Abstract—India is a country with more than 60–70% of its population dependent on agriculture. Rapid industrialization and successful green, white and blue revolutions have necessitated a large variety of chemicals/pesticides usage resulting environmental pollution which has become not only a national but a global problem. Fish occupy a prominent position in the field of aquatic toxicology and have been employed in studies concerning ecological health. In the present investigation two synthetic pyrethroids were selected namely Cyfluthrin and Fenvalerate to evaluate their toxicity on Gambusia affinis. In the present study there is significant decline in DNA content in the gill, liver and kidney tissues is observed, when the fish Gambusia affinis is exposed to the cyflthrin and fenvalerate. Decreases in the DNA content due to inhibition of the enzymes in DNA synthesis. The declines in RNA level in treated fish due to obstruction in RNA synthesis. Resulting in the swelling of Nissle bodies which are rich in RNA. Pesticides may influence directly or modify DNA, other cellular process associated with the integrity of the genome.

Keywords : India, agro-based country, pesticides, fish, DNA, RNA decline

I. INTRODUCTION

India is primarily an agro-based country with more than 60–70% of its population dependent on agriculture. Rapid industrialization and successful green, white and blue revolutions have necessitated a large variety of chemicals usage resulting environmental pollution which has become not only a national but a global problem. More than 30% of its agricultural productivity is lost due to the to pests. Excess use of insecticides to get more agriculture yield and spraying of pesticides improperly in agricultural fields cause these chemicals to accumulate in the field, along with rain water these pesticides finally reach the natural water systems like ponds, streams, lakes and rivers., it will ultimately lead to alteration in the ecosystem. [1]. Pesticides are hazardous to the biodiversity including the man. The synthetic pyrethroids act as excitatory nerve poisons, thought to poison by interfering with the ionic permeability of nerve cell membranes. Through a better understanding of toxicology and toxicity testing in relation to these biochemical parameters in non-target organisms like fish we may be able to contribute to such an inquiry and hence the present work.

Various pesticides that are commonly used in pest control programmes are generally grouped under 4 heads: They are:
(1) chlorinated hydrocarbon, the 1st generation insecticides (aldrin, DDT, chlordane, dicofol (kelthane), dieldrin, endrin, endosulphon, heptachlor, kepone, lindane, methoxychlor, mirex, and toxaphene);
(2) the second generation organophosphate insecticides include, parathion, malathion, diazinon, monocrotophos etc.,
(3) the third generation insecticides are exemplified by carbamates like, carbaryl, carbofuran, propuxur etc., and
(4) the fourth generation insecticides are the pyrethroids example, Allethrin

Fish occupy a prominent position in the field of aquatic toxicology and have been employed in studies concerning ecological health. After China, India is the second largest producer of pesticides in Asia and with 12th ranks globally. India started the production of BHC in 1952 with the establishment of a plant near Calcutta. The main use of pesticides in India is for cotton crops (45%), followed by paddy and wheat. In India the usage pattern of the of pesticide is different from the world in general. Globally 44% of the pesticide is used as insecticide, where as in India 76% of the pesticide used as insecticide. [2]. The potential toxicity of the pesticide will lead to tissue damage and change in behaviour. [3,4].
The detoxification mechanism present in the body of the organism will work to get rid of foreign substances. If the concentration of toxic substance is higher it results in damage at cellular or even at molecular level, which leads to behavioral, physiological, pathological and biochemical disorders that may prove fatal conditions to aquatic life [5,6,7,8]. The protection of the genetic diversity in natural populations maintained through DNA integrity [9]. DNA and RNA damage, health assessment of organism can be done by detection of structural/functional disturbances in the DNA [10]. The main purpose of this study is to evaluate the toxicity of cyfluthrin and fenvalerate on, Gambusia affinis with reference nucleic acid content.

