Research Article

Effect of abamectin on development of chick embryo

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Abstract
Abamectin is widely used as an insecticide and pesticide on crops. Despite of its beneficial uses it also induces some adverse effects like it causes infertility in farmers who frequently encounter abamectin. It is broadly used but little is known about its effects on avian. To check its teratogenic effects on developing chick embryo abamectin is injected into fertilized chick embryos. In chick embryo, gross anomalies and morphological changes are observed. The results of this experiment indicated that low dose of abamectin has adverse effects than that of high dose, as low dose of abamectin reduces the growth rate of developing chick embryo and delayed hatching. Moreover, hatched chicks are physically weak.

Keywords: Acetone; Chick Embryo; Growth; Pesticide; Teratogen

Introduction
Abamectin (ABM) is a macrocyclic lactone product derived from the soil microorganism Streptomyces avermitilis. ABM is used as an insecticide and acaricide in many parts of the world, and acts as an agonist of γ-aminobutyric acid [1, 2]. Abamectin is a derivative of avermectin. It acts on gamma-aminobutyric acid (GABA) receptors in both vertebrates and invertebrates [3]. Abamectin is highly lipophilic due to which it adsorbs strongly to the soil surfaces and undergoes rapid photolysis. This property of abamectin limits its mobility and transport from treated agricultural fields to surface water as well as into groundwater [4]. ABM was examined against the newly hatched larvae (neonate) and 25 days old larvae of Red Palm Weeli, Rhynchophorus ferruginous (Olivier) results showed that high concentration of abamectin induced mortality in new larvae but low mortality rate in older larvae. Abamectin also affect the hatching of eggs and development of cocoon. Consequently, the life cycle of insect is disrupted and likewise the rest of the developmental stages including egg, larva and pupa [5]. Abamectin is one of the most commonly used pesticides throughout the world. Its effect on aquatic animals had been observed, abamectin induce toxicity in D. rerio. Abamectin and difenoconazole pesticides cause greater toxicity in the fish D. rerio when they act synergistically [6]. Abamectin has characteristic of crossing the blood brain barrier in fish to cause toxicity, unlike in mammals. In both vertebrates and invertebrates, abamectin act on the receptors of gamma amino butyric acid (GABA), it
also acts on glutamatergic receptors present in the chloride channels of invertebrates [3]. The exposure to abamectin causes an increase in chloride ions which will hyperpolarize the nerve and muscle cells thus ultimately interfere with neuromuscular transmission that can lead to death [7]. The static exposure of zebrafish embryos to abamectin from 5–25 hpf suggested it inhibits the neurotransmission by the reversibly activating ligand-gated but abamectin did not affect the outgrowth of neurite from spinal motor-neurons thus making the embryos vulnerable to hypoactivity [8]. Mateus investigated that the cytotoxic effects of lower concentration of abamectin on Lithobates catesbeianus tadpoles. It ascertain the hypothesis that abamectin caused cytotoxic effects on L. catesbeianus tadpoles, even though the exposure lasted short and the concentrations were low. It revealed the likelihood for variations in the nucleus of erythrocytes circulating in amphibians [9]. In mammals, abamectin administration to male rats causes decreased sperm count, motility and increased damage in seminiferous tubule. There is also an elevated level of 4-hydroxy-2-nonenal (4-HNE)-modified proteins and poly (ADP-ribose) (PAR) expression, as markers for oxidative stress and poly (ADP-ribose) polymerase (PARP) activation, observed in rat testes exposed to abamectin [10]. In birds, the cytotoxic effects caused by low abamectin concentrations on the erythrocytes of female Japanese quails were investigated [11, 12]. It revealed distinct physical abnormalities in those birds exposed to higher abamectin levels such as nuclear shapes (asymmetric constriction nuclei in erythrocytes, notched nuclei, indented and moved nucleus) [11]. Cultured primary King pigeon brain neurons were exposed to avermectin. The results showed that AVM inhibited neuronal mitochondria activity, decreased the Δψm, and increased the activities of Caspases 3 & 9. They indicated that AVM could induce the apoptosis of the cultured brain neurons through mitochondrial injury [13]. As little is known about abamectin effect on birds therefore the present study is based on abamectin effect on chick embryos. As it acts through GABA receptors, so it can be concluded that it might affect embryo development; hence teratogenicity in chick embryos is also under consideration.

