**HEAT STRESS IN POULTRY**

***Vo Van Su (collection)***

**Animal Husbandry Association of Viet Nam**

Corresponding author: Vo Van Su. Tel: 0989382311. Email: vovansu.vcn@gmail.com

**ABSTRACT**

Heat stress is a common reality in poultry production, its effects are quite complex and harmful and depend on the intensity and duration of the exposure to high temperatures.

The gut is affected by heat stress through several pathways, including organ ischemia and hypoxia, as well as oxidative stress.

In heat stress challenges, the intestinal barrier is compromised because of lower tight junction protein expression, enterocyte damage, and microbiome unbalance, leading to gut health issues such as dysbiosis and necrotic enteritis.

At the gut level, phytomolecules such as carvacrol, cinnamaldehyde, capsaicin, silymarin, cineol, and menthol, among others, have been found to alleviate heat stress through their antioxidant capacities, leading to improved animal health and performance.

**Keywords**: *heat stress, avian, oxidative, intestinal*

**INTRODUCTION**

Stress in animals can be defined as any factor causing disruptions to their homeostasis, their stable internal balance. Stress engenders a biological response to regain equilibrium (Das et al., 2011). We can distinguish four major types of stress in the poultry industry: technological or management-related stress; environmental stress; nutritional stress, including due to heavy metals, mycotoxins, and low-quality ingredients; and internal stress, which is related to health status and health challenges. (Surai et al., 2019). All types of stress lead to molecular and cellular changes that decrease health and productivity.

**Climate change, thermoregulation, and stress**

High environmental temperatures are among the most important environmental stressors for poultry production, causing significant economic losses in the industry (St-Pierre et al., 2003). Climate change has increased the prevalence and intensity of heat stress conditions in most poultry production areas all over the world (Tellez et al., 2017;Lian et al., 2020)

The optimum temperature for poultry animals well-being and performance – the so-called thermoneutral zone – is between 18 and 22°C. When birds are kept within this temperature range, they do not have to spend energy on maintaining a constant body temperature (Akbarian et al., 2016).

Heat stress is the result of unsuccessful thermoregulation in the animals, as they absorb or produce a higher quantity of heat than they can lose. It means that there is a negative balance between the net amount of energy flowing from the animal to the environment and the energy it produces (Lara and Rostagno, 2013).

**Heat stress – contributing factors**

This energy imbalance is influenced by environmental factors such as sunlight, thermal irradiation, air temperature, humidity, and stocking density, but also by animal-related factors such as body weight, feather coverage and distribution, dehydration status, metabolic rate, and thermoregulatory mechanisms (Lara and Rostagno, 2013; Saeed et al., 2019). When the environmental temperature is above the thermoneutral zone, the animals activate thermoregulation mechanisms to loose heat through behavioral, biochemical, and physiological changes and responses (Farag and Alagawany, 2018;Quinteiro-Filho et al., 2010; Santos et al., 2015;Awad et al., 2018).

Heat stress can be classified into two main categories, acute and chronic. Acute heat stress refers to a short and fast increase in environmental temperature (a few hours), whereas under chronic heat stress the high temperatures persist for more extended periods (several days). Some studies suggest that, in some circumstances, poultry animals show a degree of resilience to acute heat stress (Lara and Rostagno, 2013;Farag and Alagawany, 2018;Quinteiro-Filho et al., 2010). However, in the long-run, their compensatory mechanisms are not sufficient to maintain tissue integrity and thus health and performance (Santos et al., 2015).

**The animal’s response to heat stress**

The exposure of poultry to heat stress changes the gene expression of cytokines, upregulates heat shock proteins (HSP), and reduces the concentration of thyroid hormones (Quinteiro-Filho et al., 2010, Awad et al., 2018). When heat stress persists, these cascades of cellular reactions result in tissue damage and malfunction.

The animals exposed to heat stress suffer adverse effects in terms of performance, which are widely known and include high mortality, lower growth and production (Figure 1), and a decline in meat and egg quality (Mujahid et al, 2005;Hu et al., 2019).



