

Effect of Adding Guanidinoacetic Acid to Semen Diluents, Duration of *in Vitro* Storage on Semen Quality, Hatchability of Iraqi Chickens

Waleed Khalid A. AL – Hayani

Department of Animal Production, College of Agriculture, University of Baghdad, Iraq.

ABSTRACT

This study was conducted to examine the effect of adding different levels of Guanidinoacetic Acid (GAA) to the semen Diluents of the Iraqi rooster in the semen quality characteristics and the hatchability of two Iraqi chicken breeds during different storage periods *in vitro*. It was used 80 Cockerels (40 white + 40 naked red), and 256 chickens (128 white + 128 naked red) at 29 age weeks. Birds of each strain were divided in to four groups, these groups including concentration (Conc), 0, 1, 2, 3% of GAA were added to semen. Then, stored for four *in vitro* storage (SP) 0, 24, 48, 72 hours at a temperature of 4 C^o. The results indicated a significant increase in mass motility (MM) and individual motility (IM) hatchability (Ha) with significant reduction of Dead sperm (DS), abnormal spermatozoa (AS) and Acrosomal abnormalities (AA) when adding GAA with concentration Br and Conc in previous qualities. While a significant effect of interaction was found between Br and SP with significant effect of interaction and Br and Conc and SP. It is therefore possible to conclude that adding GAA to sperm diluents improves sperm quality and hatchability characteristics. Therefore, GAA can be used to improve the productive qualities of Iraqi chickens.

Key Words: Guanidinoacetic Acid, Semen diluents, in vitro storage, Semen quality, Iraqi chickens.

eIJPPR 2017; 7(1):12-21

HOW TO CITE THIS ARTICLE: Waleed Khalid A. AL – Hayani. (2017). "Effect Of Adding Guanidinoacetic Acid To Semen Diluents, Duration Of *In Vitro* Storage On Semen Quality, Hatchability Of Iraqi Chickens ", *International Journal of Pharmaceutical and Phytopharmacological Research*, 7(1), pp.12-21.

INTRODUCTION

The local Iraqi breeds of chicken were characterized by a decrease in their reproductive efficiency, reflected in a decrease in local production of egg and meat [1]. Therefore, the studies recommended artificial insemination of birds in Iraq to overcome the problems of production and decrease in the local breeds, as the adoption of artificial insemination provides the possibility of election cocks and chicken with good production and rejection of the weak [2]. Then, obtaining sperm with good specifications and desirable quality and hatching rates [3].

In spite of the advantages of artificial insemination in birds, it focuses on a number of problem such as the short life of sperm *in vitro*, and the short time of

Corresponding author: Waleed Khalid A. AL – Hayani

Address : Department of Animal Production, College of Agriculture, University of Baghdad, Iraq.

e-mail 🖂 waleed.khaled@coagri.uobaghdad.edu.iq

Relevant conflicts of interest/financial disclosures: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

Received: 29 August 2016; Revised: 11 February 2017; Accepted: 26 February 2017

²

artificial insemination, which does not exceed 10 - 15 min due to the rapid rates of high metabolism of birds [4], which opened the doors to using of Diluents to storage semen, to prolong the life of sperm and prevent the causes of death, the most important are free radicals [5]. Therefore, it is important to search for every metabolite, add to the sperm diluents that are presented by energy, reduce the activity of free radicals and then extended he length of storage [6, 7]. The hypothesis of sperm diluents is based on providing an environment that allows prolongation of life intravenous insomnia provides energy substrate to sustain metabolism or antioxidants that inhibit free radicals during storage [8]. In this sense, it was necessary to search for materials that are equipped with energy – efficient and one of these materials Guadinoacetic Acid [9].

Guadinoacetic (GAA), or Guanidinoacetate or Glycocyamineo is derived from the reaction of arginine and glycine in the liver and kidney [10]. Contributes to the synthesis of creation (Ct) in the reactions of the transfer of the group of the instance of S – adenosy methionine to GAA [11], Ct is a high energy phosphate that is important in the metabolism of the body's energy, especially in the male reproductive system [12], as Vigue et al. [13] report the Ct is responsible for the transfer of ATP energy to the sperm tail (sperm), to provide adequate energy from sperm movement.

The disturbance in energy metabolism reduces the mobility of the sperm and the movement of the sperm is a key factor of fertilization [14]. Tapeh et al. [15] reported that the use of GAA in roosters' diets improved the IM and MM and the reduction of DS, AA, AS. So that, this study examines the possibility of adding GAA to sperm diluents and indicates its role in semen, fertility and hatchability at four SP for two Iraqi Br.

MATERIALS

The study was carried out in the Poultry Research center, Agricultural Research office, Ministry of Agriculture, Iraq. In the period of time from 1/2/2016 to 1/3/2016 to study the effect of adding different levels of the GAA to Semen diluents of Iraqi rooster in the semen quality and hatchability for two Br from Iraqi Chicken's during different *in vitro* storage. **Experimental design and treatments**

This study involved 80 Cockerels (40 white + 40 naked red), and 256 hens (128 white + 128 naked red) at the age of 29 weeks, these birds were housed in halls their environmental conditions perfect, and given diet contained 2850 Kcal and 16.71% crud protein. Birds were divided into four groups, included adding the levels 0, 1, 2, 3% of the GAA to sperm Diluents. **Sample collections and preparations**

Then collecting sperm by massage the abdominal area [16] by using semen suction device [17], reduced sperm using Lake diluent [18], after the preparation of

GAA solution by dissolving 1gm of GAA in 10 ml distilled water, and soluble fully complete in size to 100 ml, then added 1, 3, 5 ml of aqueous solution of GAA to semen diluted by Lake diluents [18]. And then stored semen samples in vitro under the temperature of 4 C^o for the SP 0, 24, 48, 72 h, the following tests were carried out at the previous periods. The microscope used to study the qualitative characteristics to sperm during the four SP (0, 24, 48, 72 h) to calculate the collective MM on the strength of the 40x and IM on 100x [19] and the percentage of DS by uses of Eosin nigrosin stain [16], and the percentage of AS and AA by using Fast Green - Eosin stain [20, 21]. Then, artificial insemination Hens by using 0.05 of diluents / hen [2, 17], the lap of the eggs of the product for the following days, the second day of artificial insemination and reached 50 eggs/group, after 21 days Ha calculated by the following equation:

Chicks hatched

Hatchability % = ----- × 100 Total eggs

Statistical analysis

Analysis data according to factorial experiment in Completely Randomized Design –CRD to effect of different factors (GAA, Br, SP) in qualities under study. Statistical analyses were performed using the All Statistical Package for Social Science [22], version 21.0 for windows (SPSS Inc., Chicago, IL, USA) according to a Complete Randomized Design. Means were compared by Duncan's Multiple Range Test [23] with a significance level of 5%.

