## MOLECULAR AND CELLULAR BIOLOGY Non-Invited Review

## Recent advances in avian egg science: A review

Philippe B. Wilson

Faculty of Health & Life Sciences. De Montfort University. The Gateway. Leicester, LE1 9BH, United Kingdom

**ABSTRACT** Eggs and egg products form an integral part of the food chain. As such, research into egg structure, function, and production has made an important contribution to the field of poultry science. The past decade has seen significant advances in avian egg science research, with work supplementing our understanding of the nature of the avian egg, and its biological, chemical, and physical properties.

Eggshell color, strength, and chemical composition, poultry nutrition, and genetics have all been intensively studied recently, with significant progress being made in a number of these areas. Indeed, with the prevalence of robust theoretical techniques, it is now commonplace to combine experimental investigations with theory, providing a balanced and interdisciplinary perspective.

Key words: eggshell structure, eggshell function, eggshell formation, laying hen

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## INTRODUCTION

The avian egg is a chemical and mechanical powerhouse, containing within it the ingredients for life, and is protected by a unique crystalline barrier. In poultry, the yolk and albumen are formed in the ovary and oviduct, respectively, and subsequently coated in the uterine fluid of the hen. This secretion contains organic and inorganic materials, with a high concentration of calcium carbonate (CaCO<sub>3</sub>). The calcium carbonate in an ionic form binds to membrane proteins on the outer albumen, and begins to crystallize. In healthy fowl, this process continues until an even, protective shell is formed, whereby the terminal organic layer, or cuticle, is deposited, and acts most notably as a bacterial barrier.

The main focus of this review will be on poultry eggs, examining factors influencing their color, structure, and properties. Eggs and derived products form one of the most popular classes of foodstuffs. Indeed, according to the UK Department for Environment, Food, and Rural Affairs (2017) in the fourth quarter of 2016, more than

24.9 thousand tonnes of egg products were produced in the UK alone. The inherent commercial interests in poultry egg production have a profound impact on research; most notably, work is ongoing in finding new ways to improve poultry feed for egg and meat production, developing our theoretical understanding of the biological processes of fowl, and considering the effects of the environment on laying hens.

Herein is presented a concise review of progress in the field of avian egg research, with an emphasis on species of poultry. Understanding the molecular basis of eggshell formation can have broad impact on poultry feed production, as well as impact the fields of solid state chemistry and crystallization. This review describes recent research on the factors affecting the quality of eggs and eggshells in particular, beginning with a discussion on pigmentation, moving on to nutritional and genetic factors influencing eggshell production, and finishing with a discussion on the impact of medication on eggshell formation.

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Corresponding author: philippe.wilson@dmu.ac.uk

### **EGG PIGMENTATION: AN OVERVIEW**

Eggshell color has provided a great deal of material for research. Indeed, it has been shown in numerous studies (Fulton et al., 2012) that the color of eggs significantly affects commercial retail demand. Originally, the brown pigment extracted from eggshells by Sorby (1875), was called oorhodeine; however, it was one hundred years later that protoporphyrin IX was in fact found responsible for the color (Kennedy and Vevers,

1973). Protoporphyrin IX, biliverdin, coproporphyrin, and uroporphyrin are all found in the shells of domestic egg layers, accounting for the complexity in the coloring of eggshells we see from domestic breeding. In fact, it has recently been confirmed that eggshell color has significant genetic links (Zhang et al., 2005), and a correlation has been identified between brown-pigmented eggshells, egg strength, degree of hatchability, and bacterial resistance (Eleroğlu et al., 2016).

Analytically, the color of eggs is measured using a reflectivity meter. This device detects the degree to which light is reflected from the eggshell and measures the darkness of shell color, a dark shell having a low value and light shell, high. Shell reflectivity is not the most accurate method of detecting the color of eggshells. It has been suggested that this is due to the small area on the shell where the actual sample measurement is made (Roberts et al., 2013; Samiullah et al., 2015).

Spectrophotometry is employed for more accurate results, and measures a value between 0 and 100, where 0 corresponds to black, and 100, white. In spectrophotometry, a much larger area of the shell is analyzed, providing more uniform data on the sample.

Issues remain with measurements of shell color; some shells are covered in amorphous calcium deposits, which shield the pigment from the sensor. Currently, the techniques discussed above are unable to quantify the amount of pigment in the layers of shell. Measurement of the amount of pigment present in the shell itself, as opposed to its effect on light, is often regarded as the gold standard in these studies. For more details on this, see Wang et al. (2009) and Samiullah and Roberts (2013).

In fowl, the deposits of the protoporphyrin IX pigment are mainly located in the outer epithelial cells within the shell gland (Poole, 1967). Some hormones have been found to stimulate pigment production, among them estrogen, progesterone, and prostaglandins, although work is ongoing to detect the exact location of pigment synthesis in domestic egg layers (Soh et al., 1993; Soh and Koga, 1994). Some studies were carried out, which have shown, for example, that pigments are produced in different sites in the Rhode Island Red and White Leghorn breeds (Tamura et al., 1965). Indeed, porphyrin pigments were present in Rhode Island Red uterine tissue, but absent in those of White Leghorns (Stevens et al., 1974).