Figure 1. Body weight gain of broilers exposed to chronic heat stress (35°C continuously from day 21). A marker for tight junction permeability was added to feed (FITC-d - fluorescein isothiocyanate dextran); its fluorescence (in serum) increased with heat stress exposure time, showing higher intestinal permeability. (Adapted from Ruff et al., 2020)

**Oxidative stress – a consequence of heat stress**

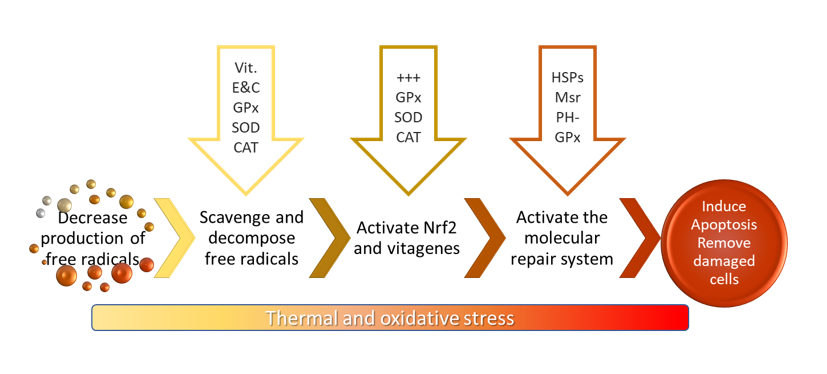
Oxidative stress, simply put, occurs when the amount of reactive oxygen species (ROS – such as superoxide anions, hydrogen peroxide, and hydroxyl radicals) exceeds the antioxidant capacity of the cells (Akbarian et al., 2016;Hu et al., 2019; Salami et al., 2015). Oxidative stress is regarded as one of the most critical stressors in poultry production as it is a response to diverse challenges affecting the animals (Surai et al., 2019;Surai and Fisinin, 2016)

At a cellular level, the metabolism of the animal – its energy production – generates ROS and reactive nitrogen species (RNS), such as hydroxyl radicals, superoxide anions, hydrogen peroxide, and nitric oxide. These usually are further processed by antioxidant enzymes produced by the cell (Surai et al., 2019; Salami et al., 2015), including superoxide dismutase (SOD), catalase (CAT), and glutathione peroxidase (GSH-Px). Nutrients such as selenium and vitamins E, C, and A also participate in antioxidant processes (Surai et al., 2019; Lian et al., 2020). When the generation of ROS exceeds the capacity of the antioxidant system, oxidative stress ensues (Surai et al., 2019; Lauridsen, 2019).

Heat stress leads to higher cellular energy demand, promoting the generation of ROS in the mitochondria (Mujahid et al., 2005.), which exceed the antioxidant capacity of the organism. As a consequence, oxidative stress occurs in several tissues, leading to cell apoptosis or necrosis (Santos et al., 2015). Among these tissues, the gastrointestinal tract can be highly affected.

Oxidative stress damages cell proteins, lipids, and DNA, and reduces energy generation efficacy (Akbarian et al., 2016). Moreover, oxidized molecules can take electrons from other molecules, resulting in a chain reaction. If not controlled, this reaction can cause extensive tissue damage (Lauridsen, 2019).

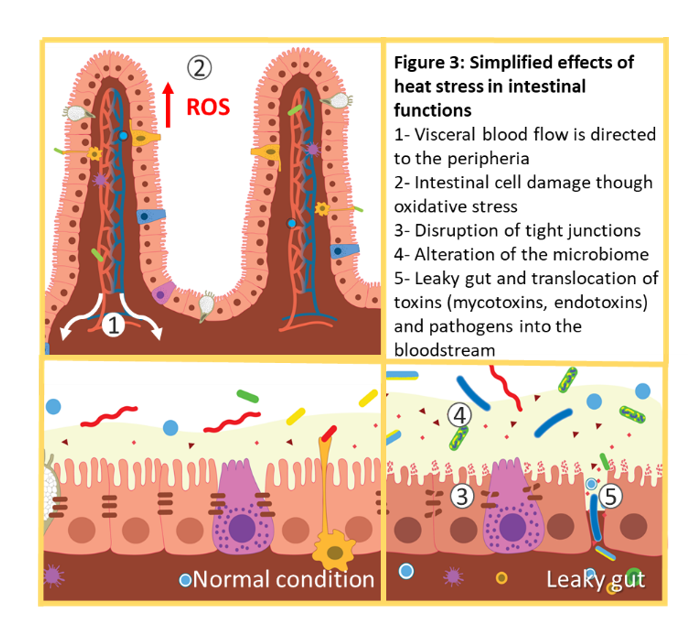
In response to oxidative stress, all antioxidants in the organism work together to re-establish homeostasis. Several steps in the oxidative stress response have been identified. Whether they take place depends on the intensity of the stressor, with ROS and RNS acting as signalling molecules. These steps include the internal synthesis of antioxidants, the activation of transcription factors or vitagenes, and the production of protective molecules (Figure 2).