RESULTS

Mass motility

Table 1 show that there is no significant effect for Br in MM, but the addition of the GAA in Conc_{1, 3, 5} to the semen diluent had significant increase of MM (P<0.01) compared to Conc_{.0}, while noting the significant decrease (P<0.05) in MM when increasing storage periods (SP), as the best MM reached at SP₀ and then SP₂₄ and SP₄₈ and SP₇₂ h respectively.

It also indicated that of interaction between Br and Conc. in MM. However, the interaction between Br and SP reached the level of significant (P<0.05), noting that the interaction between Br and SP₀ achieved the best MM, then Br With SP₂₄ trailed by interaction Br with SP₄₈, the interaction between Br and SP₇₂ has recorded less than with regard to the interaction between the Conc and SP notable deterioration of the MM with the increase of the SP significantly (P<0.05), noting that it absorbs the interaction of the four (0, 1, 3, 5%) achieved best MM when interaction with SP₀. While the interaction between Conc₀ and SP₇₂ less than MM. The interaction between it Conc_{1, 3, 5} with durations of 24, 48, 72 h achieved MM best interaction of Conc₀ with the same duration. At the same time notes that interaction each of Br and Conc and SP₀ achieved the best MM interaction with other SP, then the interactions of Br and Conc with the SP₂₄, and thus interaction SP₄₈, and

that the interaction with the SP_0 indicated lower levels of significant (P<0.05). Also, the interaction of Br and Conc₀ with the SP_{72} recorded the lowest significant (P<0.05).

Individual motility

As can be seen from table 2 that there are no significant differences between Br1 and Br2 in IM, as indicated above $Conc_{1, 3, 5}$ significantly (P<0.01) on the $Conc_0$ in IM, with the significant decrease (P<0.01) in IM when SP24, 48, 72 when compared with SP0. no significant difference of the interaction between the Br and Conc, with the interaction significant (P<0.05) between Conc and SP, noting that the Br_{1,2}SP₀ indicated significant increase (P<0.05) IM interactions with the other, followed by Br_{1,2}SP₂₄, then Br_{1,2}SP₄₈ and then Br_{1,2}SP₇₂ with regard to the interaction between the Conc and SP notes that the Conc₀, 1,3,5SP₀ indicated significant increase (P<0.01) in IM compared to other groups, while noting that the Conc_{1,3,5}SP_{48,72} indicated significant increase (P<0.01) compared with Conc₀SP_{48,72}. When the interaction of the three factors notes from the same table (table 2): Br₁Conc₁SP₀ and Br₂Conc₀SP₀ a record significant increase (P<0.05) in IM compared to other groups. The Br₂Conc₅SP₂₄ did not differ from the significant of Br_{1,2}Conc._{0,1,3,5}SP₀. While Br_{1,2}Conc₀SP₄₈ indicated significant decrease (P<0.05) in IM compared to Br₂Conc₅SP₄₈ and significant decrease (P<0.05) Br_{1,2}Conc₀SP₇₂ comparing to Br_{1,2}Conc_{1,3,5}SP_{48,72}.

Dead sperm

As can be seen from table 3 that there is no significant differences in DS between Br₁ and Br₂, as shown in the low significant (P<0.01) in the DS in favor of $Conc_{1,3,5}$ compared with Conc₀, also notes the significant decline (P<0.01) in the SD in favor of the SP_0 compared to SP0,24,48 and the SP24 recorded less SD than SP48,72 and SP₇₂ posted a significant increase (P<0.01) compared to SP₄₈ in SD. While not observed significant differences in interaction between Br and Conc. It also notes that the Br_{1,2}SP₀ recorded significant decline (P<0.05) in SD compared to groups remaining, and Br_{1,2}SP₂₄ Best of Br_{1,2}SP₄₈ and Br_{1,2}SP₇₂ recorder the highest rate(P<0.05) of SD compared with other groups. It also noted from table 3 that the ConcoSPo recorded a significant increase (P<0.05) in SD compared to all interactions, including Conc_{1,3,5}SP₇₂, as well as notes that Conc_{0,1,3,5}SP₀ the lowest SD compared to other interactions. As can be seen from table 3 that the Br_{1,2}Conc₀SP₇₂ significant decrease (P<0.05) in SD compared to all interactions especially Br_{1,2}Conc_{1,3,5}SP₇₂.

Abnormal spermatozoa

It is clear from table 4 that there is no significant difference between Br_1 and Br_2 in AS, with the existence of a significant decrease (P<0.01) in AS for $Conc_{1,3,5}$ compared $Conc_0$. At the same time noted the increase in AS significant (P<0.01) progress of the SP. No

significant differences interaction between Br and Conc in AS. While noting that the Br_{1,2}SP₀ recorded a lower average (P<0.05) AS compared to all interactions followed by Br_{1,2}SP₂₄ and then Br_{1,2}SP₄₈ and then $Br_{1,2}SP_{72}$. It also notes that the $Conc_5SP_0$ significant decrease (P<0.05) in AS compared to the groups remaining, while the Conc₀SP₇₂ the highest values (P<0.05) AS compared to the rest of the intersections. As can be seen from the same table that Br₁Conc₅SP₀ significant decrease (P<0.05) compared to Br_{1,2}Conc_{0,1,3,5}SP_{48,72} and Br₁Conc_{0,1,3}SP₂₄ and Br₂Conc_{0,3}SP₂₄. In the same time in which the Br_{1,2}Conc₀SP₇₂ higher significant (P<0.05) values of AS compared to other interactions.

