocal Microscope (TCS SP5 II, Leica, Nussloch, Germany) with 549 nm excitation and 565 nm emission.

Statistical Analysis

Fluorescent signal was quantified by ImageJ software. Statistical analyses were performed using GraphPad Prism (GraphPad Software, Inc., CA). The values were mean ± SEM. The change of mRNA levels and the Dil/DAPI intensity ratio in the EECs treated with dynasore or GSNO for different times were analyzed using one-way ANOVA followed by Tukey’s test to evaluate differences. The embryonic lethality was analyzed using a non-parametric method, Kruskal-Wallis one-way ANOVA, followed by Dunn’s multiple comparison test. The Dil/DAPI intensity ratio in the EECs treated with dynasore at different concentrations was analyzed using a t test. A P-value ≤ 0.05 was considered statistically different.

RESULTS

The mRNA Expression Levels of Lipid Utilization Genes in YSM During Embryonic Developmental of Japanese Quail

The changes of mRNA expression levels of DNM1, DNM3, Sterol O-acyltransferase 1 (SOAT1), Cell Death Inducing DFFA Like Effector A (CIDEA), Cell...
Death Inducing DFFA Like Effector C (CIDEC) and Apolipoprotein B (APOB) in the Japanese quail YSM were determined from incubation d 2 to post-hatch d 4 using real-time PCR analyses. Compared with incubation d 2, mRNA levels of DNM1 were increased from incubation d 8 to 10 and at post-hatch d 2 (Figure 1A). The mRNA levels of DNM3, the other isoform of dynamin, were increased at incubation d 10 and post-hatch d 4 (Figure 1B). The SOAT1 mRNA expression increased from incubation d 10 and reached a peak at d 12 of incubation (Figure 1C). The CIDEA mRNA expression was increased from incubation d 14 to post-hatch d 4 (Figure 1D). The expression of CIDEC was increased at incubation d 16 (Figure 1E). The APOB

**Figure 1.** The mRNA expression levels of lipid utilization genes in YSM during embryonic developmental of Japanese quail. (A) The mRNA levels of DNM1, (B) DNM3, (C) SOAT1, (D) CIDEA, (E) CIDEC, and (F) APOB in YSM were analyzed at the time points indicated. ID and PD represented incubation day and post-hatch day, respectively. The mRNA levels were detected by real-time PCR and normalized by β-actin in the same sample. The values were shown as mean ± SEM. SOAT1, n = 11 for each group; APOB, n = 3 for each group; CIDEA and CIDEC, n = 9 for each group; DNM1 and DNM3, n = 7 for each group. Values with different letters indicated significance at P ≤ 0.05 (one-way ANOVA). Abbreviations: APOB, apolipoprotein B; CIDEA, cell death inducing DFFA like effector A; CIDEC, cell death inducing DFFA like effector C; DNM1, dynamin 1; DNM3, dynamin 3; SOAT1, sterol O-acyltransferase 1.
mRNA was increased from the d 2 to 4 after hatching (Figure 1F).

Inhibition of Dynamins Led to Embryonic Lethality in Japanese Quail

Among the lipid utilization related genes analyzed in the current study, mRNA levels of **DNMs**, particularly **DNM1**, were elevated prior to the changes of **SOAT1, APOB, CIDEA, and CIDEC**. The roles of DNMs in embryonic development of Japanese quail have not been determined. Dynasore, a dynamin inhibitor, was injected into yolks of fertilized eggs to examine whether the inhibition of dynamins affects its development. Dynasore (2.538 mg per egg) was injected at incubation d 3 with observation of embryo morphology at incubation d 10. The morphology of the embryos in the sham and DMSO groups was similar to the control group (Figures 2A−2C). However, the dynasore-injected embryos showed developmental defects (Figure 2D). Statistical results indicated that the ratio of early embryonic lethality in the dynasore-injected group was higher than in the control, sham, or DMSO groups (Figure 2E). The result indicates that functions of dynamins were essential for the embryonic development of Japanese quail.

