Prospect of early vascular tone and satellite cell modulations on white striping muscle myopathy

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ABSTRACT
Polyphasic myodegeneration potentially causes severe physiological and metabolic disorders in the breast muscle of fast-growing broiler chickens. To date, the etiology of recent muscle myopathies, such as the white striping (WS) phenotype, is still unknown. White striping–affected breast meats compromise the water holding capacity and predispose muscle to poor vascular tone, leading to the deterioration of meat qualities. Herein, this review article provides insight on the complexities around chicken breast myopathies: (i) the etiologies of WS occurrence in chicken; (ii) the metabolic changes that occur in WS defect in pectoralis major; and (iii) the interactions between breast muscle physiology and vascular tone. It also addressed the effects of nutritional supplements on muscle myopathies on chicken breast meats. Moreover, the review explored breast muscle biology focusing on the early preparation of satellite and vascular cells in fast-growth chicken breeds. Transcriptomics and histological analyses revealed poor vascularity in breast muscle of fast growth chickens. Thus, we suggest in ovo feeding of nutrients promoting vascularization and satellite cells replenishment as a potential strategy to enhance endothelium-derived nitric oxide availability to promote vascularization in the pectoralis major muscle region.

Key words: pectoralis major muscle, vascular tissue, satellite cell, hypertrophy, myodegeneration

INTRODUCTION

Genetic selection has helped the poultry industry to breed fast-growing broiler chickens over the decades. Poultry producers can also meet the demand for chicken meats by improving the feed’s nutritional composition (Tallentire et al., 2018). Likewise, the modern production practices significantly reduced the time required for commercial birds to reach market sized weight. All these are subject to contributions from selective breeding and feeding technologies (Petracci et al., 2015). However, in the last few decades, some striking unintended consequences of fast growth rate are on the rise, including but are not limited to the occurrences of mild and severe muscular myopathies (Petracci et al., 2013).

Most recently, muscular defects such as white striping (WS) and wooden breast (WB) of chicken have been reported as 2 of the emerging pectoral major muscle (PMM) myopathies, and they pose higher economic risks on meat quality (Sihvo et al., 2014). Although, there are concerns about oxidative stress and improper handling; however, according to Kuttappan et al. (2013a; 2012c), Vitamin E supplements or pre-slaughter handling did not influence the appearance of WS phenotype. Therefore, poultry scientists are currently exploring ways to unravel WS and WB etiologies (Trocino et al., 2015). In fact, it is suggested that both myopathies exhibit similar molecular alterations but with distinct phenotypic characteristics (Malila et al., 2018). The causes of the WS defect are focused on the rapid growth of muscular tissues, vasodilation, hypoxia, and lipidosis in the pectoralis major region (Kuttappan et al., 2013a; Mudalal et al., 2015; Dalle Zotte et al., 2017; Lilburn et al., 2019).

Here, we reviewed the meat quality and genetic factors associated with muscle myopathies in broiler chickens. Moreover, considering the hypertrophic muscle development in fast-growing commercial broiler chickens, we speculate that the satellite cells (SC) are unable to keep up with the normal muscle growth and repair. Thus, we suggest that the enhancement of SC could support the cell’s regenerative properties. Furthermore, strategies that increase muscle fiber numbers could likely
reduce muscle hypertrophy stress. For example, hyper-plasia increases the ratio of capillary density to muscle fibers in broiler chickens (Harthan et al., 2014); therefore, we propose that in ovo feeding of L-arginine might positively affect the relaxation of capillary tissue vasodilation in the PMM of broiler chickens.

Meat Quality, Consumers Acceptability, and Economic Impact of White Striping Breast Muscle Defect

Customer choice toward poultry meat is based on its nutritional profile and the relatively low price compared with other meats. The nutritional profile of chicken meat was reported to contribute to human health (Mir et al., 2017). Nevertheless, consumers are more concerned about the quality of meat available in the market and are willing to pay extra to eat safe, healthy, and nutritious breast fillet products. Factors such as meat color and tenderness influence consumer preference. For instance, research showed that consumers would purchase fewer breast fillets with an increase in WS phenotype severity (Kuttappan et al., 2012b). High fat deposits and increased drip loss are among other quality concerns that have been reported to correspond with WS and WB phenotypes (Mazzoni et al., 2015; Huang and Ahn, 2018). These increasing concerns are global problems, especially in the US, Europe, and China. Annually, the poultry industry in the United States of America reports a loss of over $200 million because of muscle-related defects (Bunge, 2019). The other problem is that the United States Department of Agriculture (USDA) recommended in 2017 that abnormal fillets be downgraded and used for processed foods (Qin et al., 2013; USDA-FSIS, 2017).

Growth Rate and Incidence of White Striping in Modern Broilers Chicken

The growth rate of broiler chickens has recorded significant improvement in the last 50 yr. Broiler chickens’ weight at the market age has reached over a 200% increase between 1957 and 2001 (Havenstein et al., 2003). Compared with local breeds, modern fast growth broilers show a significantly higher feed to growth ratio (FCR), which explains the improvement in carcass yield. The carcass of commercial broilers (Ross 308) showed a relative increase in muscular weight than unselected genetically enhanced chickens (Havenstein et al., 2003). Thus, the focus is now on carcass yield with particular interests in increasing breast meat, thigh, and wings, constituting up to 35% of the body mass. In agreement with the study above, fast-growing birds have become more economical and sustainable than the older breeds of chickens (Zuidhof et al., 2014). Over the years, high energy feed helps achieve higher body weight and better FCR. Unfortunately, high energy feed changes the energy dynamics required for physical and metabolic activities in the body. For instance, a comparative study on broilers with fresh and abnormal fillets demonstrates the effect of growth rate on the progression of the muscular disorder and thereby suggested that increased growth rate, as seen in heavy birds, results in higher degrees of white stripes in the fillets (Kuttappan et al., 2012a).