**II. MATERIALS AND METHODS**

**Collection, maintenance and acclimatization of experimental fish**

Adult fresh water fishes Gambusia affinis with weight (0.7-1.5 gm), length (2.3 - 3.2 cm) from both sexes were selected randomly for the experimental studies were collected from water bodies and fisheries department of Kurnool district, Andhra Pradesh, India. The collected fish were transported in polythene bags filled with half water with least disturbance. They are kept in aquarium tanks containing well aerated de-chlorinated water to acclimatize to the laboratory conditions, with physico-chemical characteristics: temperature 24±2°C, pH 7.1±0.2 at 24°C, dissolved oxygen 9.6±0.8 mg/L, carbon dioxide 6.3±0.4 mg/L, total hardness 23.4±3.4 mg as CaCO₃/L, phosphate 0.39±0.002 µg/L, salinity in different cement tanks for one week. To prevent fungal infection the fish tanks were washed with potassium permanganate solution. Dead fish were removed as soon as possible to avoid water fouling.

Fishes were fed daily on commercial fish food. The unused food is removed daily and water was renewed daily. Test fishes were screened critically for any signs of stress, disease, physical damage and mortality. Injured, severely, diseased fishes were discarded. The fishes were exposed to sublethal concentrations for treated and control period of 96 hours. A control group was maintained with identical environment. The toxicant water as well as normal water is renewed every day. The fish were sacrificed from both experimental and control groups on 96 hours.

**Selection of pesticides**

In the present investigation two synthetic pyrethroids were selected namely Cyfluthrin and Fenvalerate.

**Cyfluthrin**

As per the reports pyrethroids are extremely toxic to fish and to some beneficial aquatic arthropods like lobster and shrimp. Cyfluthrin first registered by EPA in 1987, cyfluthrin is found in both restricted use (RUP) and general use insecticides [11]. Cyfluthrin is clear liquid, dark amber in color, with an oily to pasty consistency, and has a faint aromatic solvent odor at room temperature. Cyfluthrin, cyanomethyl (4-fluoro-3-phenoxyphenyl) methyl-3-(2,2 dichloroethyl) -2-dimethylcyclopropanecarboxylate is a broad spectrum synthetic type II pyrethroid insecticide. Cyfluthrin acts as contact and stomach poison it attacks the nervous system; a nonsystemic insecticide used to control chewing and sucking insects and also in public health situations [12]. Insects when in come in contact with cyfluthrin die of starvation and desiccation, ceasing to feed. It is extensively used in agricultural crops, stored products, public health situations (i.e. cockroaches, mosquitoes, and flies), ornamentals, turf, and domestic pests. Insects like cockroaches, ants, grain beetles, silverfish, , fleas, flies, European corn borer, Colorado potato beetle, and many others are the main target of cyfluthrin use.

Cyfluthrin act on non target organisms. The toxicity of cyfluthrin observed in various fish species is 0.68mg/L inrainbow trout (Oncorhyncus mykiss), 1.5mg/L in bluegill (Lepomis macrochirus), 22mg/L in carp (Cyprinus carpio), 3.2mg/L in golden orfe (Leuciscus idus), channel catfish (Ictalurus punctatus) and sheepshead minnow (Cyprinodon variegatus), 25.82mg/L for 48h [15], and 21.07mg/L for 72 h to Nile tilapia fry (Oreochromis niloticus L. 1758) [16]. Cyfluthrin accumulation in the flesh of Tilapia nilotica, exposed to a sublethal dose of 0.001 mg/L, was 0.009 mg/L after 5 days of application [17]. also pyrethroid toxicology in mammals, birds, amphibians and both terrestrial and aquatic invertebrates reviewed. [15,18].

Cyfluthrin causes oxidative damage in aquatic organisms resulting in the damage of essential cell components like DNA, protein, lipids and lipoproteins, caused by free oxygen radicals [19]. Thus inducing loss of cell integrity and functional alteration of cell receptors and enzymes [20]. The products which has cyfluthrin are classified as acute Toxicity Category II (bearing the signal word "Warning") or Toxicity Category I (bearing the signal word "Danger") by EPA based on its potential ability to cause eye damage [21].