The Effect of Dynasore on the Uptake of Dil-yVLDL in EECs

To verify whether dynamins were involved in VLDL uptake in EECs in the embryos of Japanese quail, we isolated primary EECs from the embryos at incubation d 5 to examine its ability of VLDL uptake. Dil-yVLDL, yolk VLDL labeled with red fluorescent, was utilized to detect the levels of VLDL internalized into EECs. After pretreating with 80 μM dynasore for 10 min, the red fluorescent signal in EECs exposed to Dil-yVLDL was not different from the control group (Figure 3A). Compared with control group, however, the fluorescent signals were decreased in the EECs pretreated with dynasore for 20 or 30 min (Figure 3A). Quantified results showed that the fluorescent signals in the cells pretreated with dynasore for 20 and 30 min were lower than the control groups (Figure 3B). Moreover, the pretreatments with 80 μM dynasore for 30 min decreased the imported Dil-yVLDL in the EECs exposed to 10, 20, or 30 μg/mL of Dil-yVLDL for two hours (Figures 3C and 3D). Thus, dynamins have a function in the uptake of VLDL into the EECs of Japanese quail embryos.

Uptake of Dil-yVLDL in EECs Was Enhanced by GSNO

To further confirm the role of dynamins in the VLDL uptake into the EECs of Japanese quail embryos, we examined whether an increase of dynamins activity promotes VLDL uptake. After pretreatment with a nitric oxide (NO) donor, GSNO (100 μM), for 30 min to increase the GTPase activity of dynamins (Wang et al., 2006; Kang-Decker et al., 2007), the EECs exposed to Dil-yVLDL for 5 or 10 min showed comparable fluorescent signals compared to those without GSNO pretreatment (Figure 4A). After exposure to Dil-yVLDL for 15 or 20 min, however, EECs presented increased fluorescent signals in the GSNO-pretreated group (Figure 4B). Thus, GSNO enhanced the uptake of Dil-yVLDL and that dynamins play a role in VLDL uptake in Japanese quail embryos.
DISCUSSION

Middle to late stages of incubation are critical periods for the avian embryo to utilize lipids from the yolk (Noble and Cocchi, 1990; Speake et al., 1998). In chicken embryos, the genes related to lipid transportation and lipid metabolism are upregulated in YSM from embryonic d 13 to 17 (Yadgary et al., 2014). This upregulation may enable YSM to actively absorb yolk TG and cholesterol. The increased activity of SOAT and inhibited activity of cholesteryl ester hydrolase in YSM enhanced the absorbed cholesterol converted to cholesteryl ester (Shand et al., 1993; Wang et al., 2017; Lin et al., 2020). The cholesterol esters are then transferred to embryos to support embryonic growth (Noble and Cocchi, 1990; Shand et al., 1993; Speake et al., 1998; Ye et al., 2009; Tiwari et al., 2013). Disrupted utilization of yolk lipids in YSM may lead to nutritional deficiencies resulting in embryonic mortality. EECs within YSM, functions to mediate lipid absorption, lipid remodeling, and lipoprotein assembly, are the predominate cells for transporting lipids to embryos during development (Bauer et al., 2013; Wong and Uni, 2021). Several enzymes and proteins, including SOAT1, lipoprotein lipase, APOB, and LRP8, are involved in lipid transfer to avian embryos (Noble and Cocchi, 1990; Shand et al., 1993; Speake et al., 1998; Ding and Lilburn, 2000). In the current study, we demonstrated that DNMs were required for lipid transportation into EECs of the yolk sac membrane for embryonic growth.

DNM proteins play roles in the uptake of LDL and VLDL to transport nutrients (Goldstein et al., 1982; Ferguson and De Camilli, 2012; Cocucci et al., 2014). The activity of SOAT, functions in the conversion of