Every bird within a species will secrete and deposit pigment at difference times, and these will differ in blue-, brown-, and white-egg layers (Liu et al., 2010). Japanese quail is a species that has a similar uterine structure to the hen, and research has therefore historically been linked between the 2 species (Poole, 1967). Pigment is deposited on quail eggs approximately 3 h before laying (Soh et al., 1993), whereas blue-egg-laying fowl, such as Araucana and Cream Legbars, have been found to have pigment deposition occur at approximately 2 h before oviposition (Zhao et al., 2006).

The pigment is thought to be deposited throughout the formation process; however, 50 to 74% is deposited in the last 5 h before lay (Warren and Conrad, 1942). Hormones and other factors such as the mineral content in the shell gland are responsible for the cessation of cuticle deposition, thereby affecting the incorporation of the pigment. Indeed, a number of factors have been identified, which when combined have significant bearing on the concentration of pigment in the cuticle layer, most notably diet and mineral intake (Nys et al., 1991).

## RELATIONSHIP BETWEEN NUTRITION AND EGG COLOR

Improving the color of eggs has long been of commercial and exhibition interest. With consumers now searching for quality over quantity, following the premise that free-range flocks provide superior eggs (Wang et al., 2009), it has been suggested that maintaining egg color in free-range flocks is more difficult than caged layers (Samiullah and Roberts, 2013).

Nutrition is a contributing factor to the quality and color of the egg, both internally and externally (Sekeroglu and Duman, 2011). Indeed, Hooge (2007) found that feeding probiotics to layers could improve eggshell color, particularly in brown-shelled eggs. Supplements containing Bacillus subtilis were administered to 63-week-old Lohman Brown commercial hybrids, resulting in increased pigmentation for up to 2 wk after first delivery. It is not yet clear how this affects the intensity of the color, although the relationship between certain amino acids and mediation of metal incorporation into polyporphyrin has been discussed. Indeed, the enzyme ferrochelatase catalyzes the insertion of iron into protoporphyrin IX. Site-specific mutagenesis on ferrochelatase from Bacillus subtilis identified 2 residues in the active site: histidine 183 and glutamic acid 264, which affected the metal binding of the enzyme when modified (Hansson et al., 2007).

Iron soy proteinate (**Fe-SP**) is characterized as a chelated ferrous group within soy proteinate as a supramolecular environment. Supplements containing Fe-SP can significantly improve eggshell color in brownegg layers (Seo et al., 2010), while vanadium adversely affects pigmentation of the shell (Sutly et al., 2001). Vanadium is a first-row transition metal often found in small quantities in poultry feeds. Indeed, it appears that a vitamin C dosage proportional to the vanadium intake neutralizes the effects of the metal on shell color (Odabasi et al., 2006). There have been suggestions that vitamin D could be responsible for more dilute shell coloring; however, a recent study showed no significant correlation between the 2 factors (Roberts et al., 2013).

Fe-SP supplementation to the order of 100 ppm was found to significantly increase the intensity of brown pigment in eggshells from a study of 800 26-week-old Hy-line Brown hens (Seo et al., 2010). This is in good agreement with previous work by Park et al. (2004)

and Paik et al. (2009), who suggested that increased levels of Fe led to a proportional increase in erythrocyte formation and breakdown. As erythrocytes synthesize porphyrin, this accompanied an increased intensity in pigmentation of the eggshells.

The effect of nutrition on alternative pigments also has been considered. Indeed, the Araucana breed of poultry originates from South America and lays a blue-green egg, with the color permeating throughout the shell, as opposed to being deposited mainly throughout the cuticle. Araucana chickens were treated with high levels of dietary antioxidants, namely, vitamins A retinol and E acetate, and the reflectance of the shells measured (Dearborn et al., 2012). Differences in eggshell coloration between females of the same breed were also considered on the basis of the sexual signaling hypothesis (SSH). The SSH suggests that the quality of females from a particular species or breed could be deduced from the intensity of eggshell pigmentation (Moreno and Osorno, 2003). Dearborn et al. (2012) reared birds to maturity and administered one of 2 diets, based on high and low antioxidant concentrations, respectively. These diets were reversed after 8 wk to ensure that birds were subjected to both extremes of antioxidant content. Antioxidant content of the feed was found to have little effect on the color of the eggshells produced by the Araucana in this study. Reflectance measurements, however, identified significant differences in pigmentation of eggs laid among different members of the study. It was suggested that the sensitivity of the reflectance detector did not allow for meaningful determination of the effect of antioxidant content in feed on blue-green pigmentation in eggshells of Araucana chickens.

It is clear that feed has some effect on eggshell characteristics, with individual components contributing more or less; however, the color and physical features of eggshells are also rooted in the genetic factors of the species, breeds, and individuals.

## IMPACT OF GENETIC FACTORS ON EGGSHELL COLOR AND QUALITY

The control of eggshell color by several genes that code for proteins and enzymes regulating the production and deposition of pigments was initially documented in the late 20th century (Vanbrummelen and Bissbort, 1993). Specific brown-egg genes, however, remain elusive, although the higher activity of certain key enzymes in brown-egg layers suggests that the brown egg trait is purely of genetic origin (Schwartz et al., 1980).

Wardecka et al. (2002) established that the same region on chromosomes 2, 4, 5, 6, and 11 influences eggshell color. Phenotypic observations showed that crosses of white with brown-egg layers resulted in intermediate colors—indicating a codominance effect (Hall, 1944). The same study found higher concentrations

of pigment in crosses of males from brown-egg-laying breeds with females from white-egg-laying breeds than the reverse, suggesting a degree of sex-linkage associated with this pigmentation.