Acrosomal abnormalities

Table 5 indicates the absence of significant differences between Br₁ and Br₂ in AA, with the existence of a significant decrease (P<0.05) in AS for Conc_{1,3,5} compared Conc₀, and the significant increases of the AS registered progress of SP. At the time did not observe the significant difference of the interaction between the Br and Conc in the AA as indicated the existence of a significant decrease (P<0.05) in favor of Br_{1,2}SP₀ compared to $Br_{1,2}SP_{24,48,72}$ in AA, and significant decrease (P<0.05) in favor of Br_{1,2}SP₂₄ compared to Br_{1,2}SP_{48,72} and Br_{1,2}SP₄₈ record the less significant (P<0.05) in AA of Br_{1,2}SP₇₂. The table also indicates Br_{1.2}Conc_{0.1.3.5}SP₀ had achieved a significant decrease (P<0.05) in the AA compared to $Br_{1,2}Conc_{0,1,3,5}SP_{24,48,72}$. and Br_{1,2}Conc₀SP₇₂ posted a significant increase (P<0.05) in AA compared to groups remaining. The Br_{1,2}Conc_{1,3,5}SP₄₈ recorded significant decrease (P<0.05) in AA compared to $Br_{1,2}Conc_0SP_{48}$ and the Br_{1,2}Conc_{1,3,5}SP₇₂ record low significant (P<0.05) in the AA compared to Br_{1,2}Conc₀SP₇₂.

Hatchability

Table 6 indicated that there is no significant effect Br in Ha, but that the use of GAA $Conc_{1,3,5}$ had led to a significant increase (P<0.01) in Ha compared to Conc₀. Increasing the duration of storage of 0 to 72 h led to significant decrease (P<0.01) in Ha, the SP₀ first and then the SP_{24} and SP_{48} and SP_{72} . It notes the no significant effects of the interaction between the Br and Conc. The significant effects were observed interaction between Br and SP, as Br_{1,2}SP_{0,24} excelled Significantly (P<0.05) on Br_{1,2}SP_{48,72}, and Br_{1,2}SP₄₈ than Br_{1,2}SP₇₂. Shows that Conc_{0,1,3,5}SP_{0,24} excelled significantly (P<0.05) on Conc.0,1,3,5SP48,72. The ConcoSP72 record less significant (P<0.05) than the value of Ha compared to other groups, no significant difference between Conc₅SP₇₂ and Conc₀SP₄₈, the significant decrease(P<0.01) in comparison with $Conc_{1,3,5}SP_{48}$. It is the same table clear Br₂Conc₁SP₀ significant increase (P<0.05) on Br₂Conc₀SP₂₄ and Br_{1,2}Conc_{0,1,3,5}SP_{48,72}, without significant effects differ from other interactions. In the record of Br_{1,2}Conc₀SP₇₂ less than the values per Ha compared to the rest of the intersections. The Br_{1,2}Conc_{1,3,5}SP₄₈, recorded a

significant increase (P<0.05) in Ha when compared to $Br_{1,2}Conc_0SP_{48}$, and did not differ from the interactions of Br and Conc with the SP₇₂.

DISCUSSION

It also noted that the results showed no significant effect of Br on the specific characteristics of the specific characteristics of the semen and the hatchability of the previously mentioned results. This may be due to the convergence of the productive efficiency of the two breeds. The Iraqi breeds characterized by their yield [1].

The significant influence of Conc significant improvment is evident in MM and IM, which may be due to the fact that adding GAA to semen diluents contributes to the Create of Ct [11], which provides energy substrates for the sperm [13]. Lee et al. [24] has indicates that the seminal vesicles in the reproductive system of mammals produce Ct to be a major source of energy and an important responsible for the movement of the sperm [25]. Due to the absence of the attached gonads in the roosters, GAA may be a direct source of Ct extract after supplementation with semen driers. Ct has an anti – oxidant effect [26], which may be due to low SD, AS, AA, as well as simple signs of GAA's antioxidant roles [27, 28].

The GAA's anti – properties reduce concentrations of polyunsaturated fatty acids (PUFA) in the semen diluted, as these acids increase the likelihood of peroxides and thus reduce the fertilization capacity of the sperm. Peroxides reduce the life of span of sperm within the *in vivo* during inoculation or *in vitro* [29]. Peroxide lead to significant changes in sperm composition, especially in the acrosome region, and cause a sharp decline in the rate and vitality of the sperm, resulting in the prevention of the reaction of the acrosome of the sperm with the egg membrane [30, 31, 32] reinforcing the role of Ct in maintaining the movement of the sperm [25].

That the degradation of the studied traits of the SP effect may be due to the penetration of the energy substrates important for the movement of the sperm, as well as the free radicals by the oxidative processes that affect the plasma membrane of the sperm, which contains a part of the phospholipid [3, 6, 7, 8], noting differences between each and every period in the specific characteristics of semen.

Interaction was not significantly because the effect of Br is greater than the Br Conc the effect of Conc this is illustrated by the interaction of BrSP, in which the effect of SP is equal for Br_1 and Br_2 .

The interaction between Conc. SP, which has a significant effect on semen quality and hatchability, shows an inverse relationship between the studied SP properties of the peroxides that attack the plasma membrane of the sperm, accompanied by a positive relationship between the studied traits and Conc on

reducing the negative effects of SP on the effect and resistance of peroxides caused by oxidation processes as well as protection of the plasma membrane of the sperm for GAA anti-oxidation and its importance in the production of Ct, one of the most important sources of energy for the movement of sperm [11, 27, 28].

When you notice BrConcSP interactions, the role of Conc in reducing the effect of SP for each of $Br_{1,2}$, in semen characteristics, Conc's role is confirmed when the SP is rendered as observed in the MM for example, the Conc effect is equal. When SP₀, 24 is only Conc effects, increases at SP_{48, 72}, especially at Conc₅. The Conc₅ effect is becoming clearer. When SP₇₂ is in IM that was close to its effects at SP₄₈, the same is true in DS. As shows the effect of Conc when SP₄₈. This reinforces the hypothesis of the role of GAA in improving the storage conditions of the semen of the Iraqi roosters, and the antioxidant antagonist [28].