More than 55% of broilers’ breast meat was recorded to have a WS phenotype, and this appearance becomes more severe with higher slaughter age (Kuttappan et al., 2013a, 2013b). Bechtel showed that broiler muscle fibers undergo hypertrophy, whereas the blood capillaries tissue simultaneously decreases over time (Bechtel, 1986). The theory supporting Bechtel data was that chicken breast muscle (i.e., PMM) is composed mainly of type IIb fiber, which has lower capillary density and makes the PMM more vulnerable to muscle myopathies (Verdiglione and Cassandro, 2013). In related research, physical examinations of chicken fillets were reported to have 74% of WS on the meat surface and 97% damaged muscle when examined histologically (Trocino et al., 2015). The WS incidence increased by an average of 8.7 to 75% between 2012 and 2017 (Huang and Ahn, 2018), whereas its severity has a varying intensity in different species of broiler chickens. A study confirmed the thickness variations of white stripes to be about 25.0 and 7.4% in Ross 308 and Cobb 500 birds, respectively, with a higher percentage in heavy-sized than the medium-sized broiler chickens (Lorenzi et al., 2014). Kuttappan et al., therefore, designed a classification system that ranges between normal, moderate, and severe scores based on the visual appearance of the white stripe on the breast fillet samples (Kuttappan et al., 2012a). Accordingly, the grading system has helped to solve the disparity in categorizing the WS thickness.

In summary, muscle myopathies such as WS occur because of the increasing carcass yield of hybrid chickens selected for meat purposes. On this premise, the subsequent sections will provide insights into histology, omics, autophagy, and mechanisms that characterized WS-affected breast muscle in comparison to other muscle defects. We also review recent reports on the possible metabolic disorders such as nutritional deficiencies, oxidative stress, and genetic factors that may have contributed to the alteration in the muscle composition.

CHARACTERIZATION OF WHITE STRIPING PHENOTYPE

Myopathies in the muscle are any neuromuscular diseases arising from defects in the structural composition of muscle fibers. Like other tissues, the muscle is vulnerable to degenerations and pathosis, in which WS and WB are linked with muscular disorders in broiler chickens. Specifically, muscle myopathies are characterized by necrosis and lipidosis, as seen in WS (Sihvo et al., 2014), whereas the WB is distinct by the hardening of the extracellular matrix cells (Velleman, 2020). The etiology of WS is still undefined; however, the rapid rate of muscle development in modern birds is suggested to
overwhelm the supporting biological mechanisms like the SC and blood vessel system, which cause abnormalities in the cellular and molecular activities in the muscle (Velleman, 2015). On this note, it is necessary to discuss the features of typical breast muscles affected by WS.

**Histology of Muscle Affected by White Striping Disorder**

Macroscopically (Figure 1), white striation parallel to the direction of the muscle fibers characterizes the WS myopathy (Petracci and Cavani, 2012) seen on the ventral side of the pectoralis major (Rehfeldt et al., 2011). White striping can be categorized as a polyphasic myodegeneration (Zambonelli et al., 2016). Under a severe condition, WS-affected muscles show fatty deposits, and it becomes unappealing. Histological data prove that WS-affected fillet had abnormal fiber development with gross lesion condition, which compromises the muscle composition (Mazzoni et al., 2015). For example, perimysial and endomysial tissues of abnormal muscles lose their cross-sectional structure leading to fibrosis in severe WB conditions (Velleman et al., 2017; Velleman, 2020).

The significant increase of the muscle fiber diameters in heavy type chickens reduces the interstitial space for capillary tissue networks, thus, causing a lower supply of oxygenated blood (Velleman and Clark, 2015). Considering the hypoxia in WS meat, Greene et al. (2019) reported the downregulation of oxygen transport proteins (hemoglobin subunits and hemerythrin) in the abnormal muscle. Upregulations of hypoxia-inducible factor subunits (HIF-1α and HIF-2α) combined with downregulations of oxygenated blood-transport proteins indicate that HIF-1α and HIF-2α are adaptive mechanisms for low oxygen supply in abnormal muscles (Nakazawa et al., 2016;)

**Omics Profiling of White Striping**

As previously described, the WS syndrome causes reduced space for the network of blood capillary tissues because of the larger fiber sizes in fast-growing broilers (Velleman and Clark, 2015). Considering the hypoxia in WS meat, Greene et al. (2019) reported the downregulation of oxygen transport proteins (hemoglobin subunits and hemerythrin) in the abnormal muscle. Upregulations of hypoxia-inducible factor subunits (HIF-1α and HIF-2α) combined with downregulations of oxygenated blood-transport proteins indicate that HIF-1α and HIF-2α are adaptive mechanisms for low oxygen supply in abnormal muscles (Nakazawa et al., 2016;)

Figure 1. The histomorphology of white striping myopathy. (A) Phenotypic features of White striping in fast-growing broiler chickens. (B) Microscopic image revealing the myodegeneration in the breast muscle of fast-growing broiler chickens.
Greene et al., 2019). HIF-1α, a transcriptional factor in hypoxia condition, modulates the gene expressions of glyceraldehyde-3-phosphate dehydrogenase, phosphoglycerate kinase 1, and lactate dehydrogenase A proteins (Kaluz et al., 2008; Marin-Hernandez et al., 2009; Malila et al., 2019). These alterations in glycolysis and gluconeogenesis support the notion that HIF-1α activation is essential for anaerobic metabolism in breast myopathies (Malila et al., 2019). The modulation of metabolic enzymes has pointed out by Lake and Abasht (2020) is probably not the main cause of WS and WB; however, changes of these enzymes could reprogram the cellular activities for limited oxygen supply conditions in affected breast muscle (Malila et al., 2019).