The expression of 2 genes, SLCO1A2 and SLCO1C1, in the shell gland was found to be associated with the coloring of the eggs (Zheng et al., 2014). It was suggested by Dunn (2011) that it may in fact be more productive in the short term to explore the genetic basis of pigmentation by means of selection and breeding rather than molecular methods, until such a time as these become less practically and computationally expensive.

A recent advance in genetic and proteomic studies has been the use of novel isobaric tag technology (iTRAQ); this is used to identify specific proteins regulated by functional genes and measure their expression (Ross et al., 2004). Li et al. (2016) employed this technique to identify regulatory proteins responsible for the brown pigmentation and suggest mechanisms for their action. Two-hundred-eighty-one hens laying both light and dark brown eggs were subjected to shell gland epithelial cell sampling, with 147 differently expressed proteins between the 2 groups being identified as affecting eggshell color. Hens laying light-shelled eggs were found to express proteins that decreased the synthesis of protoporphyrin IX, whereas dark-shelled egg layers possessed expression of proteins regulating the formation of hemoglobin. The identification of these proteins by Li et al. (2016) highlights the feasibility of the iTRAQ methodology and provides a basis for proteinbased studies on the mechanisms behind eggshell pigmentation.

Francesh et al. (1997) estimated the genetic parameters associated with egg color, using a subset of native Catalan poultry breeds. Penedesenca Negra, Prat Lleonada, and Empordanesa Roja populations were obtained from the Catalan Institute for Research, Technology, and Agriculture (IRTA) Poultry Genetic Program. Feed was uniformly administered across the breeds for each period of growth and lay. Eggs were collected from 3 consecutive d each wk, from 18 to 39 wk of age, and color was estimated by reflectometer. The complete pedigree for each hen had been recorded, and average egg weights and color measurements were obtained. The Empordanesa breed was found to possess different genetic characteristics and additional links to egg color and heritability than the other breeds in the study. Indeed, the Empordanesa breed showed a negative correlation among breeding birds selected for egg number, and intensity of pigmentation of the shells of eggs from these strains.

The impact of breeding for intensive laying on the color of eggshells has recently been revisited (Mulder et al., 2016). Environmental and genetic variance in purebred and crossbred laying hens was investigated at different points in the laying period. More than 167,000 eggshells from purebred and more than 79,000 eggshells from crossbred laying hens were analyzed for color. It was concluded that genetic selection could be used

to judiciously select for uniformity of eggshell color, a trait often sought out by consumers. Indeed, Mulder et al. (2016) suggest future crosses between purebred and crossbred strains in order to maximize selection for uniformity of eggshell pigmentation.

Recent genome sequencing of the Araucana breed of chicken has identified genes responsible for the blue coloration of eggs from this breed. Indeed, work by Wang et al. (2013) and Wragg et al. (2013) had identified expression of the SLCO1B3 gene in the uterus of the oviduct in Araucana as resulting in formation of blue eggshells. Jeong et al. (2016) extended investigations in the expression of this gene to alternative blue-egg layers, noticing that the SLCO1B3 gene was expressed similarly across the breeds. Moreover, the Araucana derivative breed considered by Jeong et al. (2016) displayed characteristics clearly removed from numerous popular breeds, physical size, flight ability, and egg production being a subset of these. Indeed, the origin of this breed suggests little crossing with other breeds within the investigation, such as Dongxiang from China. It is therefore likely that expression of the SLCO1B3 gene is responsible for the coloration of the eggs in breeds known to possess blue-shelled characteristics.

Genetic factors pertaining to various shell characteristics have been studied for a number of years. Indeed, although shell color is of significant commercial interest, the ability of eggshells to endure physical stress is of concern relative to the handling and packing of eggs.

## GENETIC FACTORS ASSOCIATED WITH SHELL STRENGTH

Sun et al. (2016) published data on candidate genes expressed in the uterus during calcification and their relationship with eggshell mechanical properties. They considered the mechanism for variable eggshell strength to verify if calcification-related genes were involved in the determination of shell strength, as had previously been postulated (Ahmed et al., 2005). Quantitative polymerase chain reaction (qPCR) was used to detect the expression of the selected genes in 2 groups of White Leghorns laying strong and weak-shelled eggs. The eggs were collected at 60 wk of age to obtain samples from the optimum laying period (Zita et al., 2009).

The study found that expression of the CALB1 gene increased in the strong-shelled group compared to the weak-shelled, confirming previous work in the field (Bar et al., 1992). This gene is associated with calcium transport in the intestine and uterus; it is therefore unsurprising that CALB1 expression also affects the strength of eggshell. More intriguing is the overexpression of the DMP4 gene is the weak-shelled egg group. This gene codes for a protein used to bind calcium to the shell, a function different from the transporting behavior coded for by CALB1 (Hao et al., 2007). Sun et al. (2016) therefore suggested additional work on the behavior of

DMP4 to gauge the advantageous effects of expression over strength of shell.