The improvement in Ha was due to the characteristics (tables 1, 2, 3, 4 and 5), with the effect of GAA, which provided protection against oxidizing factors, or worked on processing the tail of sperm with energy needed for the movement and fertilization [2, 17]. That decrease in the movement of the fetus reduces the sperm stored in the uterovaginal sperm - host glands, because the movement is necessary to cross the vagina and access to the glands, and that the reduction of sperm in those glands reflected negativity in the fertility of birds [33], as well as the importance of movement of the embryos to move from the uterovaginal sperm - host glands to the funnel during ovulation to fertilize the ovarian egg from the ovary [3]. In addition, protection against oxidative damage to the ovaries increases in IM, which is expressed in the rapid transfer of the embryos to the repression. The movement of the sperm is a major condition for the fertilization process, and the reduction of SD, AS and AA increases the fertility rates of fertilization [10], and the fertility ratio is positively correlated with Ha [2, 17].

CONCLUSION

According to previous results, it can be concluded that adding GAA to semen diluents improves the quality characteristics of semen and hatching rate. It limits the negative effects of the in vitro storage. It can therefore be used in the Iraqi rooster's semen diluents to improve its productive qualities and looking for better ways to prolong semen storage.

ACKNOWLEDGEMENTS

Thanks to Dr. Firas Muzahem Hussein Al – Khilani, Director General of Agricultural Research office / Ministry of Agriculture / Iraq, to provide research requirements. And Mr. Ali Hussein Khalil Al – Hilali, office of Animal resources / Ministry of Agriculture / Iraq, to review and revise the research.

REFERENCES

- [1] Al Athari, A. K, A. A. Al Rawi, F. M. Al Khilani, and Z. H. Al – Bustani. 2002. Performance of indigenous genetic lines of Iraqi chicken. IPA. J. Agri. Res. 12 (4), pp. 53 – 67.
- [2] Al Daraji, H. J. 1998. Effect of ascorbic acid supplementation on physiological and productive traits of Fawbro broiler breeders' flocks reared under hot climate. Ph. D. Dissertation, College of Agriculture, University of Baghdad.
- [3] Al Daraji, H. J. 2013. Artificial Insemination in Birds. 1st ed. Ministry of Higher Education and Scientific Research, College of Agriculture, University of Baghdad.
- [4] Bakst, M. 1993. Changes in sperm surfaces associated with epididymal transit. Oviducal sperm storage in poultry. Reprod. Fert. Dev. 5, pp. 595 – 599. <u>Article</u>.
- [5] Blesbois, E., I. Grasseau and D. Hermier. 1999. Changes in lipid content of fowl spermatozoa after liquid storage at 2 to 5 C. Theriogenology. 52, pp. 325 – 334.
- [6] Al Daraji, H. J. 2011. Effect of diluent supplementation with different levels of green tea on roosters' semen quality during *in vitro* storage. Int. J. of Plan., Anim. & Envi. Sci. 1 (3), pp. 51 – 56. <u>Article</u>.
- [7] Al Daraji, H. J. 2014. Impact of extender supplementation with tomato juice on semen quality of chicken semen during liquid storage. Int. J. of Bio. Sci. & Appl. 1 (1), pp. 19 – 23. <u>Article</u>.
- [8] Al Daraji, H. J. 2012. The use of pomegranate juice for counteract lipid peroxidation that naturally occurred during liquid storage of roosters' semen. Am. J. Pharm. Tech. Res. 2 (4), pp. 341 – 350. <u>Article.</u>
- [9] Lemme, A, J. Ringel, A. Sterk and J. F. Young. 2007. Supplemental guanidino acetic acid affects energy metabolism of broilers. 16th European Symposium on Poultry Nutrition, pp. 339 – 342. <u>Article</u>.
- [10] Dilger, R. N., K. Bryant-Angeloni, R. L. Payne, A. Lemme and C. M. Parsons. 2013. Dietary guanidino acetic acid is an efficacious replacement for arginine for young chicks. Poultry Sci. 92 (1), pp. 171 – 177. <u>Article</u>.
- [11] Walker, J. B. 1979. Creatine: biosynthesis, regulation, and function. Adv. Enzymol. Relat. Areas Mol. Biol. 50, pp. 177 – 242. <u>Article</u>.

- [12] Schmidt, A., B. Marescau, E. A. Boehm, W. K. J. Renema, R. Peco and A. Das. 2004. Severely altered guanidino compound levels, disturbed body weight homeostasis and impaired fertility in a mouse model of guanidinoacetate N – methyltransferase (GAMT) deficiency. Hum. Mol. Genet. 13, pp. 905 – 921. <u>Article</u>.
- [13] Vigue, C., L. Vigue and G. Huszar. 1992. Adenosine triphosphate (ATP) concentrations and ATP/adenosine diphosphate ratios in human sperm of normospermic, oligospermic, and asthenospermic specimens and in their swim-up fractions: lack of correlation between ATP parameters and sperm creatine kinase concentrations. J. Androl. 13, pp. 305 – 311. Article.
- [14] Etches, R. J. 2000. Reproduction in Poultry. University Press, Cambridge, UK. pp. 208 – 234.
- [15] Tapeh, R. S., M. Zhandi, M., Zaghari and A. Akhlaghi. 2017. Effects of guanidinoacetic acid diet supplementation on semen quality and fertility of broiler breeder roosters. Theriogenology, 89, pp. 178 – 182. <u>Article</u>.
- [16] . Lake, P. E. and J. M. Stewart. 1978.Artificial Insemination in Poultry. Bulletin 213, Ministry of Agriculture, Fisheries and Food, London. P: 10.
- [17] Al Hayani, W. K. A. 2012. Effect of dietary supplementation with different levels of L – carnitine on productive, physiological and reproductive performance of guinea fowl. Ph. D. Dissertation. College of Agriculture, University of Baghdad.
- [18] Lake, P. E. 1960. Studies on the dilution and storage of fowl semen. J. Reprod. Fert. 1, pp. 30 – 35. <u>Article</u>.
- [19] Sexton, T. J. 1976. Studies on the dilution of turkey semen. Br. Poultry Sci. 17, pp. 179 – 186. <u>Article</u>.
- [20] Al Daraji, H. J. 2001. Effects of holding temperature and time on acrosomal abnormalities of fowl sperms. Indian J. Anim. Sci. 71 (1), pp. 32 – 34. <u>Article</u>.
- [21] Al Daraji, H. J, B. T. O. Al Tikriti, K. H. Hassan and A. A. Al – Rawi. 2002. New techniques for determination of avian spermatozoa abnormalities. Res. J. Bio. Tech. 4 (1), pp. 47 – 64. <u>Article</u>.
- [22] SPSS. 2010. User guide statistic version, 18th ed. SPSS, statistical package for social science, user guide statistical version, 6th ed.
- [23] Duncan, D. B. 1955. Multiple range and Multiple F test. Biometrics. 11: 1 – 42.