The RNA-Seq analysis of myopathic muscles revealed several differentially expressed genes depend on HIF-1 transcription (Malila et al., 2019; Marchesi et al., 2019). HIF-1 transcription regulatesangiogenesis, myogenesis, vasodilation, and apoptosis (Marchesi et al., 2019). Among the 20 most-affected genes (e.g., CRH, MDK, PTX3, COQ10 B, IGLL1) that were differentially expressed are genes involved in hypoxia, amino acid (AA) metabolism, muscle tissue regeneration, and inflammatory activities (Marchesi et al., 2019). The study also confirmed the downregulation of angiogenesis transcription genes (such as bone morphogenetic protein-4, reticulon-4, and fibroblast growth factor-1) (Marchesi et al., 2019). We could deduce from their analyses the significant impacts that the altered genes have on the development of capillary tissue damage, contributing to polyphasic myodegeneration in WS-affected muscles (Figure 2). Under this premise, it is pertinent to investigate other molecular factors linked to muscular distress.

**Autophagy and Muscle Protein Turnover**

Autophagy is among the biological processes activated as innate cellular response mechanisms in damaged cells (such as apoptosis). During metabolic disorders, reactive oxidative stress causes oxidative damage and triggers cellular apoptosis (Schwartzman and Cidlowski, 1993; Chen et al., 2016). In an attempt to mitigate the oxidative damage in muscle, evidence showed that autophagy systems are deployed as repair mechanisms in regulating muscular protein degradations induced by oxidation (Rabinowitz and White, 2010). The autophagy process is critical to maintaining cellular integrity by ensuring a balance between cellular synthesis and death (Ohsumi, 2001). However, the synthesis and degradation imbalance of protein cells in muscular tissue could contribute to degenerative damage seen in WS or WB poultry meat. This notion suggests that rapid growth outpaces the protein turnover mechanisms in broiler chickens (Vignale et al., 2017; Tallentire et al., 2018).

Avian species express autophagic genes that are tissue specific (Piekarski et al., 2014). Some of the autophagy-related genes highlighted in previous studies include LC3, Beclin 1, Bcl2, and Bax (Kabeya et al., 2000; Marino and Lopez-Otin, 2004). An enhanced LC3-II/LC3-I ratio expression was revealed to alleviate induced oxidative stress in the broiler’s muscular cells (Chen et al., 2018).

Figure 2. Putative mechanisms of polyphasic myodegeneration in fast-growing broilers. In a white striping affected breast muscle, multiple cellular processes target satellite cell proliferation, oxidative stress alleviation, and enzymatic reprogramming to repair the damaged muscles. The dysfunction of these mechanisms purportedly causes the myopathic defect in the pectoralis major muscle in broiler chickens. Abbreviations: HIF, hypoxia-inducible factor; ROS, reactive oxygen species; SC, satellite cell; GADPH, glyceraldehyde-3-phosphate dehydrogenase, GK, phosphoglycerate kinase 1; LDH, lactate dehydrogenase; NFIL, nuclear factor interleukin.
The sequences of mRNA transcription obtained from breast muscles of high feed efficiency birds (Rehfeldt et al., 2011) showed upregulations of both the mammalian target of rapamycin (mTOR) and AMP-activated protein kinase (AMPKα1) genes (Piekarski-Welscher et al., 2018). Surprisingly, the upregulation of both genes contradicts their functions, as mTOR and AMPKα1 regulate protein synthesis and inhibit energy usage, respectively.

The RNA seq database for the high feed efficiency broilers documented the upregulation of an autophagy pathway–related gene (Bottje et al., 2017), meaning that fast growth broilers exhibit autophagic enrichment genes. During metabolic distress in the muscle, the autophagy mechanism was triggered to restore homeostatic conditions in the cell (Slupecka et al., 2014; Jokl and Blanco, 2016; Cui et al., 2020). Indeed, these studies support the idea that muscular disorders emanate from the impairment in cell regeneration. Consequently, we suggest that impairment of the autophagy mechanism in breast muscles may potentially contribute to deficiency in protein turnover, lipidosis, and fibrosis linked with chicken meat myopathic tissues.

**White Striping and Wooden Breast Degeneration Response Mechanisms**

The PMM is exclusively composed of type IIb fibers. This implies that the primary source of ATP synthesis in the pectoralis major is via anaerobic glycolytic metabolism. Phosphoglucone isomerase, lactate dehydrogenase, and GNE genes in the glycolytic pathway were altered in WS-affected breast muscles (Zambonelli et al., 2016). On this account, muscle degeneration in WS-affected breast meat could result from the progressive inhibition of the genes encoding the enzymes that regulate anaerobic glycolysis (Pampouille et al., 2019). Lactate dehydrogenase, the key enzyme that regulates pyruvate conversion to lactate, was reportedly downregulated in a chronic hypoxia condition (Kuttappan et al., 2017). The downregulation of lactate dehydrogenase and phosphofructokinase indicate that the anaerobic glycolysis pathway is impaired (Mutryn et al., 2015).