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Fenvalerate

Fenvalerate is a clear viscous yellow liquid with a mild odor, a synthetic pyrethroid. First developed in 1974 by Sumitomo Chemical Co. of Japan (Sumicidin™), it is most widely used in agricultural crops such as cotton, paddy, jowar, maize, soyabean, tomato, lady’s finger, cauliflower, tobacco and tea. The biology of the non target species is affected by the use of insecticide along with pests. [22,23,24,25,26,27,28]. In India, the pesticide is used primarily to control pests of cotton and vegetables [29].

The LC$_{50}$ for 48 h of fenvalerate exposure to carp, *C. carpio* (8 cm) at pH 8.88 was reported to be 0.0210 mgL$^{-1}$ [30]. Acute toxicity (LC$_{50}$) calculated over 96h for catla fingerlings (2.02 ± 0.1 g) was observed to be 6 μgL$^{-1}$ [31]. LC$_{50}$ value of fenvalerate at 96 hours to *Channa punctatus* (20 ± 3 g) is reported to be 2.13 μg L$^{-1}$ [32]. Fish have a poor ability to metabolise and excrete fenvalerate [33] and thus are susceptible to even minute concentration of the pesticide.

Fenvalerate have half life of approximately two to four weeks in mineral and organic soils [34]. Deltamethrin is more persistent with a half-life of more than two months in soils [35]. Fenvalerate is recently developed synthetic pyrethroids, which is highly toxic to fish but shows very low avian and mammalian toxicity. [36,37].

**Experimental design**

**Group 1** : Fish exposed to tap water observed for 24 to 96 Hours (Untreated control).

**Group 2** : Fish exposed to sublethal concentration of cyfluthrin of 24 to 96 hours.

**Group 3** : Fish exposed to sublethal concentration of fenvalerate of 24 to 96 hours.
Calculation of LC$_{50}$
The LC$_{50}$ values were obtained by probit analysis method based on percentage of test animals surviving at concentrations that were lethal to more than half and less than half of the test subjects [38].

Acute toxicity studies: The toxicity studies will conducted by using the technical grade formulation available formulation of Fenvalerate and Cyfluthrin. Solutions of desired concentrations were prepared in 95% acetone to get the stock solution as well as working solution as 100mg/100ml and 1mg/1ml of toxicant chemical Fenvalarate and Cyfluthrin in both technical formulation use of acetone in control as recommended by EPA will be followed. The the acclimatized fishes were kept in cement tanks and ten fishes in two groups control fishes will be kept in dechlorinated water only. Each set of experiment was replicated six times. Mortality was recorded every 24h during the observation period of 96h.

The LC$_{50}$ values were calculated for Gambusia affinis exposed to of cyfluthrin and fenvalerate at 3, 6,12,24, 48, 72 and 96 hours and they were found to be 36.12, 34.41, 33.76, 31.32, 28.97, 26.69, 24.73 $\mu$g/L – cyfluthrin and 19.67, 18.98, 17.29, 16.98, 15.93, 14.75, 13.87 $\mu$g/L - fenvalerate respectively. The toxicity curve showed the mode of action and the toxicity of cyfluthrin and fenvalerate for different concentrations at different periods. A similar result with regard to fenvalerite toxicity in Gambussia affinis, the LC$_{50}$ value of 15.0 $\mu$g/L (48 h) was reported [39].

In the present study, it was observed that the mortality rate increased with the higher concentration of cyfluthrin and fenvalerate over a short period of time and diminished with decreasing concentration during the subsequent hours of treatment.

Statistical Analysis
The data obtained from the quantitative study were expressed as the mean ± S.E. The mean values were calculated from 6 individual observations. P—values were calculated by the students ‘t’ test. The two mean values obtained were considered significant from each other with standard error.

Estimation of nucleic acids
Nucleic acids were extracted from the tissues by the method described by Schneider (1957). [40].

Principle: Nucleic acids are separated from the other tissue components by exploiting their property of preferential solubility in hot trichloroacetic acid (TCA) solution.