- [24] Lee, H. J, W. S. Fillers and M. R. Iyengar 1988. Phosphocreatine, an intracellular high – energy compound, is found in the extracellular fluid of the seminal vesicles in mice and rats. Proc. Natl. Acad. Sci. 85, pp. 7265 – 7269. <u>Article</u>.
- [25] Sidhu, R. S., R. K. Sharma and A. Agarwal. 1998. Relationship between creatinekinase activity and semen characteristics in subfertile men. Int. J. of fert. 43 (4), pp. 192 – 197. <u>Article</u>.
- [26] Wang, X. F., X. D. Zhu, Y. J. Li, Y. Liu, J. L. Li, F. Gao, G. H. Zhou and L. Zhang. 2015. Effect of dietary creatine monohydrate supplementation on muscle lipid peroxidation and antioxidant capacity of transported broilers in summer. Poultry Sci. 94 (11), pp. 2797 – 2804. <u>Article</u>.
- [27] Ostojic, S. M., M. D. Stojanovic and G. Olcina. 2015. Oxidant-Antioxidant Capacity of Dietary Guanidinoacetic Acid. Ann. Nutr. Metab. 67, pp. 243 – 246. <u>Article.</u>
- [28] Wang, L. S., B. M. Shi, A. Shan and. Zhang Y. Y. 2012. Effects of Guanidinoacetic Acid on Growth Performance, Meat Quality and Antioxidation in Growing – Finishing Pigs. J. of Anim. & Vete. Adva. 11 (5), pp. 631 – 636. <u>Article</u>.

- [29] Agarwal, A., A. S. A. Prabakaran and T. M. Said. 2005. Prevention of Oxidative Stress Injury to Sperm. Minireview. J. Andrology, 26 (6), pp. 654 – 660. <u>Article</u>.
- [30] Aitken, R. J. and J. S. Clarkson. 1987. Cellular basis of defective sperm function and its association with the genesis of reactive oxygen species by human spermatozoa. J. Reprod. Fertil. 81, pp. 459 – 469. <u>Article</u>.
- [31] Aitken, R. J., D. Harkiss and D. W. Buckingham. 1993. Analysis of lipid peroxidation mechanisms in human spermatozoa. Mol. Reprod. Dev. 35, pp. 302 – 315. <u>Article</u>.
- [32] Samuel, F. O and M. O. Abiola. 2013. Free Radical Scavenging Activity of Methanolic Extract of Calotropis. procera Roots in Ethanol-Induced Oxidative Stress Male Wistar Rats. Int. J. of Pharmaceutical and Phytopharmacological Res. 2 (5), pp. 312 – 318. <u>Article.</u>
- [33] Murray, A. M. B., R. Denis and B. K. Speake. 1999. Acyltransferase activities in the yolk sac membrane of the chick embryo. Lipids, 34, pp. 929 – 935. <u>Article</u>.

17

Table 1. Effect of Adding Guanidinoacetic Acid to the Semen Diluents, and Duration of *in vitro* Storage on mass motility (%) of two Iraqi chickens' breeds.

Dread	Concentratio		Storage I	Periods (h)		Breed *	Maan, Draad
Breed	n (%)	0	24	48	72	Concentration	Mean: Breed
	0	75.06± 1.59 ^A	57.81±4.96 ^B	33.95±5.11 DE	5.15±0.87 ^G	42.99±8.08	
White	1	$75.64 \pm 1.45^{\text{A}}$	58.80 ± 5.53^{B}	35.47 ± 3.21 DE	6.45±0.67 ^G	49.94±6.77	48.24 ±3.42
white	3	75.87 ± 1.81 ^A	63.48 ± 1.72^{B}	43.60±2.20 CD	16.81±1.39 F	49.77±6.40	40.24 ±3.42
	5	76.58 ± 3.81^{A}	61.83±1.97 ^B	41.40±4.07 CDE	17.48±1.22 ^F	50.28±6.64	
	0	76.80 ± 0.93^{A}	59.14±4.37 ^B	43.03±2.78 ^{CD}	20.10±1.32 F	44.09±7.96	
Naked	1	77.81 ± 0.85 ^A	62.97 ± 1.53^{B}	41.20±1.83 CDE	20.97±2.10 ^F	49.32±6.82	49.00 ±3.39
brown	3	74.56 ± 3.47 ^A	64.78 ± 2.67 ^B	43.89±1.96 ^{CD}	17.89±1.70 F	50.74±6.54	49.00 ±3.39
	5	76.45 ± 1.02^{A}	64.93±3.23 ^B	44.88±1.49 ^C	21.07±0.27 ^F	51.83±6.40	
W	hite	75.57 ± 0.96^{A}	61.30±1.79 ^B	41.12±1.87 ^C	14.99±1.84 D		
Nakeo	d brown	76.62 ± 0.94^{A}	62.14±1.61 ^B	40.74±1.58 ^C	16.49±1.88 D		
Concent	ration (%)					Mean: Concer	ntration
concent						(%)	
	0.00	$75.35 \pm 0.97^{\text{A}}$	58.31±3.33 ^C	34.71±2.72 ^E	$5.80 \pm 0.57 \text{ G}$	43.54±5.55 ^B	
	.00	76.22±1.89 ^A	62.66±1.23 ^{BC}	42.50±2.13 ^D	$17.14 \pm 0.84 \text{ F}$	49.63±4.70 A	
3	8.00	$77.30 \pm 0.61^{\text{A}}$	61.06±2.24 ^{BC}	42.12±1.54 ^D	20.53 ± 1.13 ^F	50.25±4.48 A	
5	5.00	$75.50 \pm 1.67^{\text{A}}$	64.86±1.87 ^B	44.38±1.12 ^D	19.48 ± 1.05 F	51.06±4.51 ^A	
Mean Stora	ge Periods (h)	76.10 ± 0.67 ^A	61.72 ± 1.18^{B}	40.93±1.20 ^C	15.74 ± 1.30 ^D		
S.	0. V	P – Value	R-Square (%)	CV (%)	Over All Mean	Over All Sta	nder Error
H	Breed	N.S					
Conc	entration	0.01					
Stora	ge Periods	0.05					
Breed* C	Concentration	N.S					
Breed* St	orage Periods	0.05	97.3	9.49	48.62	2.4	40
	ation* Storage eriods	0.05					
	oncentration* ge Periods	0.05					