Materials and methods
Test compound
Abamectin (95% pure) is supplied by National Engineering Services Pakistan Pvt. Stock solution is made with 0.01g abamectin dissolved in solvent acetone (50 ml) and distilled water (50 ml), working solutions was made from stock solution, 5 different doses of abamectin 0.5µg, 1.25µg, 2.5µg, 5µg and 10µg were given to embryos.

Experiment model
Fertilized eggs of vingo layer chicken were purchased from Conimpex Hatchery and Breeding Farm. Total eggs were 60, all the eggs were healthy and pathogens free. Eggs were weighed that ranges from 51 - 62 g. Then eggs were cleaned with 70 % ethanol for aseptic condition and labeled them. Place eggs in incubator and maintain the temperature of incubator at 37.5°C. Eggs were rotated regularly.

Dose administration
Eggs were divided into 6 groups; each group consist of 10 eggs (n=10). One is control group and remaining five are experimental groups. Candled the eggs to mark the yolk sac and a hole is made in the shell at 45° from air sac cavity. The experimental groups were injected with different concentrations of abamectin solution (Graph 1). On the third day of incubation, doses of abamectin 0.5µg, 1.25µg, 2.5µg, 5µg and 10µg were injected to all the experimental groups A, B, C, D, E respectively. Dose was given to all experimental groups by inserting syringe
through the hole after that hole was sealed with paraffin. The eggs were placed back in incubator. Eggs were rotated twice per day.

Observation of embryos
Eggs were opened on the 8th day of drug administration. Two eggs were sacrificed from each group while placing them in Petri dishes. Wash the embryos with tap water and development was observed. The eye size of embryo and total length of embryo were measured with the help of digital Vernier caliper enlisted in (Table 1). The remaining eggs were opened on 23rd day of incubation, growth rate of all chick embryos varies with abamectin dose shown in (Graph 2).

Results
According to Hamilton stages on the 8th day certain characteristics like beak formation, limbs formation, blood vessels formation, development of feather germs was observed by comparing them with control group embryos. Embryo of control group when opened at 8th day showed normal growth i.e development of limbs, beak, and feather germs, heart and blood vessels were also present as displayed in (Fig. 1 & 2) and there is no significant effect of abamectin on the weight of embryo at 8th day (Graph 3) but low dose of abamectin affected the total length (size) of chick embryos as displayed in (Graph 4). The ANOVA analysis between the dose and eye size of embryos on day 8 proved that reducing the amount of abamectin shortened the eye size (Graph 5). The eggs of group A injected with 10 mg/L dose of abamectin show similarities with normal chicks of control groups, many of them hatched on normal time and few eggs were assisted in hatching, yet their development is near to complete with proper formation of all organs and feathers also covered the body but in some chicks’ yolk is still present outside the body (Fig. 10). Embryos injected with 5 mg dose of abamectin show slow growth, poor yolk sac retraction (Fig. 11) although the chick size is normal. In group C of experimental groups that were injected with 2.5 mg/l of abamectin chicks shows delaying in hatching, failure of yolk-sac retraction and have reduce body weight, embryo length smaller eyes size (Fig. 12). Embryos injected with slightly increase in dose to 1.25 m/L showed ectopia viscera, which only small eyes and heart developed, and took long time for hatching (Fig. 13).

While in embryos given Low dose of
abamectin 0.5 mg show in many eggs retardation in development occurred in development and their growth ceased at initial stage of development only few blood vessels were formed, heart developmental arrest occurred (Fig. 14 & 15).