Figure 2. First, decrease free radical production by decreasing oxygen availability and reducing the activities of enzymes responsible for ROS production (NADPH oxidase). Second, scavenge and decompose free radicals through natural antioxidants (vitamins E & C, GSH, SOD, GPx, and CAT). Third, activate Nrf2 and vitagenes to further stimulate the synthesis of antioxidants. Fourth, activate enzymatic systems responsible for damaged molecule repair (HSP, Msr, DNA-repair enzymes) and removal (PH–GPx). Fifth, induce apoptosis and other processes to deal with terminally damaged cells. (Adapted from Surai  
et al., 2019)

**Oxidative stress’ effects on the gut**

In the gastrointestinal tract, oxidative stress and the consequent tissue damage lead to increased intestinal permeability. This facilitates the translocation of toxins and pathogens from the intestinal tract into the bloodstream (Figure 3).

Under oxidative stress conditions in the gut, there is a demand for antioxidants to counteract the excess of ROS; hence, dietary antioxidants can help reduce ROS and improve animal performance (Salami et al., 2015). Research shows that certain phytomolecules have antioxidant properties and improve performance under conditions of oxidative stress (Hu et al., 2019; Arab Ameri et al., 2016; Saadat Shad et al., 2016; Mishra and Jha, 2019).



**Thermoregulation: changes in blood flow**

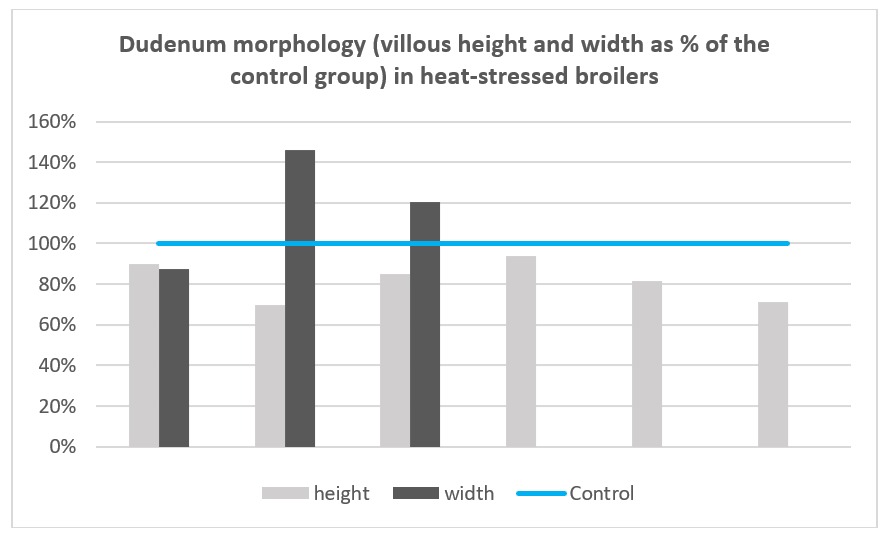
The gastrointestinal tract is profoundly affected by heat stress: to help with heat dissipation, the thermoregulatory mechanism of the animal shifts visceral blood flow towards peripheral circulation. Organ ischemia and hypoxia follow, limiting gut motility, nutrient utilization, and feed intake (Lian et al., 2020;Hu et al., 2019). Enterocytes are particularly sensitive to hypoxia and nutrient restriction, which leads to oxidative stress (Surai et al., 2019;Awad   
et al., 2018).

**Changes in intestinal barrier’s tight junctions**

Several studies indicate that both acute and chronic heat stress increase gut permeability, partly by increasing oxidative stress and by disrupting the expression of tight junction proteins (Lian et al., 2020; Ruff et al., 2020). Heat and oxidative stress in the gut result in intestinal cell injury and apoptosis. When the tight junction barrier is compromised, luminal substances leak into the bloodstream, which constitutes the condition described as “leaky gut”( Tellez et al., 2017; Ruff et al., 2020).

**Changes in intestinal morphology**

Heat stress affects intestinal weight, length, barrier function, and microbiota, resulting in animals that have lower total and relative weight of the small intestine, with shorter jejunum and duodenum, shorter villi (Figure 4), and reduced absorption areas, in comparison to non-stressed animals (Santos et al., 2015; Awad et al., 2018; Abdelqader and Al-Fataftah, 2016; Jahejo et al., 2016; Wu et al., 2018; Santos et al., 2019).