Recent work by Sun et al. (2016) employed genomewide association analysis to understand the genetic basis of eggshell structure and mechanical properties. Sixty-six-week-old hens (n = 927) were subject to analvsis, with eggshell thickness, effective layer thickness, mammillary density, and mammillary layer thickness being measured. Four key genes coding for ion transport, ABCC9, ITPR2, KCNJ8, and WNK1, were found to contribute significantly to the regulation of eggshell thickness. The ITM2C and KNDC1 genes also were identified as affecting mammillary thickness and mammillary density, with chromosome lengths found to be proportional to eggshell thickness variation per chromosome. ITM2C conventionally controls the assembly of protein signaling complexes, and the KNDC1 gene is best known for regulating neuronal dendrite growth (Huang et al., 2007). These robust identifications of gene expression relating to mechanics properties provide a meaningful contribution to the understanding of shell ultrastructure.

Although nutritional and genetic factors contribute significantly to numerous eggshell properties, environmental influences have been demonstrated to affect shell characteristics and, in particular, color.

## EFFECTS OF AGE, STRESS, AND MEDICATION ON LAYING AND PIGMENTATION

It is well known that uniform, dark-brown eggs are the goal of many commercial breeders. It has been noted that there are significant, quantifiable shell color differences among breeds laying brown eggs, and among individuals within those breeds (Grover et al., 1980). Although seemingly evident, the fact that blue and brown pigment concentrations are higher within the glands of each respective egg layer is an indication of the pigments being breed specific, as opposed to being unilaterally produced in all breeds (Liu et al., 2010).

Odabasi et al. (2007) followed a flock of commercial hybrids at different ages, noting the differences in degree of pigmentation of the shell. They found that color remained relatively constant with the bird, most specimens having lighter eggs at the start of the laying period and correspondingly light eggs at the end.

A general trend of decreased pigmentation with age often has been observed (Samiullah et al., 2016). A number of hypotheses suggest that the increase of egg size with age results in a dilution of the pigment on the shell, a symptom of pigment concentration cover over the entirety of the egg.

Samiullah and Roberts (2013) conducted a longitudinal study on a flock of commercial hybrids. Longitudinal studies differ from horizontal studies in that factors from the same flock are considered at different ages, as opposed to different flocks at different ages.

They found no significant difference in eggshell color between 35 and 75 wk of age, noting, however, that the color at 25 wk was quantifiably darker than for other age groups. The color of shells became lighter as age increased; the amount of protoporphyrin IX in one gram of whole eggshell from 33-, 50-, and 67-week-old Hyline Browns did not differ significantly. Conversely, the concentration of the pigment in the cuticle layer itself was found to be significantly higher in 50-week-old birds than that of 33 and 67 weeks. The study therefore concluded that following the initial optimum laying period, eggshell color became paler, suggesting that age was proportional to paleness of shell in brown-egg layers.

The effect of stress from such stimuli as handling and relocation on the laying trends of brown-egg layers has been of particular commercial interest (Reynard and Savory, 1999). Indeed, for 4.5 h prior to oviposition, stress was found to cause a delay in laying of up to 3 hours. A stress threshold was observed whereby hens became unable to lay if the levels and duration of the stress reached or surpassed this.

Molting stress is an example of a condition significantly affecting the variations observed in eggshell color once laying resumes. Aygun (2013) showed that the response to molting with respect to eggshell color varied significantly among individuals in a flock. For caged flocks, high cage densities were proportional to the level of stress exhibited by the birds. Cage design, fear, and frequency of disturbance were also identified as contributing to changes in egg color through the effects of stress.

It has been noted that abnormalities such as paleness, calciferous deposits, and some structural defects were related to environmental influences (Mills et al., 1987). This was explained by the retention of the egg in the shell gland, where additional calcium deposition onto the formed egg results in masking of the ground color. Physical stress such as feather removal and extremes of temperature also were confirmed to adversely affect pigment deposition on the shell.

Drugs have been identified to have a profound effect on both laying and the color of the shell. Prostaglandins are active lipid compounds with biological effects similar to those of hormones. The effects of intrauterine injections of prostaglandin  $F2\alpha$  (PGF<sub>2 $\alpha$ </sub>) on pigmentation of brown eggs laid in the time following administration has been determined (Soh and Koga, 1994). The PGF<sub>2 $\alpha$ </sub> caused oviposition to occur on a reduced timescale as compared to the control group, but resulted in the production of paler eggs—likely due to the time reduction of the egg residing in the shell gland. Indomethacin also was independently administered and was found to entirely inhibit pigment secretion.

Nicarbazin, an anti-coccidial medication, has been noted to result in production of unpigmented eggs for up to a wk after first dose (Hughes et al., 1991). The concentration and duration of treatment with the medication affected pigmentation, but not synthesis of

the pigments themselves. Indeed, the effect was reversed within a wk of completion of the treatment, the birds returning to producing eggs of the same coloration as previously. It has been concluded that although pigment is produced under treatment, it is the medication that affects deposition onto the shell and cuticle, as opposed to its synthesis (Schwartz et al., 1975).

## IMPORTANCE OF COLOR AND QUALITY OF SHELLS

The weight of the egg itself, as well as that of the shell, its strength, abnormalities in structure, yolk color, and properties of the albumen, are indicators for uniformity of the shell physical properties, and internal standard of the egg.

A significant correlation between brown-shelled eggs and the strength of the shell has been previously reported (Yang et al., 2009), with a similar study (Joseph et al., 1999) noting that the specific gravity of the eggshell was higher in brown eggs than blues and whites.