Graph 1. Percentage of different type of Growth in all chick embryos

Table 1. Data about Eggs opened on 8th day after drug administration

| Egg no | Group            | Dose mg/Kg | Embryo's weight | Embryo size in mm | Eye size in mm |
|--------|------------------|------------|-----------------|-------------------|---------------|
| 2      | Experimental A   | 10 mg Kg   | 53.60 g         | 34.41             | 7.47          |
| 11     | Experimental B   | 5 mg Kg    | 57.54 g         | 30.49             | 9.43          |
| 12     | Experimental B   | 5 mg Kg    | 58.68 g         | 31.94             | 8.78          |
| 21     | Experimental C   | 2.5 mg Kg  | 55.71 g         | 29.73             | 7.22          |
| 22     | Experimental C   | 2.5 mg Kg  | 54.66 g         | 27.47             | 6.4           |
| 31     | Experimental D   | 1.25 mg Kg | 62.85 g         | 22.41             | 5.42          |
| 32     | Experimental D   | 1.25 mg Kg | 60.93 g         | 24.13             | 4.86          |
| 51     | Control          | Nil        | 53.63 g         | 40.69             | 7.25          |
| 52     | Control          | Nil        | 53.81 g         | 38.27             | 5.54          |

Graph 2. Dose quantity given to specific experimental group
Figure 1. Control group embryos opened on 8th day of drug Administration show proper allantois formation

Figure 2. Control group Embryo after washing

Figure 3. In Control group embryos opened on 8th occurred with Proper Feather germ layer formation

Figure 4. Embryo of experimental group B showing slow growth

Figure 5. Growth arrest occurs in experimental group C

Figure 6. Damaged blood vessels seen in Experimental group embryo
Figure 7. Normal hatching seen in control group

Table 2. General information about all the eggs, body weight and eye size of chicks hatched normally

| Group          | Control | Experimental A | Experimental B | Experimental C | Experimental D | Experimental E |
|----------------|---------|----------------|----------------|----------------|----------------|----------------|
| No. Of hatched eggs | 5       | 3              | 1              | 2              | 1              | 0              |
| Dose mg/Kg      | 0       | 10             | 5              | 2.5            | 1.25           | 0.5            |
| Time of incubation | 22 days | 22 days         | 22 days        | 22 days        | 22 days        | 22 days        |
| Body weight range | 53-57 g | 52-54 g        | 57-59 g        | 54-56 g        | 60-63 g        | 57-59 g        |
| Body size mean value | 168 mm   | 165.2 mm       | 166 mm         | 148 mm         | 142 mm         | 140 mm         |
| Eye size mean value | 10.2 mm    | 8.4 mm         | 6.8 mm         | 5.9 mm         | 6 mm           | 5.7 mm         |

Figure 8. Physically weak chick of experimental group

Figure 9. Assisted hatching was provided to un-hatched eggs of Experimental group
Table 3. Data of eggs in which hatching was assisted

| Group    | Control | Experimental A | Experimental B | Experimental C | Experimental D | Experimental E |
|----------|---------|----------------|----------------|----------------|----------------|----------------|
| No. Of hatched eggs | 1       | 3              | 4              | 3              | 2              | 1              |
| Dose mg/Kg | 0       | 10             | 5              | 2.5            | 1.25           | 0.5            |
| Time of incubation | 23 days | 23 days        | 23 days        | 23 days        | 23 days        | 23 days        |
| Body weight | 53-57 g | 52-54 g        | 57-59 g        | 52-56 g        | 60-63 g        | 57-59 g        |
| Body size range | 166-170 mm | 163-166.4 mm | 164-168 mm | 146-150 mm | 140-144 mm | 0 |
| Eye size range | 9-11 mm | 7.4-8.8 mm | 7.0-6.3 mm | 5-5.9 mm | 5.6 mm | 0 |

Table 4. Eggs opened on 23rd day after drug administration

| Chicks | Group       | Given dose in mg/Kg | Weight in grams | Embryo's length in mm | Eye size in mm |
|--------|-------------|---------------------|-----------------|-----------------------|----------------|
| 16     | Experimental A | 10 mg/Kg            | 52.44 g         | 163.68                | 8.30           |
| 25     | Experimental A | 10 mg/Kg            | 54.85 g         | 164.98                | 8.48           |
| 27     | Experimental B | 5 mg/Kg             | 58.65 g         | 162.89                | 6.98           |
| 39     | Experimental B | 5 mg/Kg             | 57.58 g         | 146.70                | 5.94           |
| 44     | Experimental C | 2.5 mg/Kg           | 53.30 g         | 145.55                | 5.85           |
| 47     | Experimental D | 1.25 mg/Kg          | 61.66 g         | 141.50                | 4.74           |
| 48     | Experimental D | 1.25 mg/Kg          | 62.59 g         | 143.78                | 5.47           |