Table 2. Effect of Adding Guanidinoacetic Acid to the Semen Diluents, and Duration of *in vitro* Storage on individual motility (%) of two Iraqi chickens' breeds

Breed	Concentration		Storage Pe	eriods (h)		Breed *	Mean: Breed
Diccu	(%)	0	24	48	72	Concentration	
	0	81.73 ± 1.76 AB	66.09±1.68 ^D	21.26±2.77 ^F	6.01±1.20 G	43.77 ± 9.42	
White	1	83.74±3.40 ^A	66.98±3.83 ^D	31.99±2.33 ^E	30.83±2.11 ^{EF}	53.38 ± 6.98	49.84±3.74
white	3	77.33±4.40 ABC	65.25±3.36 ^D	27.77 ± 2.58^{EF}	29.19±0.21 EF	49.88±6.71	49.04±3.74
	5	$81.70 \pm 1.27 ^{AB}$	69.23±2.39 ^{CD}	29.52 ± 1.76^{EF}	28.91±1.99 EF	52.34 ± 7.14	
	0	83.50±3.17 ^A	68.61±1.44 ^{CD}	21.16±2.51 ^F	5.23±1.45 G	44.63 ± 9.80	
Naked	1	79.88 ± 4.04 AB	67.16±6.26 ^D	$30.40 \pm 4.24^{\text{EF}}$	32.35±0.85 E	52.45 ± 6.75	50.71±3.84
brown	3	81.12 ± 4.06 AB	67.70±2.61 ^D	29.53±3.89 ^{EF}	28.75±3.38 EF	51.78 ± 7.13	50./1±5.64
	5	82.08 ± 4.26 AB	73.25±0.84 ^{BCD}	33.86±3.69 ^E	26.77±1.86 EF	53.99 ± 7.35	
	White	81.13±1.46 ^A	66.89±1.33 ^B	27.63±1.57 ^C	23.73±3.17 ^C		
Nak	ed brown	81.65±1.71 ^A	69.18±1.66 ^B	28.74±2.09 ^C	23.28±3.32 ^C		
Conce	ntration (%)					Mean: Concen	tration (%)
	0.00	82.62±1.67 A	67.35±1.14 ^B	21.21±1.67 ^D	5.62±0.86 E	44.20 ± 6.65	В
	1.00	81.81±2.51 A	67.07±3.28 ^B	31.19±2.19 ^c	31.59±1.07 ^C	52.92 ± 4.75	А
	3.00	79.23±2.81 ^A	66.47±1.98 ^B	28.65±2.12 ^C	28.97±1.52 ^C	50.83 ± 4.79	А
	5.00	81.89±1.99 ^A	71.24±1.45 ^B	31.69±2.07 ^C	27.84±1.31 ^C	53.17 ± 5.02	А
Mean St	orage Periods (h)	81.39±1.10 ^A	68.03±1.07 ^B	28.18±1.29 ^c	23.50±2.24 D		



S. O. V	P – Value	R-Square (%)	CV (%)	Over All Mean	Over All Stander Error
Breed	N. S				
Concentration	0.01				
Storage Periods	0.01				
Breed* Concentration	N. S				
Breed* Storage Periods	0.05	97.4	10.26	50.28	2.66
Concentration* Storage Periods	0.01				
Breed* Concentration* Storage Periods	0.05				

Table 3. Effect of Adding Guanidinoacetic Acid to the Semen Diluents, and Duration of *in vitro* Storage on Dead sperm (%) of two Iraqi chickens' breeds.