On the other hand, pathways of the glycolytic intermediates are also changed (Zambonelli et al., 2016), leading to lipolysis and fibrosis (Soglia et al., 2015), as evident in WB-affected muscles. Because of these reasons, we still need to determine if the shift in the pathways of the substrate could potentially reduce the energy production in the type II fibers of the PM.

Muscle hypertrophy is favored by upregulations of growth factors like IGF-1 (Musarò et al., 2001; Sandri, 2008; Schiaffino et al., 2013) and myogenic factors like MYC (Pampouille et al., 2019). Because muscle development, degeneration, and repair take place simultaneously, it depicts crosstalk events regulated by multiple mechanisms controlled by the interactions between noncoding RNA and messenger RNA (Petracci et al., 2013). The interaction between miRNA and mRNA induces differential expression of FHL2 (a postinflammatory gene) during the late developmental phase of chicken muscle growth (Li et al., 2019). A dysregulated interaction between miRNA and mRNA will have considerable effects on muscle qualities (Khatri et al., 2018). Enhanced transcription of mRNA and protein translation in fast-growing birds is suggested to contribute to the degenerative WS phenotype. Therefore, elucidating the associated miRNA and mRNA interaction will provide a better understanding of the mechanistic etiology of myopathic muscle in fast-growth chickens.

Li et al. demonstrated that Forkhead Box O transcription (FOXO) signaling pathways are crucial for muscle fiber hypertrophy (Li et al., 2019). During muscle growth, FOXO3 targets several myoblast differentiation genes upregulated by AKT, an upstream gene in the FOXO pathway (Xue et al., 2017). It is not surprising that FOXO3 could maintain homeostasis through negative feedback loops because of the metabolic changes by the intermediates of both Akt and mTOR pathways (Chen et al., 2010; Stefanetti et al., 2018). Moreover, FOXO3 proteins enhance SC proliferation via the notch pathway during the regeneration in muscle cells (Stefanetti et al., 2018). However, a proteomics study revealed upregulations of mTOR and eIF-2 signaling pathways in WS-affected breast muscles (Kuttappan et al., 2017). Under such conditions, WS breast muscles could be triggered by the ubiquitin–proteasome system leading to protein degradation in the muscle fibers. Aberrant protein synthesis in hypertrophic muscles could cause posttranslational modification of E3 ubiquitin–protein ligase 1 in chickens with muscular dystrophy (Matsumoto et al., 2008). Besides, there is a possibility that the intermediates of the P13k/Akt pathway are stimulated because this pathway is involved in an increased protein translation in severe myopathic breast muscles (Deng et al., 2014). Such multifactorial mechanisms make it more challenging to unravel the cause of the abnormal phenotype appearing on chicken breast muscles.

Inflammation functions as one of the biological responses that inhibit oxidative damage caused by excess reactive compounds in tissues. As mentioned in previous studies, cytokines molecules related to muscle myopathies include IL-6, TNF-α, and NK-kB (Lundberg et al., 1995; Salomonsson and Lundberg, 2006). Cytokines regulate both proinflammatory and antiinflammatory cellular responses and may trigger the alleviation of muscle myopathies. Most cytokines produced by monocytes, macrophages, neutrophils, and many other immune cells in the endothelium (Goetz et al., 2004) complicate the pathogenesis of idiopathic inflammatory myopathies (Scheurich et al., 1987; Seko et al., 1993). The genes that encode the inflammatory cytokines in
the abnormal breast muscle mediate cellular activities like proliferation and apoptosis. Data from WS-affected breast meat showed that upregulation of nuclear factor interleukin 3-coding genes demonstrates the effort of the tissue to alleviate apoptotic and necrotic cell activities (Zambonelli et al., 2016). Cytokines like IL-6 and TNF-α might play significant roles in myopathic cell regeneration in the breast muscle tissue. For example, the differential expression of TNFAIP6, an isoform of TNF-α in fast-growing muscles, might degrade the linkage between the extracellular matrix component and cytoskeleton caused by the matrix metalloproteinase expression (Pampouille et al., 2019). TNF-α also mediates MuRF1 protein expression (linked with muscle necrosis and disorder activities) (Minetti et al., 2011).

NK-kB transcription cascade enhances the signals for distressed muscles by increasing the inflammatory lesions in the muscle tissues. Therefore, it is interesting to know that cytokine/NK-kB signaling affects cell growth and regeneration in dystrophic muscles. There is an interplay among the cytokines linked with muscle myopathies, most importantly IL-6, TNF-α, and NK-kB, where they initiate SC growth and proliferation, as well as myoblast differentiation in damaged muscles (Guerci et al., 2012; Yang and Hu, 2018). The development of SC proliferation in the damaged myofibers is an indicator for the initiation of cell regeneration. Another significant cytokine, that is IL-4, participates in alleviating hypertrophic-related defects in the muscles (Guerci et al., 2012). Hypertrophic muscles may use cytokine molecules as a signal for damaged tissue, which leads to regenerative processes in WS-affected muscles (Pampouille et al., 2019).

### RECENT EFFORTS TO ALLEVIATE WHITE STRIPING AND WOODEN BREAST MYOPATHY

Different efforts, such as nutritional supplements and managerial practices, have improved carcass quality and production performance in broiler chickens. Above all, AA are the main component for protein synthesis in the body. This relation has inspired several studies to focus on AA’s role in WS and WB development. Also, AA is critical to major metabolic pathways that regulate muscular growth and development (Wu, 2009). Considering the reports that underlying factors of WS and WB could originate from polyphasic degenerative activities in the breast muscle, modern fast growth birds may require changes in the feed composition. Improper adjustments could significantly affect the birds’ performance, although most researchers believe that nutritional abnormalities do not cause WS and WB disorders. However, this review provides an overview of nutritional effects on WS/WB myopathies (Table 1).