Graph 3. One-way ANOVA analysis between dosage and body weight of Chicks on 8th day
Graph 4. One-way ANOVA analysis between dosage and embryo size of chicks on 8th day.

Graph 5. One-way ANOVA analysis between dosage and eye size of chicks on 8th day.
Graph 6. One-way ANOVA analysis between dosage and embryo weight of chicks on 23rd day

Graph 7. One-way ANOVA analysis between dosage and embryo length on 23rd day
Graph 8. One-way ANOVA analysis between dosage and eye size of chicks on 23rd day

Figure 10. Failure of yolk sac retraction

Figure 11. Outside the body yolk is still present

Figure 12. Ruptured yolk sac in experimental group

Figure 13. Ectopia viscera seen in chicks of experimental group
Discussion

Chicks are preferred as model organisms due to their genetic similarity and analogous homology of structure with humans also they have short development period so that results can be elaborated in short time [14, 15]. In recent years use of abamectin is increased besides of its multi benefits it also has some harmful impacts [16]. In many animals, Abamectin is found to have adverse effect [17]. Previous studies have proved the toxicological impact of abamectin on other animals but nothing is known about its teratogenic effect on chicks that's why we selected chick to observe the teratogenic effect of abamectin on them. In experiment of Abol-El-Saad and colleagues, On treatment with abamectin total count of RBCs was decreased while that of WBCs was non-significantly increased as compared to control ones in male albino rats [5]. In our study little adverse effects were observed in group B injected with 5mg/L dose of abamectin like Blood vessels were damaged, heart was ruptured and further development of blood vessels were arrested. In our experiment, embryos treated with 1.25mg/L show poor limb formation, poor development of blood vessels, heart, feather germ layer and small size of embryo, with decreasing drug doses, effects were more prominent. Further Celik-Ozenci observed that AVM causes the inhibition of neuronal mitochondria activity, increase in activities of CASPASES-3 and 9 which indicates that AVM induces the apoptosis of the culture brain neurons through mitochondria injury [10]. In our study, embryos injected with increase in dose to 1.25mg/L showed ectopia viscera, with small eyes and heart developed, without limbs formation. In group C injected with 2.5mg/ Kg of abamectin chicks shows delaying in hatching, failure of yolk sac retraction and has reduced body weight and embryo length.

In other research, after abamectin exposure, increase in chloride ions hyperpolarizes the nerve and muscle cells, ultimately interfering with neuromuscular transmission, leading to death [7]. In our experiment, embryos injected with 5mg/ Kg dose of abamectin showed slow growth, poor yolk retraction. while group A injected with 10mg/ Kg dose of abamectin showed similarities with normal chicks, many of them hatched on normal time and few eggs were assisted in hatching yet their development is near to complete with proper formation so we can conclude that abamectin affects the development rate time of hatching of growing chick embryo.

Conclusion

Abamectin being used as pesticide over crops on broad level any type of mishandling can drag bad effects so it is prerequisite to
educate farmers in regard to its safe use. From our study, it can be concluded that abamectin imparts teratogenic effect in chick embryos that’s why its use should be limited. As results showed that chicks of experimental group had more cases of growth retardation, poor yolk sac retraction and defects in size of eye and limb as compared to control group. Comparatively low doses proved to be more toxic and imparts defects in development and teratogenic effects in embryo.

Authors’ contributions
Conceived and designed the experiments: S Perveen, Performed the experiments: K Zaffar & U Khalid, Analyzed the data: N Sattar, Contributed materials/ analysis/ tools: Z Mustabshira, Wrote the paper: K Sardar.

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