Procedure: Known weight of tissue was rinsed finely and homogenized in distilled water, to homogenate 5 ml of 10% ice cold TCA was added and was kept in an ice bath for 30 minutes to allow complete precipitation of proteins and nucleic acids. The mixture was centrifuged and the precipitate was later treated with absolute alcohol and centrifuged to remove lipid materials. The precipitate free of lipids was suspended in required amount of 5% TCA and placed in a water bath maintained at 90°C for 15 minutes with occasional stirring which facilitates the quantitative separation of nucleic acids from the precipitated proteins. This was centrifuged and the supernatant was used for the estimation of Ribonucleic acid (RNA) and Deoxyribonucleic acid (DNA). [41].
Estimation of DNA

**Principle:** Estimation of DNA was performed by using diphenylamine reagent (Schneider, 1957). This method is based on the property of deoxyribose moiety of DNA to form hydroxy laevulaldehyde in TCA solution. This reacts with diphenylamine to give a coloured complex. The intensity of colouration is presumed to be proportional to the pentose (deoxyribose) concentration in the DNA hydrolyzate. Total DNA content was expressed as mg/100g wet tissue.

Reagents A. Dische’s diphenylamine reagent 1 g of diphenylamine was dissolved in 100 ml of glacial acetic acid. After adding 2.75 ml of Conc. H2SO4, the solution was mixed and stored in dark. B. Standard DNA Solution Freshly prepared by dissolving 100 mg of commercially available DNA with 100 ml of distilled water.

**Procedure:** 1 ml of isolated nucleic acid TCA hydrolyzate was taken and mixed well with 0.5 ml of perchloric acid. The solution was kept for boiling after adding 2 ml of reagent A. Boiling was done in water bath for 10 minutes. The solution was then cooled under tap water and optical density was measured at 565 nm against blank. [41].

2.12.2. Estimation of RNA

**Principle:** Total RNA content was estimated by the methods of Schneider (1957) using orcinol reagent. The ribose moiety of RNA in the TCA extract is converted to furfural which condenses with orcinol to yield a blue coloured complex. The colour intensity is presumed to be proportional to RNA concentration in the extract. Total RNA content was expressed as mg/100g wet tissue.

Reagents A. Orcinol reagent 500 ml of 30 % HCl was taken and 1 g of orcinol was dissolved well in it. 4-5 ml of 10 % ferric chloride was added and mixed in it. The solution was stored in a brown bottle. B. Standard RNA Solution dissolved 100 mg of RNA in 100 ml of distilled water

**Procedure:** 2 ml of nucleic acid TCA extract was taken and added with 2 ml of orcinol reagent. The solution was mixed and kept for boiling for 20 minutes and cooled thereafter. Absorbance was read at 670 nm in UV- spectrophotometer. [41].

III. RESULTS

**Deoxyribonucleic acid (DNA)**

**Gill**
The content of DNA in gill tissues are decreased significantly in the present study when *Gambusia affinis* exposed to cyfluthrin (group-2) and fenvalerate (group-3), when compared with the control group. The percentage of decrease over the control with cyfluthrin (group-2) are - 16.73, - 22.22, - 27.08, - 32.35 for 24, 48, 72 and 96 hours respectively. Whereas in group-3 percent decreased in DNA content over the control are -20.21, 30.05, - 39.19, - 43.94 for 24, 48, 72 and 96 hours respectively. The levels of DNA content for 3 groups are recorded are statistically significant at 1% and 5% levels (Table 1: Fig.4).

**Liver**
The observed value of DNA content in the liver tissues of *Gambusia affinis* fish exposed to cyfluthrin (group-2) and fenvalerate (group-3) showed a decline, when compared with the control group. The percent decline in DNA content with cyfluthrin (group-2) are - 19.48, - 27.88, - 30.37, - 38.63 for 24, 48, 72 and 96 hours respectively. The percent decrease in DNA content with fenvalerate (group-3) are - 23.84, - 32.38, - 45.33 for 24, 48, 72 and 96 hours respectively. The levels of DNA content for 3 groups are recorded are statistically significant at 1% and 5% levels ((Table 1: Fig.4)).

