DNA-Based Toxicity Assay for Pesticides in the Environment

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Abstract: Increased use of domestic and agricultural pesticides has become a serious threat to the environment. Prolonged exposure to pesticides is capable of affecting the genetic integrity of humans and other animals. The aim of this study is to access the effect of dichlorvos (DDVP), a widely used pesticide in Nigeria, on the DNA of poultry birds (Gallus domesticus). This study explored different special representation of treated groups using a three-patch matrix model incorporating dichlorvos contamination (0.01, 0.02 and 0.4%). Exposure was carried out for ten weeks after which the birds were sacrificed and the liver was extracted. Thermal denaturation of the DNA from the exposed birds resulted in a significant reduction (p< 0.01) in the DNA melting temperature from 87.2°C to 81.7°C while the GC/AT ratio was also significantly reduced (p=0.01) from 0.77 in the control to 0.42 in exposed birds respectively. Electrophoresis of isolated DNA in 0.8% agarose gels gave variations in band intensity between the control DNA sample and DNA from exposed birds. These variations in band intensity were more pronounced in the RAPD-PCR products amplified with OPE-01 primer, where there is complete disappearance of DNA bands in the birds exposed to 0.04% pesticide. Thus deletion of DNA segments of birds exposed to dichlorvos can be modelled as a molecular biomarker of genotoxicity. This may also suggest that insecticides are capable of impacting genotoxic effects on non-target populations with consistent, long-term use.

Keywords: Pesticides, genotoxicity, RAPD-PCR, melting temperature, DNA strand breaks.

RUNNING TITLE: Genotoxic effect of dichlorvos pesticide on poultry birds.

INTRODUCTION

Pesticides are one of the most potentially harmful chemicals introduced into the environment. Although they have contributed considerably to human welfare, their adverse impacts on non-target organisms are significant (Hazari and Das, 1998; John, 2007). Widespread use of pesticides in agriculture and domestic pest control has contributed to the pollution of the environment (Partanen et al. 1999). Bioaccumulation of pesticides in the food chain can lead to potentially adverse effects in humans and useful animals due to their apparent toxic effect. Some pesticides are highly persistent in nature, causing contamination of soil, ground and surface water (Wolfe et al. 1973; Frank et al. 1990). DNA is one of the most critical cellular targets for hazardous chemicals and wastes (Birnboin and Jevcak, 1981). Exposure to environmental xenobiotics, including pesticides, has the ability to cause DNA damage through the formation of strand breaks and DNA-adducts. Many in vitro and in vivo studies, as well as epidemiological approaches, have demonstrated the ability of certain chemical pesticides to produce genetic effects including cancer and other chronic pathologies in humans (Bolognesi et al. 1981). A number of DNA based assays have been used for in vitro study of effects of pesticides. In this study DNA melting characteristics and random amplified polymorphic DNA (RAPD) were used to access the genotoxic effects of pesticide exposure.

MATERIALS AND METHODS

Test Sample
Dichlorvos (2,2 – dichlorovinylidimethyl phosphate) –DDVP, was purchased from an agrochemical shop in Owerri.

Formulation of Contaminated Poultry feeds
Commercially available poultry feed was contaminated by weighing out a definite amount of the feed and mixed with a graded percentage of the pesticide to give 0.01, 0.02 and 0.04 % (w/v) contamination respectively. Feed for the control contained no pesticide.
Experimental Animals
Day old black pullets were obtained from Zartec Farms, Ibadan, Nigeria. The birds were brooded under appropriate conditions until they were seven weeks old. The seven weeks old pullets with an average weight of 557.5 ± 9.5 g were divided into four groups containing 10 birds each and housed in poultry pens at the livestock unit of the Department of Animal Science and Technology, Federal University of Technology, Owerri, Nigeria. Three groups received a diet containing 0.01, 0.02 and 0.04 % dichlorvos respectively, while the control was fed on pesticide-free diet. The experiment lasted for a period of ten weeks.