There was no correlation between internal quality of the egg, and the color of the shell (Sekeroglu and Duman, 2011); however, shell structure strength and thickness were found to be related to pigmentation (Schreiweis et al., 2006).

Mroz et al. (2007) reported on the relationship between surface ultrastructure and hatchability of turkey eggs. The hatchability of the eggs from Broad-breasted White Turkeys was considered as a function of shell surface character. In this report, eggs (n=17,590) were split into 3 groups. Group 1 contained regular eggs with no visible calciferous deposits or deformities, group 2 consisted of rough eggs with some calcium deposition and surface nodules, and group 3 contained eggs with significant pigment-spotting.

Each of the eggs was individually marked and all were obtained at the peak of the laying season. Scanning electron microscopy (SEM) was used to observe the shell microstructure, and hatchability was estimated based on fertility rate and number of dead embryos—specifically of healthy poults. The main differences between groups 1 and 2 included varying cuticle size, size of crystal, palisade and mammillary layers, curved mammillary walls, and thicker inner shell membrane fibers. Group 3 was found to contain more eggs with thicker crystal layers and curved fibers of inner membranes, in particular, compared to group 1.

Of all eggs included in the study, those in group 1 had the highest hatch rate, at 77%. Groups 2 and 3 had gradually higher embryo mortalities at just under and just over 5%, respectively. The findings therefore suggested that, certainly in Broad-breasted White Turkeys, smoother eggs with less pigmentation tend to have higher hatchability than their rougher, more pigmented counterparts.

### EGGS AS A BARRIER TO INFECTION

It is generally thought that the eggshell itself serves as both a barrier to physical impact, as well as to pathogens (Sá e Silva et al., 2016). There has therefore been a great deal of research carried out to understand the factors influencing the ability of the shell to act as a preventative medium to infection, as well as methods of treatment to avoid consumer infection (Berrang et al., 1999: Messens et al., 2005).

The penetration of 7 common pathogenic agents, Staphylococcus warneri, Acinetobacter baumannii, Alcaligenes sp., Serratia marcescens, Carnobacterium sp., Pseudomonas sp., and Salmonella enteritidis, through the eggshell has previously been reviewed (De Reu et al., 2006). This research correlated bacterial penetration, with structural features of the eggshell and particular strains of the bacteria.

Shells were inoculated with selected strains and stored at 20°C and 60% humidity for a period of 3 wk, with eggshell penetration being regularly monitored by the Berrang agar method, and whole egg contamination sampled after 3 weeks. The method consists of draining the egg contents and filling the shell with treated agar before being closed with commercial silicone. Among the constituents of the treated agar was triphenyl tetrazolium chloride (**TTC**). As bacterial colonies grew, the TTC was reduced to formazan, which has a distinctive red coloration. Through candling the eggs daily during the first wk and 3 times the following wk, it was possible to record bacterial penetration through the shell (Berrang et al., 1998).

It was found that shell thickness and number of pores did not influence penetration. For each bacterial strain, the average cuticle deposition was lower for penetrated than unpenetrated shells, confirming that some strains digest the cuticle itself before penetrating. Nonclustering strains and those capable of movement were found to penetrate more frequently than others, with Pseudomonas sp. and Alcalignenes sp. showing highest contamination through the shells, most samples being penetrated after 4 to 5 days.

Shell thickness and the influence of age on penetration were not statistically significant; however, eggs with high specific gravities (good quality) were less penetrated than inferior examples. The number of pores did not correlate with penetration, but pore sizes were proportional to inner contamination of the shell, suggesting that perhaps a threshold diameter is necessary for additional penetration.

The cuticle on the shell serves as the first line of defense against pathogens, and acts as a waterproofing agent. Removal of the cuticle increased penetration by 20 to 60%. Eggs with poor-quality cuticles were found to have 40% more contamination than examples with more intact cuticles (26%). Finally, as a useful and non-invasive measure, a statistically significant correlation was found between the level of contamination of the outside of the shell and degree of penetration,

suggesting that eggs can be at least partially judged in terms of contamination from an outer inspection of the eggshell itself.

#### **CONCLUSIONS AND FUTURE RESEARCH**

Recent scientific developments in the area of avian egg research have been discussed. This review has covered a broad spectrum of applications and practice related to the science of eggs, and in particular those of laving hens. Numerous factors have been shown to affect eggshell structure and color, implicating fields from enzyme reactivity to calcite formation (Wilson and Williams, 2015; Wilson et al., 2015; Leitch et al., 2016; Wilson and Williams, 2016). It has, however, been noted that a substantial number of questions remain in fully understanding the avian egg; the precise chemical formation of the crystalline structure of the eggshell, the distribution of pores, and processes behind termination of calcification all remain relatively unknown. Indeed, in an era in which hybrid research is becoming the norm, the greatest discoveries are likely to be made by incorporating powerful theoretical methodologies, such as computer simulations (Williams and Wilson, 2017), into biological research, as has been the subject of this review. Our understanding of the eggs and eggshells is continuously developing; however, the avian egg has not vet yielded all of its secrets.

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#### REFERENCES

Ahmed, A. M. H., A. B. Rodriguez-Navarro, M. L. Vidal, J. Gautron, J. M. Garcia-Ruiz, and Y. Nys. 2005. Changes in eggshell mechanical properties, crystallographic texture and in matrix proteins induced by moult in hens. Br. Poult. Sci. 46:268–279.