### Nutritional Supplements

#### Amino Acids

Although there are limited reports on the intervention of dietary supplements to mitigate the muscular defect caused by enlarged muscle fibers, no study has concluded that WS/WB is related to

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Table 1. Effect of supplements on breast muscle myopathies.

| Supplements      | Concentration | Effects                                      | Deduction                                      | References                  |
|------------------|---------------|----------------------------------------------|-----------------------------------------------|-----------------------------|
| Glutamine        | 1%            | Exacerbate wooden breast (WB) and White striping (WS) in fillets—raises blood H<sub>2</sub>CO<sub>3</sub>, CO<sub>2</sub>, and decreases O<sub>2</sub> | Hypoxia leads to fatty acid and Ca accumulations | (Livingston et al., 2018)   |
| Arginine         | 0.25%         | Reduced average woody breast score without final BW decrease | Allows satellite cells to recuperate from fast growth in early stage | (Bodle et al., 2018)        |
| Digestible amino acid | 15% reduction at grower’s phase | Alleviate WB score by 32% | Protect against peroxidation in tissues | (Abasht et al., 2016; Bodle et al., 2018) |
| Ascorbate        | 94.4 mg/kg    |                                              |                                               |                             |
| Magnesium        | 3000 mg/kg    | Decrease WB and WS scores by half Upregulate catalase activities in plasma | Protect against peroxidation in tissues | (Estevez and Petracci, 2019) |
| Vitamin E        | 50 mg/kg      | Increased plasma Vitamin E No significant alleviation of muscle damage | WS/WB not related to nutritional deficiency | (Guetchom et al., 2012)    |
| Vitamin E        | (15, 50, 100, 200, 400) mg/kg | No significant correlation in any of the dietary levels | WS occurrence is peculiar to heavy fillet weight | (Kuttappan et al., 2012c)  |
| DL-α-tocopherol acetate | 200 IU/kg | Reduces the severity of WS and wooden breast at growers’ phase | Alluded to early Vitamin E inclusion in the diets | (Wang et al., 2020) |
nutritional deficiencies. Instead of glutamine and arginine supplements to alleviate the muscle defects in a previous study, they improved both the BW and FCR of the experimental broilers, contributing to the WS condition exacerbation (Livingston et al., 2018). From their results, the dietary supplement had an indirect effect on the development of muscle defects in heavy breed chickens. On the contrary, Bodle et al. (2018) decreased the AA ratio in the diet fed to broilers during the grower phase and aimed to replenish SC development from the rapid growth at the starter phase (Powell et al., 2014). Their report demonstrated that feeding lower AA at the grower’s stage could possibly alleviate WB because of reduced SC activities (Bodle et al., 2018). Thus, it is okay to deduce that the PMM progressive degeneration may be reduced by minimizing the feeds’ nutritional content at the grower’s phase.

The implication of hypoxic conditions in myopathic breast muscle is equivocally agreed upon by researchers (Malila et al., 2019). In a previous trial, dietary arginine supplements improved hypoxia in the broiler muscles (Bodle et al., 2018). According to their result, Bodle et al. (2018) suggested that WB severity may be reduced by supplementing a higher ratio of arginine in broilers’ diet without any significant negative impact on the overall breast meat yield. This could be because of the effect of arginine on the capillary vasodilation (Bautista-Ortega and Ruiz-Feria, 2010). On the contrary, TRIM63/MuRF1 upregulations involved in the atrophy of muscle cells was suggested to decrease the weight gain in broiler chickens fed modified methionine diets (Bodine and Baehr, 2014; Sachs et al., 2019). In a similar report on the modification of feed regimen, higher lysine supplement increased WS and WB occurrence in boilers (Cruz et al., 2016). Lysine is an essential amino acid for broiler growth performance and proper muscle development (Sterling et al., 2006). The explanations here emphasized that muscle outgrows its physiological system like vascular tone, therefore leading to muscular damage. Altogether, these results indicated that AA has no direct effect on muscle defects.

**Minerals** Another notable disorder in myopathic muscles is the swelling of the tissues resulting from an intracellular build-up of inflammation, which induces osmotic dysregulations (Livingston et al., 2019). In a previous study, potassium’s role in electrolyte balance was investigated in broiler chickens (Livingston et al., 2019). The study concluded that decreased WB incidence in birds fed potassium supplement might result from improved vascular circulation and hydrogen residue reduction in the muscle cells. Similarly, Magnesium supplement also significantly reduced the appearance of WS and WB phenotypes in broiler chickens (Estevez and Petracci, 2019). Their result corroborates with the study that showed magnesium deficiency exacerbates lipid peroxidation linked with muscular myopathy (Liu et al., 2007). Accumulation of these oxidative alterations confirmed excess free radical formation, oxidative degradation, and damage to cells and tissues as apparent in muscle fiber myopathies (Abasht et al., 2016). For example, a metabolomic study on WS muscle samples suggested an increase in calcium level results from the accumulation of unsaturated fatty acid in a deprived oxygen environment (Boerboom et al., 2018). Therefore, minerals imbalance is potentially a consequence of myodegeneration in broiler breast muscles.