**Kidney**
The DNA content of the in the tissues of kidney in *Gambusia affinis* fish exposed to cyfluthrin (group-2) and fenvalerate (group-3) decreased in comparison with control group. The percentage of alterations are -9.92, - 16.83, - 19.62, - 23.90 for 24, 48, 72 and 96 hours respectively in fish exposed to cyfluthrin (group-2).

The percentage of alterations in fish exposed to fenvalerate (group-3) are - 11.26, - 19.57, - 28.07, - 34.39 for 24, 48, 72 and 96 hours respectively. The levels of DNA content for 3 groups are recorded are statistically significant at 1% and 5% levels (Table 1: Fig.4).
| Tissues | Groups                              | Hours of exposure | 24    | 48    | 72    | 96    |
|---------|-------------------------------------|------------------|-------|-------|-------|-------|
|         |                                     |                  | 3.310 ± 0.045 | 3.311 ± 0.043 | 3.312 ± 0.042 | 3.313 ± 0.040 |
| Gill    | Group –II- Cyfluthrin % COC          |                  | 2.756* ± 0.065 | 2.575** ± 0.064 | 2.415** ± 0.065 | 2.241** ± 0.066 |
|         |                                    |                  | % - 16.73    | % - 22.22    | % - 27.08    | % - 32.35    |
|         | Group –III- Fenvalerate % COC        |                  | 2.641* ± 0.065 | 2.316** ± 0.064 | 2.014** ± 0.065 | 1.857** ± 0.066 |
|         |                                    |                  | % -20.21     | % -30.05     | % -39.19     | % -43.94     |
|         | Group-I Control                      |                  | 3.312 ± 0.043 | 3.312 ± 0.043 | 3.312 ± 0.043 | 3.313 ± 0.040 |
|         | Group- II - Cyfluthrin % COC         |                  | 4.606* ± 0.733 | 4.129** ± 0.737 | 3.984** ± 0.739 | 3.512** ± 0.740 |
|         |                                    |                  | % -19.48      | % -27.88      | % -30.37      | % -38.63      |
|         | Group- III - Fenvalerate % COC       |                  | 4.357* ± 0.733 | 3.869** ± 0.737 | 3.128** ± 0.739 | 2.567** ± 0.740 |
|         |                                    |                  | % -23.84      | % -32.38      | % -45.33      | % -55.14      |
|         | Group-I Control                      |                  | 5.721± 0.674  | 5.722± 0.672  | 5.722± 0.671  | 5.723± 0.669  |
| Liver   | Group- II - Cyfluthrin % COC        |                  | 4.606* ± 0.733 | 4.129** ± 0.737 | 3.984** ± 0.739 | 3.512** ± 0.740 |
|         |                                    |                  | % -19.48      | % -27.88      | % -30.37      | % -38.63      |
|         | Group- III - Fenvalerate % COC       |                  | 4.357* ± 0.733 | 3.869** ± 0.737 | 3.128** ± 0.739 | 2.567** ± 0.740 |
|         |                                    |                  | % -23.84      | % -32.38      | % -45.33      | % -55.14      |
|         | Group-I Control                      |                  | 4.412 ± 0.049  | 4.413 ± 0.047  | 4.413 ± 0.046  | 4.414 ± 0.048  |
| Kidney  | Group- II - Cyfluthrin % COC        |                  | 3.974* ± 0.670 | 3.670* ± 0.672 | 3.547** ± 0.674 | 3.359** ± 0.680 |
|         |                                    |                  | % -9.92       | % -16.83      | % -19.62      | % -23.90      |
|         | Group- III - Fenvalerate % COC       |                  | 3.915* ± 0.542 | 3.549* ± 0.546 | 3.174* ± 0.550 | 2.896* ± 0.553 |
|         |                                    |                  | % -11.26      | % -19.57      | % -28.07      | % -34.39      |

Values are mean ± S.E Mean of six individual observations and student t-test. significant at * P<0.05; Significant at * P<0.01 levels (+,-) denotes decreased and increased. %COC (change over control);

**Fig. 4**: Changes in the content of Deoxyribonucleic acid (DNA) (mg/g wet. wt. of tissue) in fresh water fish Gambusia affinis exposed to Cyfluthrin and Fenvalerate for 96 hours.