Breed Co	oncentration		Storage Pe	riods (h)		Breed *	Mean:
ыееи	(%)	0	24	48	72	Concentration	nBreed
	0	14.88±1.43 ^H	24.81±3.98 ^G	69.46±3.56 ^{CDE}	93.87±1.43 ^A	50.76 ± 9.81	
White	1	15.92±1.18 ^H	22.02±2.81 ^{GH}	66.62±2.79 ^{DE}	77.19±2.11 ^{BC}	45.44 ± 8.15	44.45±4.12
white	3	14.83 ± 1.77 ^H	20.61 ± 1.98 ^{GH}	68.05 ± 2.11^{DE}	74.65±3.05 ^{BCD}	44.54 ± 8.20	44.45±4.12
	5	13.87 ± 2.28 ^H	14.03 ± 1.96^{H}	53.73 ± 2.72^{F}	66.71±3.53 ^{DE}	37.08 ± 7.20	
	0	14.30±0.90 ^H	25.74±2.19 ^G	71.60±1.03 ^{CDE}	97.77±1.23 ^A	52.35 ± 10.23	
Naked	1	15.50±1.76 ^H	19.41±3.25 ^{GH}	65.08 ± 4.15^{E}	70.33±4.14 ^{CDE}	42.58 ± 7.75	45 60+4 22
brown	3	13.73±1.74 ^H	18.68 ± 4.00 GH	56.81 ± 2.32 F	71.19±4.53 CDE	40.10 ± 7.52	45.60±4.22
	5	14.46±1.52 ^H	24.26±3.16 ^G	69.82±0.02 ^{CDE}	80.95 ± 0.17^{B}	47.37 ± 8.63	
V	White	14.88±0.76 ^D	20.36±1.69 ^{CD}	64.46 ± 2.25^{B}	78.11±3.19 ^A		
Nake	ed brown	14.50±0.68 ^D	22.02±1.65 ^c	65.83±2.01 ^B	80.06±3.59 ^A		
Concen	tration (%)					Mean: Concer	ntration (%)
	0.00	14.59±0.77 ^F	25.27 ± 2.04^{E}	70.53±1.72 ^{BC}	95.82±1.21 ^A	51.55 ± 6.93	А
	1.00	15.71±0.95 ^F	20.71 ± 2.01^{EF}	65.85±2.26 ^{CD}	73.76 ± 2.58^{B}	44.01 ± 5.51	В
	3.00	14.28±1.14 ^F	19.64 ± 2.04^{EF}	62.43±2.88 ^D	72.92 ± 2.56^{B}	42.32 ± 5.46	В
	5.00	14.16±1.23 ^F	19.14±2.83 ^{EF}	61.78±3.80 ^D	73.83 ± 3.55^{B}	42.23 ± 5.60	В
	n Storage iods (h)	14.69±0.50 ^D	21.19±1.17 ^C	65.15 ± 1.48^{B}	79.08±2.36 ^A		
	5. 0. V	P – Value	R-Square (%)	CV (%)	Over All Mean	Over All Sta	ander Error
-	Breed	N. S					
	entration	0.01					
	ge Periods	0.01					
В	reed* entration	N. S					
	l* Storage eriods	0.05	98.3	10.02	45.03	2.9	93
Stora	entration* ge Periods breed*	0.05					
Conce	entration* ge Periods	0.05					

Breed	Concentration		Storage Per	riods (h)		Breed *	Mean:
Dieeu	(%)	0	24	48	72	Concentration	Breed
	0	16.69±2.00 ^{DEFG}	24.63±2.14 ^{DE}	60.11±1.49 ^c	95.22±1.74 ^A	49.16±9.45	
White	1	$16.93 \pm 2.04^{\text{DEFG}}$	23.70±2.90 ^{DEF}	53.22±2.86 ^C	85.41 ± 2.08^{B}	44.81±8.25	45.98±4.23
white	3	16.19±2.03 ^{FG}	23.52±3.23 ^{DEF}	56.45±1.70 ^C	87.71±1.33 ^B	45.97±8.64	45.90±4.25
	5	15.15 ± 2.04^{G}	22.41±3.95 ^{DEFG}	52.23±3.85 ^C	86.11 ± 1.80^{B}	43.97±8.55	
	0	$16.67 \pm 2.51^{\text{DEFG}}$	24.86±1.71 ^D	58.92±1.71 ^C	95.39±1.31 ^A	48.96±9.42	
Naked	1	17.00±3.03 ^{DEFG}	$22.58 \pm 2.53^{\text{DEFG}}$	52.96±3.39 ^c	86.03 ± 1.64^{B}	44.64±8.38 44.90+8.24	45 45 + 4 20
brown	3	16.44 ± 2.08^{EFG}	24.44±3.38 ^{DE}	53.63±3.74 ^C	85.09 ± 2.14^{B}	44.90±8.24	45.45±4.20
	5	15.99±2.47 ^{FG}	19.52±0.25 ^{DEFG}	52.17±2.60 ^C	85.54 ± 1.93^{B}	43.30±8.54	
	White	16.24±0.89 ^D	23.57±1.35 ^C	55.50 ± 1.46^{B}	88.61±1.40 ^A		
Nak	ked brown	16.52±1.09 ^D	22.85±1.16 ^C	54.42 ± 1.50^{B}	88.01±1.49 ^A		
Conce	ntration (%)					Mean: Concentr	ation (%)
	0.00	16.68±1.43 ^{FG}	24.74±1.23 ^E	59.52±1.05 ^C	95.31±0.97 ^A	49.06±6.52	А
	1.00	16.97±1.63 ^{FG}	23.14 ± 1.74^{E}	53.09 ± 1.98^{D}	85.72 ± 1.19^{B}	44.73±5.75	В
	3.00	16.31±1.30 ^{FG}	23.98±2.10 ^E	55.04 ± 1.94^{D}	86.40 ± 1.27^{B}	45.43±5.84	
	5.00	15.57 ± 1.44^{G}	20.96±1.88 ^{EF}	52.20 ± 2.08^{D}	85.82 ± 1.19^{B}	43.64±5.91	В
Mean St	torage Periods (h)	16.38±0.69 ^D	23.21±0.88 ^C	54.96±1.03 ^B	88.31±1.00 ^A		
	S. O. V	P – Value	R-Square (%)	CV (%)	Over All Mean	Over All Stan	der Error
	Breed	N. S					
Con	centration	0.01					
Stora	age Periods	0.01					
Breed*	Concentration	N. S					
Breed* Storage Periods		0.05	98.6	9.24	45.72	2.96	,
	centration* age Periods	0.05					
Cond	Breed* centration* age Periods	0.05					

Table 4. Effect of Adding Guanidinoacetic Acid to the Semen Diluents, and Duration of *in vitro* Storage on abnormal spermatozoa (%) of two Iraqi chickens' breeds

Table 5. Effect of Adding Guanidinoacetic Acid to the Semen Diluents, and Duration of *in vitro* Storage on Acrosomal abnormalities (%) of two Iraqi chickens' breeds