**Antioxidants**

In myopathic muscles, hypertrophy and hypoxia cause tissue injuries and oxidative stress, respectively (Mutryn et al., 2015; Kuttappan et al., 2017; Malila et al., 2019). According to the omics report, there is little doubt that hypoxia could potentially be a cause of WS/WB in chicken breasts (Malila et al., 2019). This proposition suggests that hypoxia could predispose muscle to reactive oxygen species (ROS) accumulations in muscle tissue cells (Malila et al., 2019). During the onset of WS, increased secretion of antioxidant markers was observed to decrease free radical molecule accumulation (Salles et al., 2019). On the contrary, the glutathione S-transferase and glutathione peroxidase activities were reduced in severe WS-affected breast muscles (Malila et al., 2019). Indeed, the increase in free radicals in the pectoral muscle contributes to WS in chicken (Zambonelli et al., 2016; Salles et al., 2019). Excess ROS accumulation exacerbates oxidative damage in muscle cell lesions, DNA, and protein denaturation. Similarly, it could further aggravate inflammatory molecules, which might initiate the appearance of WS and WB on chicken fillets (Powers et al., 2010). Besides, broiler chickens are most vulnerable to increased oxidative stress because their meats contain a relatively higher polyunsaturated fatty acid found in WS/WB-affected muscles (Decker et al., 2010; Abasht et al., 2016). Nevertheless, it is still uncertain if oxidative stress is the cause or the effect of WS and WB phenotype in fast-growth type chickens (Zambonelli et al., 2016; Boerboom et al., 2018).

Antioxidants, both biological and synthetic, prevent oxidative damage in adipose tissues and the gut immune system (Surai, 2007). Vitamin E represents an essential functional component in poultry feed and a potent antioxidant against lipid peroxidation in both plasma cells and skeletal muscle tissues (Smet et al., 2008; Gao et al., 2010; Guetchom et al., 2012). However, according to studies, antioxidants have negligible effects on excess free radical reduction in the muscle cells. Important antioxidants like vitamins C, E, and omega have been studied, but their results are inconsistent in mitigating the myopathy caused by oxidative stress. For instance, Kuttappan et al. (2012c) study showed an increased Vitamin E level did not decrease WS appearance in abnormal breast muscles. This might happened because WS is not directly linked with vitamin deficiency (Kuttappan et al., 2012c; Guetchom et al., 2012). On the contrary, Vitamin E (200 IU/kg) supplement in starter diets reduced two-thirds of WS severe and moderate scores from starter to growers’ phase, whereas the WB severity score was alleviated only at the starters phase (Wang et al., 2020). The disparity in these studies could be from the difference in strains of experimental
birds, management practices, or the accuracy level during grading of the phenotypic scores. From Wang et al. (2020) report, Vitamin E supplements could increase metabolic activities in broiler chickens at the starter phase. It might further alleviate excess ROS caused by oxidative stress, possibly through its scavenging properties (Miller et al., 1993; Volj et al., 2011; Mutryn et al., 2015). In all, the bioavailability of the vitamin E in muscle depends on the vascular tone (Velleman et al., 2003; Petracci et al., 2019; Velleman, 2019; Wang et al., 2020), and it may be argued that capillary tissue damage could also its availability in breast muscles (Rocha et al., 2016; Papah et al., 2017).

Even though Bodle et al. (2018) reported that vitamin C could alleviate WB, on the contrary, the treatment did not decrease the appearance of WS in the same study. From their study, Vitamin C detoxified oxygen-derived free radicals only in a severe degree (WB). On the other hand, in the less severe condition like WS, it is evident that vitamin C supplement has no impact on its appearance in breast fillets. The explanation for the observed disparity could be because the required level of vitamin C is biologically synthesized in normal metabolic conditions, and WS disorder occurs at the severe stage of hypoxic stress (Xiang et al., 2002), whereas white stripes phenotype starts to show on meat during the onset of these 2 breast muscle disorders (Kuttappan et al., 2017). This notion means that the effect of Vitamin C supplement against breast muscle myopathy can only be seen at the later degenerative stage of muscle myopathy (Bodle et al., 2018).

**BROILERS PECTORALIS MUSCLE COMPOSITION**

Skeletal muscle is among the most abundant tissues in the body. Moreover, skeletal muscle growth, like any other tissue, also depends on protein turnover (Greising et al., 2012; Anthony, 2016). Therefore, this section provides insight into muscle growth and repair systems relating to myopathies in broiler chickens. It also addressed the prospect of SC in maintaining muscle integrity.

**Muscle Development and Breast Muscle Myopathy**

Myogenesis in broiler chickens involves the embryonic proliferation and differentiation of mesodermal cells to form multinucleated primary muscle fibers. After primary myofiber formation, the secondary muscle fibers get attached to the adjacent of the primary myofibers. Thus, these muscular cell syntheses make up the hyperplasia phase, and it biologically terminates at the ends of the incubation stage. Generally, it is assumed that hyperplasia decline till the early posthatch stage (Smith, 1963).

The fusion of SC and adjacent muscle fibers serve as the basis of increased genetic material in postnatal muscles (Daughtry et al., 2017). Note, the muscle fiber diameter increased to more than five-folds in heavy chicken type (Dalle Zotte et al., 2017). However, the physiology of skeletal muscle can only sustain the overgrowth to a certain threshold. As a result, it becomes a challenge for vascular cells to adjust to hypertrophic muscle in the later phase of chicken breast muscle growth (Petracci et al., 2013). For this reason, the skeletal muscle compositions are vulnerable to muscular damage, as seen in abnormal muscles. Muscle defects occur because of altered muscle components in WS-affected breast muscles (Pampouille et al., 2019). From a previous study, creatine kinase, domicile in the muscle fiber, was found at an increased level in the plasma sample of heavy strain turkey breeds (Wilson et al., 1990). Histological data showed a near-complete absence of connecting tissue spacing between the perimysial and endomysial fascicles (Wilson et al., 1990; Velleman, 2015). These data support the hypothesis that SC replenishment is impaired to compensate for myofiber degradation in myopathic muscles.