Aygun, A. 2013. Effects of force molting on eggshell colour, egg production and quality traits in laying hens. Rev. Med. Vet. (Toulouse). 164:46–51.

Bar, A., E. Vax, and S. Striem. 1992. Relationships between calbindin (Mr 28,000) and calcium transport by the eggshell gland. Comp. Biochem. Physiol. A Physiol. 101:845–848.

Berrang, M. E., J. F. Frank, R. J. Buhr, J. S. Bailey, N. A. Cox, and J. Mauldin. 1998. Eggshell characteristics and penetration by Salmonella through the productive life of a broiler breeder flock. Poult. Sci. 77:1446–1450.

Berrang, M. E., N. A. Cox, J. F. Frank, and R. J. Buhr. 1999. Bacterial penetration of the eggshell and shell membranes of the chicken hatching egg: a review. J. Appl. Poult. Res. 8:499–504.

De Reu, K., K. Grijspeerdt, W. Messens, A. Heyndrickx, M. Uyttendaele, J. Debevere, and L. Herman. 2006. Eggshell factors influencing eggshell penetration and whole egg contamination by different bacteria, including Salmonella enteritidis. Int. J. Food Microbiol. 112:253–260.

Dearborn, D. C., D. Hanley, K. Ballantine, J. Cullum, and D. M. Reeder. 2012. Eggshell colour is more strongly affected by maternal identity than by dietary antioxidants in a captive poultry system. Funct. Ecol. 26:912–920.

- Department for the Environment, Food, and Rural Affairs. 2017. United Kingdom Egg Statistics Quarter 1, 2017. Office of National Statistics Publications. Accessed May 2017. https://www.gov.uk/government/uploads/system/uploads/attachment\_data/file/612150/eggs-statsnotice-04may17.pdf
- Dunn, I. 2011. Poultry breeding for egg quality: Traditional and modern genetic approaches. Pages 245–260 in Improving the Safety and Quality of Eggs and Egg Products, Vol 1: Egg Chemistry, Production and Consumption. Y. Nys, M. Bain, and F. Van Immerseel eds. Woodhead Publishing, United Kingdom.
- Eleroğlu, H., A. Yildirim, M. Duman, and N. Okur. 2016. Effect of eggshell color on the egg characteristics and hatchability of guinea fowl (Numida meleagris) Eggs. Rev. Bras. Ciênc. Aví. 18:61–68.
- Francesch, A., J. Estany, L. Alfonso, and M. Iglesias. 1997. Genetic parameters for egg number, egg weight, and eggshell color in three Catalan poultry breeds. Poult. Sci. 76:1627–1631.
- Fulton, J. E., M. Soller, A. R. Lund, J. Arango, and E. Lipkin. 2012. Variation in the ovocalyxin-32 gene in commercial egglaying chickens and its relationship with egg production and egg quality traits. Anim. Genet. 43:102–113.
- Grover, R. M., D. L. Anderson, and R. A. Damon. 1980. The correlation between egg shell color and specific gravity as a measure of shell strength. Poult. Sci. 59:1335–1336.
- Hall, G. O. 1944. Egg shell color in crosses between white- and brown-egg breeds. Poult. Sci. 23:259–265.
- Hansson, M. D., T. Karlberg, M. A. Rahardja, S. Al-Karadaghi, and M. Hansson. 2007. Amino acid residues His183 and Glu264 in Bacillus subtilis ferrochelatase direct and facilitate the insertion of metal ion into protoporphyrin IX. Biochem. 46:87–94.
- Hao, J., K. Narayanan, T. Muni, A. Ramachandran, and A. George. 2007. Dentin matrix protein 4, a novel secretory calcium-binding protein that modulates odontoblast differentiation. J. Biol. Chem. 282:15357–15365.
- Hooge, D. M. 2007. Bacillus subtilis spores improve brown egg colour. World Poult. 23:14–15.
- Huang, J., A. Furuya, and T. Furuichi. 2007. Very-KIND, a KIND domain-containing RasGEF, controls dendrite growth by linking Ras small GTPases and MAP2. J. Cell Biol. 179:539–552.
- Hughes, B. L., J. E. Jones, J. E. Toler, J. Solis, and D. J. Castaldo. 1991. Effects of exposing broiler breeders to nicarbazin contaminated feed. Poult. Sci. 70:476–482.
- Jeong, H., K. Kim, K. Caetano-Anollés, H. Kim, B.-K. Kim, J.-K. Yi, J.-J. Ha, S. Cho, and D. Y. Oh. 2016. Whole genome sequencing of Gyeongbuk Araucana, a newly developed blue-egg laying chicken breed, reveals its origin and genetic characteristics. Nat. Sci. Rep. 6:26484.
- Joseph, N. S., N. A. Robinson, R. A. Renema, and F. E. Robinson. 1999. Shell quality and color variation in broiler breeder eggs. J. Appl. Poult. Res. 8:70–74.
- Kennedy, G. Y., and H. G. Vevers. 1973. Eggshell pigments of Araucano Fowl. Comp. Biochem. Phys. 44:11–25.
- Leitch, J. A., P. B. Wilson, C. L. McMullin, M. F. Mahon, Y. Bhonoah, I. H. Williams, and C. G. Frost. 2016. Ruthenium(II) catalyzed C-H functionalization using the oxazolidinone heterocycle as a weakly coordinating directing group: Experimental and computational insights. ACS Catal. 6:5520–5529.
- Li, G., C. Sun, G. Wu, F. Shi, A. Liu, and N. Yang. 2016. iTRAQ-based quantitative proteomics identifies potential regulatory proteins involved in chicken eggshell brownness. PLoS ONE. 11: e0168750. http://dx.doi.org/10.1371/journal.pone.0168750.
- Liu, H. C., M. C. Hsiao, Y. H. Hu, S. R. Lee, and W. T. K. Cheng. 2010. Eggshell pigmentation study in blue-shelled and whiteshelled ducks. Asian-australas. J. Anim. Sci. 23:162–168.
- Messens, W., K. Grijspeerdt, and L. Herman. 2005. Eggshell penetration by Salmonella: A review. World Poultry Sci. J. 61:71–86.
- Mills, A. D., M. Marche, and J. M. Faure. 1987. Extraneous egg shell calcification as a measure of stress in poultry. Brit. Poult. Sci. 28:177–181.
- Moreno, J., and J. L. Osorno. 2003. Avian egg colour and sexual selection: Does eggshell pigmentation reflect female condition and genetic quality? Ecol. Lett. 6:803–806.
- Mroz, E., K. Michalak, and A. Orlowska. 2007. Hatchability of turkey eggs as dependent on shell ultrastructure. Pol. J. Natural Sci. 22:31–42.