Breed	Concentration		Storage F	Breed *	Mean: Breed		
bleeu	(%)	0	24	48	72	Concentration	Mean: Dreeu
	0	7.39±0.42 ^I	24.40 ± 2.06 GH	56.11±6.46 ^E	99.14±0.39 ^A	46.76 ± 10.63	
White	1	7.29±0.89 ^I	19.23 ± 0.80 ^H	49.26 ± 0.84^{F}	92.53 ± 2.36^{B}	42.08 ± 9.94	42.90±4.80
white	3	7.95±0.55 ^I	21.21±1.21 ^{GH}	49.13 ± 1.42 F	91.42 ± 3.16^{B}	42.43 ± 9.67	42.9014.00
	5	7.22±0.59 ^I	$18.40 \pm 0.30^{\mathrm{H}}$	$49.03 \pm 1.07 {}^{F}$	86.69±3.33 ^{BCD}	40.34 ± 9.33	
	0	7.24±0.57 ^I	26.36±2.14 ^G	56.22 ± 4.66^{E}	99.19±0.54 ^A	47.25 ± 10.52	
Naked brown	. 1	7.60 ± 0.81 ^I	19.86±1.09 ^{GH}	49.61±1.33 ^F	83.69±2.42 ^{CD}	40.19 ± 8.89	41.95±4.59
Nakeu DIOWI	3	7.15±0.61 ^I	18.70 ± 0.34^{H}	$49.42 \pm 0.91^{\mathrm{F}}$	88.87 ± 4.67 ^{BC}	41.03 ± 9.60	41.9514.59
	5	7.61±1.18 ^I	19.71±1.19 ^{GH}	48.06 ± 0.01^{F}	81.88±2.23 ^D	39.31 ± 8.65	
W	hite	7.46±0.29 ^D	20.81±0.88 ^C	50.88 ± 1.70^{B}	92.45±1.74 ^A		
Naked	l brown	7.40±0.36 ^D	21.16±1.08 ^c	50.83 ± 1.42^{B}	88.41±2.37 ^A		
Concenti	ration (%)					Mean: Concentr	ation (%)

20

0.00	7.31±0.32 ^H	25.38±1.40 ^F	56.17±3.56 ^D	99.16±0.30 ^A	47.01 ± 7.31 A
1.00	7.45 ± 0.54 ^H	19.55±0.62 ^G	49.44 ± 0.71^{E}	88.11±2.49 ^{BC}	41.14 ± 6.52 ^B
3.00	7.55 ± 0.41 ^H	19.95±0.80 ^G	49.28 ± 0.76^{E}	90.14 ± 2.59^{B}	41.73 ± 6.66 ^B
5.00	7.42±0.59 ^H	19.06±0.62 ^G	48.54 ± 0.53^{E}	84.29±2.09 ^C	39.83 ± 6.22 ^B
Mean Storage Periods (h)	7.43±0.22 ^D	20.99±0.68 ^c	50.86±1.08 ^B	90.43±1.50 ^A	
S. O. V	P – Value	R-Square (%)	CV (%)	Over All Mean	Over All Stander Error
Breed	N. S				
Concentration	0.05				
Storage Periods	0.01				
Breed* Concentration	N.S				
Breed* Storage Periods	0.05	99.11	8.77	42.43	3.30
Concentration* Storage Periods	0.05				
Breed* Concentration* Storage Periods	0.05				

Table 6. Effect of Adding Guanidinoacetic Acid to the Semen Diluents, and Duration of *in vitro* Storage on hatchability (%) of two Iraqi chickens' breeds

Breed	Concentration	-	Storage Per	riods (h)		Breed *	Mean: Breed
breeu	(%)	0	24	48	72	Concentration	Mean: Dreeu
	0	88.00±3.46 ^{ABC}	81.33±1.76 ABCD	43.33±4.06 GH	7.33±2.67 ^I	55.00 ± 9.85	
Mileito	1	92.67 ± 1.76^{AB}	88.67±5.46 ABC	60.00±1.15 ^F	33.33±1.33 ^H	68.67 ± 7.34	((17)200
White	3	$90.67 \pm 4.37 \text{AB}$	87.33±4.37 ABC	64.67±6.77 ^{EF}	34.67 ± 5.81 ^H	69.33 ± 7.13	66.17±3.88
	5	90.00 ± 3.06^{AB}	84.67±5.46 ABCD	73.33±4.81 ^{DE}	38.67 ± 2.40 GH	71.67 ± 6.28	
	0	84.00±2.31 ^{ABCD}	79.33±4.67 BCD	48.00±5.03 G	10.00±5.03 ^I	55.33 ± 9.12	
Naked	1	93.33±1.76 ^A	86.67±3.53 ABC	66.67±4.37 ^{EF}	36.67 ± 2.91 GH	70.83 ± 6.79	6750+261
brown	3	92.00 ± 4.16^{AB}	84.67±3.71 ABCD	66.00±3.46 ^{EF}	40.67±.67 GH	70.83 ± 6.14	67.58±3.61
	5	90.67±5.33 ^{AB}	85.33±2.91 ABCD	76.67±3.33 CDE	40.67 ± 4.37 GH	73.33 ± 6.14	
I	White	90.33±1.49 ^A	85.50±2.11 ^A	60.33±3.84 ^B	28.50±4.02 ^C		
Nake	ed brown	90.00±1.91 ^A	84.00±1.81 A	64.33±3.57 ^B	32.00±4.16 ^C		
Concer	ntration (%)					Mean: Concent	ration (%)
	0.00	86.00 ± 2.07 AB	80.33±2.28 ^{BC}	45.67±3.07 ^E	8.67±2.62 ^G	55.17 ± 6.57	В
	1.00	93.00±1.13 ^A	87.67±2.94 AB	63.33±2.51 ^D	35.00±1.61 ^F	69.75 ± 4.90	A
	3.00	91.33±2.72 ^A	86.00±2.63 AB	65.33±3.41 ^D	37.67±2.94 ^F	70.08 ± 4.61	Α
	5.00	90.33±2.75 ^A	85.00±2.77 AB	75.00±2.72 ^C	$39.67 \pm 2.28 \text{ EF}$	72.50 ± 4.30	A
Mean Stor	age Periods (h)	90.17±1.19 ^A	84.75±1.37 ^B	62.33±2.60 ^C	30.25±2.85 ^D		
S	S. O. V	P – Value	R-Square (%)	CV (%)	Over All Mean	Over All Sta	ander Error
I	Breed	N.S					
Conc	centration	0.01					
Stora	ge Periods	0.01					
Breed* Concentration		N.S					
Breed* Storage Periods		0.05	95.34	10.16	66.88	2.	54
Р	ation* Storage eriods	0.01					
	oncentration* ge Periods	0.05					