Isolation of liver DNA
After ten weeks of exposure, two birds each were taken from each group and sacrificed by decapitation and the liver extirpated. Liver DNA was isolated using the method of Sambrook et al. 1989). The concentration of DNA was determined spectrophotometrically at O.D 260nm. Purity was determined based on A260/280 ratio.

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\text{Conc of DNA} = \frac{A260 \times \text{dilution factor} \times 50 \text{ug/ml}}{1}\]

\[
\text{Total yield} = \text{Conc of DNA} \times \text{total purified sample volume}
\]

Agarose gel electrophoresis of DNA
The molecular pattern of the isolated DNA samples was analysed using Agarose gel electrophoresis. Molten agarose [0.8%] gel (2.4 g of agarose in 300ml of Tris-acetate buffer) + 30 µl of ethidium bromide was poured into a horizontal electrophoresis plate and allowed to set after which the electrophoresis buffer (0.04M Tris-acetate, 0.002M EDTA, pH 8.0) was poured into the tank. Five microlitre (5µl) of gel loading buffer (2.5 mg bromophenol blue, 4 g sucrose, in 6 ml 10mM Tris-HCl, pH 8 and 1mM EDTA) was added to each 25 µl of DNA sample and carefully loaded into the wells. The gel was run at 30-40 volt overnight and the ethidium bromide stained bands were observed and photographed using a UV transilluminator.

Thermal denaturation of DNA
Thermal denaturation of DNA was carried out by dissolving 10ml of DNA solution in 990 ml of saline sodium citrate (SSC) buffer. The DNA solution was placed in a stoppered special thermal cell (A.H. Thomas Co. Philadelphia, PA, USA), and overlaid with mineral oil to prevent evaporation. The samples were incubated at 70°C, 75°C, 80°C, 90°C and 95°C for 30 minutes. The rate of temperature increase was controlled by a thermocouple type F4301 (Haake Berlin Karsruhe, Germany). The melting profile was generated by measuring absorbance at 260nm every 0.5°C. A graph of absorbance against temperature was plotted and the melting temperature (Tm) was determined from the graph. The percentage GC (%GC) was calculated using the formula: %GC = \((Tm - 70°C) \times 2.5\), where 2.5 is a constant.

RAPD-PCR Analysis
The RAPD PCR was carried out using 4 primer sets: OPA-02 (5’-TGCCGAGCTG-3’), OPB-06 (5’-TGCTCTGCC-3’), OPC-04 (5’-CCGCATCTAC-3’) and OPE-01 (5’- CCAAGGTCC-3’). The RAPD protocol used was the one described by Ferrero et al. [5]. The PCR was performed in 25 µl of a reaction mixture containing DNA (10-200 ng), 200 µM of each deoxynucleoside triphosphates (dNTP) (Promega), 2.5 mM MgCl\(_2\), 1X PCR Buffer, 20 µM primer, 2.5 units of Taq DNA polymerase (Promega) and sterile distilled water. Thermal cycling was conducted in an Eppendorf Master Cycler Gradient for an initial denaturation at 94°C for 5 minutes followed by 40 amplification cycles of 1 minute at 94°C; 1 minute at 28 °C and 1 minute at 72°C. This was followed by a final extension step for 10 minutes at 72°C. The amplification product was separated on 1% agarose gel electrophoresis and visualized by ethidium bromide staining. 1kb DNA ladder was used as DNA molecular weight standard.

Statistical Analysis
The results were analyzed statistically using a one way analysis of variance (ANOVA). The results were expressed as mean ± SEM. The means were separated using Turkey’s test and considered different at p<0.01 and p<0.05.

RESULTS
The molecular pattern of isolated liver DNA in agarose gel shows that there were distinct bands for different liver tissues were found to be relatively pure (Table 1). All DNA samples isolated from the birds fed on 0.04% pesticide contaminated diet were incubated at 70°C. The amplification product was separated on 1% agarose gel electrophoresis and visualized by ethidium bromide staining. The melting profile was generated by measuring absorbance at 260nm every 0.5°C. A graph of absorbance against temperature was plotted and the melting temperature (Tm) was determined from the graph. The percentage GC (%GC) was calculated using the formula: %GC = \((Tm - 70°C) \times 2.5\), where 2.5 is a constant.