- Mulder, H. A., J. Visscher, and J. Fablet. 2016. Estimating the purebred–crossbred genetic correlation for uniformity of eggshell color in laying hens. Genet. Sel. Evol. 48:39.
- Nys, Y., J. Zawadzki, J. Gautron, and A. D. Mills. 1991. Whitening of brown-shelled eggs: Mineral composition of uterine fluid and rate of protoporphyrin deposition. Poult. Sci. 70: 1236–1245.
- Odabasi, A. Z., R. D. Miles, M. O. Balaban, and K. M. Portier. 2007. Changes in brown eggshell color as the hen ages. Poult. Sci. 86:356–363.
- Odabasi, A. Z., R. D. Miles, M. O. Balaban, K. M. Portier, and V. Sampath. 2006. Vitamin C overcomes the detrimental effect of vanadium on Brown eggshell pigmentation. J. Appl. Poult. Res. 15:425–432.
- Paik, I., H. Lee, and S. Park. 2009. Effects of organic iron supplementation on the performance and iron content in the egg yolk of laying hens. J. Poult. Sci. 46:198–202.
- Park, S. W., H. Namkung, H. J. Ahn, and I. K. Paik. 2004. Production of iron enriched eggs of laying hens. Asian-australas. J. Anim. Sci. 17:1725–1728.
- Poole, H. K. 1967. A microscopic study of uterine eggshell pigment in Japanese quail. J. Hered. 58:200–203.
- Reynard, M., and C. J. Savory. 1999. Stress-induced oviposition delays in laying hens: Duration and consequences for eggshell quality. Brit. Poult. Sci. 40:585–591.
- Roberts, J. R., K. Chousalkar, and S. Samiullah. 2013. Egg quality and age of laying hens: Implications for product safety. Anim. Prod. Sci. 53:1291–1297.
- Ross, P. L., Y. N. Huang, J. N. Marchese, B. Williamson, K. Parker, S. Hattan, N. Khainovski, S. Pillai, S. Dey, S. Daniels, S. Purkayastha, P. Juhasz, S. Martin, M. Bartlet-Jones, F. He, A. Jacobson, and D. J. Pappin. 2004. Multiplexed protein quantitation in saccharomyces cerevisiae using amine-reactive isobaric tagging reagents. Mol. Cell. Proteomics. 3:1154-1169.
- Sá e Silva, M., L. Susta, K. Moresco, and D. E. Swayne. 2016. Vaccination of chickens decreased Newcastle disease virus contamination in eggs. Avian Pathol. 45:38–45.
- Samiullah, S., A. S. Omar, J. Roberts, and K. Chousalkar. 2016. Effect of production system and flock age on eggshell and egg internal quality measurements. Poult. Sci. 96: 246–258.
- Samiullah, S., and J. R. Roberts. 2013. The location of protoporphyrin in the eggshell of brown-shelled eggs. Poult. Sci. 92:2783– 2788.
- Samiullah, S., J. R. Roberts, and K. Chousalkar. 2015. Eggshell color in brown-egg laying hens - A review. Poult. Sci. 94:2566– 2575
- Schreiweis, M. A., P. Y. Hester, P. Settar, and D. E. Moody. 2006. Identification of quantitative trait loci associated with egg quality, egg production, and body weight in an F2 resource population of chickens. Anim. Genet. 37:106–112.
- Schwartz, S., W. A. Raux, B. A. Schacter, B. D. Stephenson, and R. N. Shoffner. 1980. Loss of hereditary uterine protoporphyria through chromosomal rearrangement in mutant Rhode Island red hens. Int. J. Biochem. 12:935–940.
- Schwartz, S., B. D. Stephenson, D. H. Sarkar, and M. R. Bracho. 1975. Red, white, and blue eggs as models of porphyrin and heme metabolism. Ann. N. Y. Acad. Sci. 244:570–590.
- Sekeroglu, A., and M. Duman. 2011. Effect of egg shell colour of broiler parent stocks on hatching results, chickens performance, carcass characteristics, internal organ weights and some stress indicators. Kafkas Univ. Vet. Fak. Derg. 17:837–842.
- Seo, Y. M., K. S. Shin, A. R. Rhee, Y. S. Chi, J. Han, and I. K. Paik. 2010. Effects of dietary Fe-soy proteinate and MgO on egg production and quality of eggshell in laying hens. Asian-australas. J. Anim. Sci. 23:1043–1048.
- Soh, T., N. Fujihara, and O. Koga. 1993. Observations of pigment accumulation in the epithelium of the shell gland and superficial pigmentation on the egg shell in Japanese quail. J. Fac. Agr. Kyushu U. 38:73–80.
- Soh, T., and O. Koga. 1994. The effects of sex steroid hormones on the pigment accumulation in the shell gland of Japanese quail. Poult. Sci. 73:179–185.