**Satellite Cells and Muscle Regeneration**

The myopathies in chicken breast muscles occur because of the suppression of SC mediating the cell regenerative process, a peculiar feature of hypertrophied muscle (Daughtry et al., 2017). Satellite cell, known to be the precursor for myogenesis, supports muscular growth (Rehfeldt et al., 2011). Satellite cells also function as regenerative cells during muscular injury. Unfortunately, in vitro analysis indicates that SC lose their abilities to proliferate and differentiate with age and hypertrophy, leading to myodegeneration (Daughtry et al., 2017). The expression of the paired box protein (PAX7) gene supports the idea that SC activations participate in muscle repair against myopathy (Daughtry et al., 2017). PAX7 gene has a specific role as a proliferating SC (Zammit et al., 2004) and is required to maintain muscle homeostasis by regulating *myf5* and *myoD* expressions (Relaix et al., 2005).

Satellite cells are, in part, regulated by the physiological state of muscles. In a homeostatic condition, SC is found to be quiescent. However, when homeostasis is disturbed in hypertrophic muscle, SC proliferates and differentiates to support muscle repair. For SC regeneration to occur, vascular cells and other extrinsic factors like nutrients regulate its regenerative power. Evidence showed that SC proliferation relies on the vascular tone in the muscle (Christov et al., 2007; Rhoads et al., 2009). However, owing to the sparse density of mitochondrial in the breast (a fast-twitch) muscle, vascularization decreases in hypertrophic muscles. Accretion of muscle fibers also reduces the available space for connective tissue, contributing to poor vascularization. In all, increased muscular repair and poor vascular tone impede the SC proliferation to support muscular regeneration in fast-twitch muscle growth (Velleman, 2015). The inability of SC to proliferate might cause necrotic lesions (Velleman et al., 2018). Therefore, we believe that the
impaired SC compromises the biological mechanisms for fast growth in chicken under hypertrophic conditions. A possible way to enhance the SC proliferation in hypertrophic muscle is by identifying potential molecular markers linked to muscular tissues. Most of the isolated genes from abnormal muscles implicate adiposis, muscle fiber regeneration, and repair processes (Petracci et al., 2013). Gene upregulations such as CSRP3 and platelet-derived growth factor receptor genes (PDGFRα) in myopathic breast muscles (Zambonelli et al., 2016; Pampouille et al., 2019) indicated that muscle regeneration processes were activated in response to the histological changes in breast muscles. During muscle repair activities in abnormal muscles, the PDGFRα gene helps muscle cells differentiate into fibroblasts and adipocytes. Another important marker in the muscle is the collagen type VI α 3-chain. Collagen type VI α 3-chain is a collagen molecular marker for extracellular matrix, and any perturbation in their expressions may cause necrosis as apparent in WS-affected muscles (Pampouille et al., 2019; Papah and Abasht, 2019). Such genes play critical roles in preserving the structural integrity of muscle tissues. Over-expression of the proteoglycan gene (IMPG2) peculiar to collagen was also reported to be a causative agent of fibrosis in WS-affected breast muscles (Zambonelli et al., 2016). These perturbations proved that alterations in the cell repair gene expressions could aggravate progressive muscular tissue degenerations (Spence et al., 2002).

Satellite cell differentiation, a multipotential cell population, also relies on muscle fiber types and myogenic expression changes in abnormal muscle cells. Muscle switching from type IIb to type I fibers is suggested as one of the many responsive mechanisms in myopathic muscles (Mutryn et al., 2015). The presence of type I fiber in the PM region indicates that regeneration occurs in dystrophic muscles. The switching process could help relieve muscular stress because type-I fibers contain a higher proportion of SC numbers. For example, myoglobin—a globular protein that carries oxygen to oxidative muscles—was expressed in a high proportion in WB-affected breast meat (Mutryn et al., 2015). Not only myoglobin but also troponin I and myosin binding protein C1 were reportedly upregulated in the damaged muscle caused by WB myopathy. For these reasons, type I fibers’ presence may be justified because they contain more capillary cells and thus relieve the stress from the rapid growth of the pectoralis region. This idea aligns with fiber-type switching previously linked with pale soft exudative muscle in turkey (Malila et al., 2013). In our opinion, we suggest that the focus of future research should be on strategies that can extend the threshold of SC in fast-growth type chickens.

**Muscle Vascularization**

The PMM primarily consists of fast-twitch fibers characterized by large muscle fiber diameter and metabolic stress (Hoving-Bolink et al., 2000; MacRae et al., 2006; Lisrat et al., 2016). Indeed, the myofiber size determines the available space for total vascular density within the muscular region (Joiner et al., 2014; Radaelli et al., 2017). During hypertrophy, the insufficient space of blood vessels further exacerbates oxidative stress in muscles. Histologically, the early stage of WB samples was characterized by inflammation of the blood vessels that increased progressively over time (Papah et al., 2017; Silvo et al., 2017). During muscular damage, angiogenesis promotes SC’s ability to proliferate (Yin et al., 2013). On the contrary, poor vascularity, as seen in both WS and WB muscles, cannot support SC regenerations during muscle repair processes (Luque et al., 1995; Christov et al., 2007; Rhoads et al., 2009). Studies have shown that the fate of SC and capillary cells are intertwined (Luque et al., 1995; Christov et al., 2007; Rhoads et al., 2009), such that SC stimulates vascular cell development in the muscle region (Rhoads et al., 2009). However, when the muscle fiber is enlarged, it constrains the blood vessels’ vascularity and affects SC response (Siller, 1985). Accordingly, it is possible to argue that impaired vascularity predisposes WS-affected breast muscles to polyphasic myodegenerations (Hoving-Bolink et al., 2000; Berri et al., 2007), meaning that the solution to enhanced muscle vascularity combined with SC proliferation could potentially solve the concerns of alleviating WS-affected breast muscles.