Figure 3. The effect of dynasore on the uptake of DiI-yVLDL into EECs. (A) The EECs were pretreated with 80 μM dynasore in DMSO for 10, 20, or 30 min, respectively; the EECs without any treatments were a control group (0 min). Then the EECs were treated with 20 μg/mL of yolk VLDL labeled with DiI (DiI-yVLDL) for 2 h. The EECs were photographed using confocal microscopy and the accumulated DiI-yVLDLs in the EECs were photographed using confocal microscopy. DAPI was used to stain the nucleus. Red signal, DiI-yVLDL; blue signal, DAPI. (B) The quantitative results of the fluorescence intensities of DiI/DAPI ratio in DiI-yVLDL-treated EECs with the pretreatments of dynasore for 0, 10, 20, or 30 min. Data presented as mean ± SEM (n = 3). Statistical analysis was performed by one-way ANOVA. Groups with different superscripts indicated a significant difference (P ≤ 0.05). (C) EECs were pretreated with 80 μM dynasore or vehicle (control) for 30 min and then treated with 10, 20, or 30 ng/mL of Dil-yVLDL for 2 h, respectively. Red signal, Dil-yVLDL; blue signal, DAPI. (D) The quantitative results of the fluorescence intensities of DiI/DAPI ratio in EECs with various treatments in (C). Data were analyzed by t-test. ** P ≤ 0.01, *** P ≤ 0.001. (n = 3). Abbreviation: EEC, endodermal epithelial cells.
cholesterols to cholesteryl esters, in YSM can be regulated by nutrients and hormones (Wang et al., 2017). Cholesteryl esters are actively produced in YSM at mid stages of avian embryonic development (Noble and Cocchi, 1990; Shand et al., 1993; Speake et al., 1998; Ding and Lilburn, 2000). Consistent with previous studies, our results showed expression levels of SOAT1 peaked at d 12 of incubation. Microsomal triglyceride transfer protein (MTP), a VLDL synthesizing enzyme, is highly expressed at 9, 11, and 16 incubation days in chicken YSM (Hermann et al., 2000). The CIDE protein controls lipid metabolism in the liver or adipose tissue (Xu et al., 2012) and interacts with APOB in the VLDL transport vesicles (Ye et al., 2009; Tiwari et al., 2013). Our results indicated that levels of APOB and CIDEA mRNA expression were elevated late and suggested that the synthesis of VLDL in YSM was primarily at late embryonic stages in Japanese quail. The expression level of DNM1 mRNA reached a peak earlier than those for SOAT1, APOB, and CIDEA do, suggesting that the machinery to transport lipids was present prior to the enzymes to produce the nutrients for transportation. The CME are important for the import of LDL and VLDL (Goldstein et al., 1979; Harding et al., 1983). The LDL receptor is known as a classical receptor in CME (McMahon and Boucrot, 2011); however, CME is also involved in mediating VLDL transportation in the avian embryonic yolk (Burley et al., 1993). Previous studies indicate that the mid-stage of incubation is an important period for the YSM to absorb lipids in chicken embryos (Yadgary et al., 2010; Uni et al., 2012; Yadgary and Uni, 2012). It was found that DNM1 and DNM3 mRNA was increased at incubation d 8 and 10, respectively. DNMs also function in the regulation of exocytosis (Jaiswal et al., 2009; Anantharam et al., 2011; Jackson et al., 2015). It is still not clear whether DNMs also play roles in the transport of the nutrients from the yolk to avian embryos. Based on the current data, we depicted a potential mechanism for lipid utilization in EECs (Figure 5); in short, DNMs were highly expressed during incubation d 8 to 10 and a large amount of VLDL was endocytosed into EECs. Thus, substrates were provided to the SOAT1 to esterify cholesterol in ER at the mid-stage of incubation. At late stages of incubation, APOB and CIDEA participate to synthesize VLDL and transport it to the embryonic circulation system.

In the dynasore injection study, all the embryos in the control groups developed normally, implying that the candling inspection at incubation d 3 can help us precisely detect unfertilized eggs or early dysplasia embryos. Injection of a large dose of DMSO into chick egg yolk affects the embryonic growth performance (Wyatt and Howarth, 1976), we made a similar observation. Since dynasore is able to bind to the lipid rafts to affect cell signaling and substrate recognition by their receptors (Preta et al., 2015), it is used as a DNM inhibitor to study cellular endocytosis (Macia et al., 2006; Mohanakrishnan et al., 2017). Because dynasore treatment of EECs reduced lipid accumulation and dynasore injection into embryos inhibited development, we speculated that dynasore inhibited DNM activity resulting in embryonic lethality in Japanese quail. NO stimulates the GTPase and self-assembly activity of DNMs by S-nitrosylation at a cysteine residue to promote endocytosis in endothelial cells (Wang et al., 2006; Kang-Decker et al., 2007). The NO donated by GSNO does not target DNMs specifically; GSNO decreases the augmentation index and improves hemodynamics to alleviate pre-eclampsia (Root et al., 2004; Everett et al., 2014). In spite of these alternative possibilities, GSNO
activated DNM activity and stimulated DiI-yVLDL uptake in EECs.

In conclusion, this is the first research to show that DNM serves a critical role in lipid absorption in YSM of avian embryos. In primary EECs, our results showed activating DNMs increased VLDL import; conversely, inhibiting DNMs suppressed VLDL import. The DNMs participated in the regulation of lipid transportation during embryonic development. Our results showed the injection of DNM inhibitors in fertilized eggs led to embryonic death. Thus, the activity of DNMs was vital during embryonic development. Accordingly, we hypothesized that the functions of DNMs in lipid transportation were required in avian embryonic development.

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DISCLOSURES

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