Stevens, E. V., L. K. Miller, S. Weinstein, and A. Kappas. 1974. Biosynthesis of delta-aminolevulinic acid and porphobilinogen in the domestic fowl (Gallus domesticus). Comp. Biochem. Physiol. B. 47:779–786.

- Sun, C., Z. Duan, L. Qu, J. Zheng, N. Yang, and G. Xu. 2016. Expression analysis for candidate genes associated with eggshell mechanical property. J. Integr. Agric. 15:397–402.
- Sutly, J. P., R. D. Miles, C. W. Comer, and M. Balaban. 2001. The influence of vanadium on pigmentation of brown-shelled eggs. Poult. Sci. 80:1039–1039.
- Tamura, T., S. Fujii, H. Kunisaki, and M. Yamane. 1965. Histological observations on the quail oviduct with reference to pigment (porphyrin) in the uterus. J. Fac. Fish. Anim. Husb. Hiroshima Univ. 6:37–57.
- Vanbrummelen, R., and S. Bissbort. 1993. Chicken eggshell porphyrins and the glyoxalase pathway: Its possible physiological role. Comp. Biochem. Physiol. B. 104:657–662.
- Wang, X. T., C. J. Zhao, J. Y. Li, G. Y. Xu, L. S. Lian, C. X. Wu, and X. M. Deng. 2009. Comparison of the total amount of eggshell pigments in Dongxiang brown-shelled eggs and Dongxiang blue-shelled eggs. Poult. Sci. 88:1735–1739.
- Wang, Z., R. Liu, and A. Wang. 2013. Comparison of HMOX1 expression and enzyme activity in blue-shelled chickens and brown-shelled chickens. Genet. Mol. Biol. 36:282–286.
- Wardecka, B., R. Olszewski, K. Jaszczak, G. Zieba, M. Pierzchala, and K. Wicinska. 2002. Relationship between microsatellite marker alleles on chromosomes 1–5 originating from the Rhode Island Red and Green-legged Partrigenous breeds and egg production and quality traits in F2 mapping population. J. Appl. Genet. 43:319–329.
- Warren, D. C., and R. M. Conrad. 1942. Time of pigment deposition in brown shelled hen eggs and in turkey eggs. Poult. Sci. 21:515– 520.

- Williams, I. H., and P. B. Wilson. 2017. SULISO: The Bath suite of vibrational characterization and isotope effect calculation software. Software X. 6:1–6.
- Wilson, P. B., P. J. Weaver, I. R. Greig, and I. H. Williams. 2015. Solvent effects on isotope effects: Methyl cation as a model system. J. Phys. Chem. B. 119:802–809.
- Wilson, P. B., and I. H. Williams. 2015. Critical evaluation of anharmonic corrections to the equilibrium isotope effect for methyl cation transfer from vacuum to dielectric continuum. Mol. Phys. 113:1704–1711.
- Wilson, P. B., and I. H. Williams. 2016. Influence of equatorial CHO interactions on secondary kinetic isotope effects for methyl transfer. Angew. Chem. Int. Edit. 55:3192–3195.
- Wragg, D., J. M. Mwacharo, J. A. Alcalde, C. Wang, J. L. Han, J. Gongora, D. Gourichon, M. Tixier-Boichard, and O. Hanotte. 2013. Endogenous retrovirus EAV-HP linked to blue egg phenotype in Mapuche fowl. PLoS ONE. 8:e71393. https://doi.org/10.1371/journal.pone.0071393.
- Yang, H. M., Z. Y. Wang, and J. Lu. 2009. Study on the relationship between eggshell colors and egg quality as well as shell ultrastructure in Yangzhou chicken. Afr. J. Biotechnol. 8:2898–2902.
- Zhang, L. C., Z. H. Ning, G. Y. Xu, Z. C. Hou, and N. Yang. 2005. Heritabilities and genetic and phenotypic correlations of egg quality traits in brown-egg dwarf layers. Poult. Sci. 84:1209–1213.
- Zhao, R., G. Y. Xu, Z. Z. Liu, J. Y. Li, and N. Yang. 2006. A study on eggshell pigmentation: Biliverdin in blue-shelled chickens. Poult. Sci. 85:546–549.
- Zheng, C., Z. Li, N. Yang, and Z. Ning. 2014. Quantitative expression of candidate genes affecting eggshell color. Anim. Sci. J. 85:506– 510
- Zita, L., E. Tůmová, and L. Štolc. 2009. Effects of genotype, age and their interaction on egg quality in brown-egg laying hens. Acta Vet. Brno. 78:85–91.