**FUTURE PERSPECTIVES ON IN OVO FEEDING, SATELLAITE CELLS, AND VASCULAR TONE**

In line with other studies, we believed that poor vascular tone affects WS and WB developments in broiler chickens. Therefore, it is necessary to share ideas on possible ways to improve vasodilation in fast-growth type chickens. Nitric oxide (NO), an endothelium-derived relaxing factor, can be synthesized biologically from L-arginine (Vallance et al., 1989b) by nitric oxide synthases (Louis et al., 1987; Vallance et al., 1989a). Biologically, NO is a signaling molecule that dilates the blood vessel and relaxes smooth muscle cells, thus favoring increased blood flow in the blood vessels (Latchman, 1995). During vasodilation, NO metabolic role is to activate soluble guanylyl cyclase. Then, the soluble guanyl cyclase triggers the activation of cGMP protein kinase. Subsequently, the protein kinase stimulates a cascade of metabolic processes that decrease intracellular calcium and promote breast muscle vascularity (Marchesi et al., 2019). According to this mechanism, NO might alleviate the poor vascularization in fast-growth chickens. Moreover, Klinger and Kadowitz had previously demonstrated that L-arginine stimulated the nitric oxide synthase pathway to enhance NO biosynthesis (Klinger and Kadowitz, 2017). On this principle, we may hypothesize that enhanced NO could alleviate WS phenotype (Figure 3) (Andriantsitohaina et al., 2012). Nitric oxide low bioavailability has been previously linked with hypoxia-related ischemic diseases (Kevil and Patel, 2008; Napoli and Ignarro, 2009). Fortunately, L-arginine supplementation can biosynthesize
endothelial nitric oxide synthase, an enzyme that contributes to the relaxation of the vascular tone in breast muscle fiber. L-arginine, a precursor of NO, reportedly relieved hypoxic conditions by improving vascular tone in broilers (Bautista-Ortega and Ruiz-Feria, 2010). A related study also showed that dietary arginine improved flow-dependent vasodilation and reduced pulmonary vascular resistance (Wideman et al., 1995). These results indicated that L-arginine could promote vasodilation and, in turn, increases blood pressure and blood flow. Another concern is the impairment of SC; however, muscle repair processes during degeneration have been suggested to resemble embryonic myogenesis (Yin et al., 2013). Therefore, by considering myogenesis at the early stage, SC could be improved to sustain hypertrophy in broiler chickens. On this account, enhanced SC proliferation in ovo could potentially support the growth pace in broiler chickens. This idea calls for the need to devise in ovo strategy to achieving efficient growth in broiler chickens.

In ovo feeding is designed to improve muscle development and poultry production efficiency (Zhao et al., 2017). In fact, in ovo injection of L-arginine has previously been used to enhance posthatch growth performance and regulate protein synthesis and deposition through the mTOR pathway and its downstream target proteins intermediate (Gao et al., 2017; Yu et al., 2018). The efficiency of in ovo injection of L-arginine has also been investigated in Japanese quail’s physiological trait (Al-Daraji et al., 2012), small intestine immune barrier function, and lymphoid organ development (Gao et al., 2017). In a similar study, L-arginine in ovo injection stimulated the myogenin gene expression in posthatch chicken cultured tissue through the mediation of NO (Li et al., 2016). Therefore, we suggest that in ovo supplementation of L-arginine could be relevant in solving muscle myopathies by enhancing SC and angiogenesis during in ovo and early posthatch phases. Future studies should focus on new ways to design in ovo feeding to improve the meat quality and maintain the desired growth rate in broiler chickens.

CONCLUSION

Because of better feeding efficiency, fast-growing chickens have proven profitable on a commercial scale for meat products. Unfortunately, the growth rate of broiler chickens leads to muscle physiology perturbation; in other words, the rapid growth rate could not be sustained by the chicken biological system, especially the SC and vascular system. Consequently, some muscle defects, including growth-induced myopathies like WS phenotype, have become a great concern for broiler chicken farmers. Hence, this article provides an extensive review of possible breast muscle myopathy etiologies. This article, in fact, buttress research studies on muscle degeneration in the pectoralis major of fast-growing broilers, with the idea that early preparation of both vascular system and SC could potentially promote vasodilation and support processes of muscle regeneration, respectively. For these reasons, more studies should explore how nutritional strategies can help mitigate the defect in meat fillets. In particular, consideration

![Figure 3. Conceptualization of improved vascularity in fast-growing broiler chickens. (A) During the incubation phase, the number of muscle cells increases (hyperplasia) in ovo and then forms the myofi bers. (B) The enlargement of muscle fi bers in fast-growing chickens hinders the vasodilation of vascular tissues and thus disrupts the biological support mechanism of pectoralis major muscle. (C) Satellite cells proliferate synergistically with the development of blood capillaries. The secretion of endothelial-derived relaxing factors improves muscle vascularity. The enhancement of breast muscle vascularization suggests promoting satellite cell proliferation that can support the muscle growth pace in broiler chickens.](image-url)
should be given to in ovo feeding trials such as L-arginine in future research, as argued in this article.

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DISCLOSURES

The authors declare no conflicts of interest.

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