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RESULTS
The molecular pattern of isolated liver DNA in agarose gel shows that there were distinct bands for each DNA isolated from the different birds (Plate 1). The highest yield of DNA was observed with the birds fed on 0.04% pesticide contaminated diet (Table 1). All DNA samples isolated from the different liver tissues were found to be relatively pure as shown by their A260/A280 ratios. Generally a high quality DNA sample has an A260/A280 ratio between 1.7 and 2.0. There was significant reduction (p< 0.01) in the DNA melting temperature from 87.2% in the control to 81.7% in birds exposed to the pesticide. The Guanine-cytosine/Adenine-Thymine (GC/AT) ratio was also significantly reduced (p<0.01) from 0.77 in the control to 0.42% in birds exposed to the pesticide (Table 2). Results of the RAPD-PCR analysis shows that there were variations in band intensity between the control DNA sample and those exposed to pesticide. Variations were particularly observed in the products amplified with OPA-02 (Plate 2), OPC-04 (Plate 3) and OPE-01.

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CONCLUSION

This study highlights the potentials of using DNA based analysis in evaluating the environmental impact of pesticide exposure. The study has shown that melting temperature as well as DNA profile on agarose gel can be used as biomarkers of pesticide (dichlorvos) exposure.

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CONFLICT OF INTEREST: Authors have no conflict of interest.

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Plate 1: Agarose gel electrophoresis of liver DNA Samples. Lane 1 = control, lane 2 = 0.01% DDVP, Lane 3 = 0.02% DDVP, lane 4 = 0.04% DDVP, M = marker.

Table 1: Quantitation of DNA isolated from the liver of poultry birds exposed to DDVP
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| Experimental Group | A260/280  | Yield (ng/UL) |
|--------------------|----------|--------------|
| Control            | 1.72 ± 0.01 | 2507.0 ± 394.14 |
| 0.01% DDVP         | 1.67 ± 0.05 | 2417.5 ± 138.16 |
| 0.02% DDVP         | 1.72 ± 0.01 | 3271.3 ± 255.26 |
| 0.04% DDVP         | 1.75 ± 0.04 | 3978.1 ± 149.34 |

Each sample was assayed twice for DNA quantitation at 260nm. Purity was determined based on A260/280 ratio.

Table 2: Summary of melting temperature, percentage nucleotide bases of liver DNA samples

| Experimental group | Tm (°C) | % GC  | % AT | GC/AT ratio |
|--------------------|---------|-------|------|-------------|
| Control            | 87.2a   | 43.5  | 56.5 | 0.77a       |
| 0.01% DDVP         | 82.1b   | 31.3  | 68.7 | 0.46b       |
| 0.02% DDVP         | 82.0b   | 30.0  | 70.0 | 0.43b       |
| 0.04% DDVP         | 81.1b   | 27.8  | 72.2 | 0.39b       |

Values expressed as standard error values ± of the samples (N=3).

a-b Means within a column with no common superscript differ significantly at p<0.01

Plate 2. RAPD profile showing DNA pattern of liver of poultry birds exposed to 0.01% [2], 0.02% [3], and 0.04% [4] DDVP. The control [1] contained no pesticide. The primer used for amplification is OPA 02.
Plate 3: RAPD profile showing DNA pattern of liver of poultry birds exposed to 0.01% [2], 0.02% [3], and 0.04% [4] DDVP. The control [1] contained no pesticide. The primer used for amplification is OPC 04.

Plate 4: RAPD profile showing DNA pattern of liver of poultry birds exposed to 0.01% [2], 0.02% [3], and 0.04% [4] DDVP. The control [1] contained no pesticide. The primer used for amplification is OPE 01.