![Graph showing changes in DNA content in Gambusia affinis exposed to Cyfluthrin and Fenvalerate for 96 hours](image-url)
Table 2: Variations of Ribonucleic acid (RNA) (mg/g wet. wt of tissue) content in fresh water fish *Gambusia affinis* exposed to Cyfluthrin and Fenvalerate for 96 hours

| Tissues | Groups       | Hours of exposure |
|---------|--------------|-------------------|
|         |              | 24                | 48                | 72                | 96                |
| Gill    | Group-I Control | 2.43± 0.121      | 2.43± 0.122      | 2.44± 0.122      | 2.44± 0.123      |
|         | Group –II- Cyfluthrin % COC | 2.31* ± 0.145 % - 4.93 | 2.28* ± 0.146 % - 6.17 | 2.18** ± 0.145 % - 10.65 | 2.01** ± 0.145 % - 17.066 |
|         | Group –III- Fenvalerate % COC | 2.28* ± 0.453 % - 6.17 | 2.21* ± 0.456 % - 9.05 | 2.09** ± 0.458 % - 14.34 | 1.96** ± 0.457 % - 19.6 |
| Liver   | Group-I Control | 4.98 ± 0.221     | 4.99 ± 0.222     | 4.98 ± 0.224     | 4.99 ± 0.223     |
|         | Group –II- Cyfluthrin % COC | 4.81* ± 0.323 % - 3.41 | 4.67* ± 0.324 % - 6.41 | 4.49** ± 0.324 % - 9.83 | 4.38** ± 0.326 % - 12.24 |
|         | Group –III- Fenvalerate % COC | 4.75* ± 0.452 % - 4.61 | 4.53* ± 0.457 % - 9.03 | 4.37** ± 0.458 % - 12.24 | 4.21** ± 0.459 % - 15.63 |
|         | Group-I Control | 3.10 ± 0.115     | 3.11 ± 0.114     | 3.11 ± 0.115     | 3.12 ± 0.116     |

*Note: * indicates significant difference compared to control group.
Kidney Group –II- Cyfluthrin % COC 3.01*± 0.145 % - 2.90 2.93* ± 0.147 % - 5.78 2.84 **± 0.148 % - 8.68 2.74** ± 0.149 % - 12.17

Group –III- Fenvalerate % COC 2.98*± 0.261 % - 4.18 2.90* ± 0.263 % - 6.75 2.81 **± 0.267 % - 9.64 2.69** ± 0.298 % - 13.78

Values are mean ± S.E Mean of six individual observations and student t-test. significant at * P<0.05; Significant at * P<0.01 levels(+,-) denotes decreased and increased.%COC (change over control);

Ribonucleic acid (RNA)

Gill

In the present study RNA content in the tissues of gill are decreased in the present study Gambusia affinis exposed to cyfluthrin (group-2) and fenvalerate (group-3), when compared with the control group. The percentage of decrease over the control with cyfluthrin (group-2) are - 4.93, - 6.17, -10.65, - 17.066, for 24, 48, 72 and 96 hours respectively. Whereas in group-3 percent decreased in RNA content over the control are - 6.17, - 9.05, - 14.34, - 19.6 for 24, 48, 72 and 96 hours respectively. The levels of RNA content for 3 groups are recorded are statistically significant at 1% and 5% levels (Table 2: Fig.5).

Liver

The observed value of RNA content in the liver tissues of Gambusia affinis fish exposed to cyfluthrin (group-2) and fenvalerate (group-3) showed a decline, when compare with the control group. The percent decline in RNA content with cyfluthrin (group-2) are - 3.41, - 6.41, - 9.83, - 12.24 for 24, 48, 72 and 96 hours respectively. The percent decrease in RNA content with fenvalerate (group-3) are - 4.61, - 9.03, - 12.24, - 15.63 for 24, 48, 72 and 96 hours respectively. The levels of RNA content for 3 groups are recorded are statistically significant at 1% and 5% levels (Table 2: Fig.5).