Figure 4. Villous height and width of broilers exposed to heat stress in relation to the control group (100%). Villous height is always shorter than the control group, but width can increase when the organism shows resilience to the stressful situations and aims to recover intestinal surface. (Adapted from Jahejo et al., 2016; Santos et al., 2019; Wu et al., 2018; Abdelqader et al., 2016 ; Santos et al., 2015 and Awad et al., 2018 – by order of appearance in the graph, from left to right)

**Changes in intestinal microbiome**

Due to reduced feed intake and impaired intestinal function, the presence and activity of the commensal microbiota can also be modified. Heat stress can lead to reduced populations of beneficial microbes. At the same time, it can boost the growth of potential pathogens and lead to dysbiosis, increased gut permeability, as well as immune and metabolic dysfunction (Shi et al., 2019). Burkholder et al. (2008) and Rostagno (2020) point out that pathogens such as *Clostridia*, *Salmonella*, and coliform bacteria increase in poultry exposed to heat stress, while the populations of beneficial bacteria such as *Lactobacilli* and *Bifidobacteria* decrease.

**Necrotic enteritis**

Heat stress causes damage in the gut microbiota, intestinal integrity, and villus morphology, as well as immunosuppression. Consequently, feed digestion and absorption decline (Santos et al., 2015; Awad et al., 2018; Burkholder et al., 2008; Quinteiro-Filho et al., 2012.). These factors increase the risk of necrotic enteritis outbreaks (Lian et al., 2020; Burkholder et al., 2008; Antonissen et al., 2014; Tsiouris et al., 2018.), one of the most problematic bacterial diseases in modern poultry production.

In a study by Tsiouris et al. (2018), cyclical acute heat stress was found to increase the incidence and severity of necrotic enteritis in broilers challenged with *C. perfringens*, and to produce the disease in animals that were not exposed to the bacteria. Other signs, such as growth retardation and a reduced pH of the intestinal digesta, were also observed in the heat-stressed birds.

By lowering feed digestibility, increasing gut permeability, and compromising immunity, heat stress leaves animals more susceptible to gut-health related issues such as dysbacteriosis and necrotic enteritis – and thus increases the need to use antibiotics.

**Mitigation strategies**

Most intervention strategies deal with heat stress through a wide range of measures, including environmental management, housing design, ventilation, sprinkling, and shading, amongst others (Saeed et al., 2019). Understanding and controlling environmental conditions is always a part of heat stress management: it is crucial for ensuring animal welfare and achieving successful poultry production.

Feed management and nutrition interventions are also recommended, together with environmental management, to reduce the effects of heat stress. They include feeding pelletized diets with increased energy, higher fat inclusions, reduction of total protein, supplemental amino acids, higher levels of vitamins and minerals, and adjusting the dietary electrolyte balance (Das et al., 2011; Awad et al., 2018; Arab Ameri et al., 2016). Nutrition is crucial, and the use of the right diets aids in attenuating heat stress in birds.

**Phytomolecules: powerful antioxidants**

It is practically impossible to avoid stress in commercial poultry production; hence it is common for animals to experience oxidative stress at times. Phytomolecules are natural antioxidants with anti-inflammatory and digestive properties (Saeed et al., 2019; Hu et al., 2019), which have been shown to improve poultry performance, [including during challenging periods](https://ew-nutrition.com/challenging-times-for-broilers/). The antioxidant capacity of phytomolecules manifests itself in free radical scavenging, increased production of natural antioxidants, and the activation of transcription factors (Surai et al., 2019;Abd El-Hack et al., 2019; Surai, 2020).

As compounds that have low bioavailability, they can remain at high concentrations within the intestine, when provided at the appropriate dosage and through encapsulation technology. Research has found that phytomolecules can effectively reduce intestinal ROS and thus alleviate heat stress in poultry (Salami. et al., 2015;Arab Ameri et al., 2016; Saadat Shad et al., 2016., Mishra and Jha, 2019), specifically mitigating oxidative stress in the intestine.

One heat stress study, for example, found that carvacrol elevates serum GSH-PX activity, compared to non-supplemented broilers (Saadat Shad et al., 2016). Other studies demonstrate that cinnamaldehyde also increases the activities of natural antioxidants in heat-stressed broilers (Abd El-Hack et al., 2019;El-Maaty et al., 2014). A study by Prieto and Campo (2016) showed that dietary supplementation of capsaicin effectively alleviated heat stress, as indicated by a lower H/L ratio in supplemented animals.

Silibinin, a flavonolignan present in silymarin (milk thistle extract), is another powerful antioxidant. In the gastrointestinal tract, it can come into direct contact with cells, activating transcription factors such as Nrf2, and thus helping to upregulate the antioxidant protection (Surai, 2015). Other phytomolecules, such as menthol and cineol, also aid animals under heat stress by simulating the sensory cold receptors of the oral mucosa. This gives them a cooling sensation and reduces heat stress behavior (Arab Ameri et al., 2016.).

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