Kidney

In the present investigation the RNA content of the in the tissues of kidney in Gambusia affinis fish exposed to cyfluthrin (group-2) and fenvalerate (group-3) decreased in comparison with control group. The percentage of alterations are - 2.90, - 5.78, - 8.68, - 12.17 for 24, 48, 72 and 96 hours respectively in fish exposed to cyfluthrin (group-2). The percentage of alterations in fish exposed to fenvalerate (group-3) are - 4.18, - 6.75, - 9.64, - 13.78 for 24, 48, 72 and 96 hours respectively. The levels of RNA content for 3 groups are recorded are statistically significant at 1% and 5% levels (Table 2: Fig.5).

Fig. 5: Changes in the content of Ribonucleic acid (RNA) (mg/g wet. wt of tissue) in fresh water fish Gambusia affinis exposed to Cyfluthrin and Fenvalerate for 96 hours
IV. DISCUSSION

In the present study there is significant decline in DNA content in the gill, liver and kidney tissues is observed, when the fish *Gambusia affinis* is exposed to the cyflthrin and fenvalerate. Pesticides appears as a potential inhibitor of DNA synthesis, which might result in reduction of RNA level. In fish, DNA repairing at a much lower speed than in mammals [42,43]. The toxicant decreases the DNA content due to inhibition of the enzymes in DNA synthesis. It is known that DNA functions as a primer in DNA and RNA polymerase reactions [44], and the inhibition in DNA content can result in the inhibition of both DNA and RNA synthesis. Alteration in nucleic acid content leads to variations in protein profile [45,46]. The process of transcription control the activities of all the enzymes, when the transcription process is curtailed, mRNA and protein synthesis does not occur, due to which metabolism is impaired.

In the present study there is significant decline in RNA content in the gill, liver and kidney tissues is observed, when the fish *Gambusia affinis* is exposed to the cyflthrin and fenvalerate. The production of RNA plays a vital role in protein synthesis. The inhibition of RNA synthesis at transcription level may affect the protein content [47]. Significant decrease of RNA observed in the present study. Pesticides may influence directly or modify DNA, other cellular process associated with the integrity of the genome. The pesticides interacts with the cellular DNA produces a variety of primary lesions such as single strand breaks, double strand breaks, DNA protein cross-link and damage to purine and pyrimidine bases [48].

Similar results were also reported by various investigations: (*Cyprinus carpio*); [49]. (*Cirrhinus mrigala*); [50]. (*Channa punctatus*); [51]. (*Labeo rohita*); [52]. (*Colisa lalia*); [53]. (*Cirrhinus mrigala*); [54]. (*Channa striatus*); [55]. (*Channa punctatus*); [56]. which substantiate the results of this study suggesting that cypermethrin is a potent inhibitor of nucleic acid synthesis even at sub lethal concentrations.

Pesticides are electrophilic nature, the organophosphate (OP) compounds may attack many enzymes responsible for normal metabolic pathway. Protein level is depleted in tissues of fish because RNA plays significant role in protein synthesis [57]. The decrease in RNA level will affect the protein synthesis. [58]. reported similar results. Under stressed conditions animals...
require more energy to overcome the effect of pesticide. As a result, glucogenesis increases in comparison with protein synthesis. The significant declines in RNA level in treated fish due to obstruction in RNA synthesis. Nissle bodies are rich in RNA, the swelling and chromatolysis of Nissle bodies is caused by pesticides. [59]. Decrease in protein synthesis, impairment of nucleic acid metabolism, the degradation of cells upon exposure to pesticides may be due to, resulting in the reduction in the DNA and RNA content.

V. CONCLUSION

It can be concluded that there is correlation between fish cyfluthrin and fenvalerate exposure and alterations in the nucleic acid contents in selected tissues. The DNA and RNA contents in tissues of gill, liver and kidney are gradually decreased in experimental fish in comparison with the control group. Decreases in the DNA content due to inhibition of the enzymes in DNA synthesis. The declines in RNA level in treated fish due to obstruction in RNA synthesis, resulting in the swelling and chromatolysis of Nissle bodies which are rich in RNA.

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