Application of Neutral Red Uptake Assay Using EPC Cells as an Alternative to the Fish Acute Toxicity Test for Pesticide

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Abstract This study evaluated in vitro cytotoxicity of 5 pesticides, including 2 herbicides, 2 germicides, and an insecticide, as an alternative to the fish acute toxicity test. The in vitro cytotoxicity was tested using a neutral red uptake (NRU) assay with epithelioma papulosum cyprini (EPC) cells that originated from the epidermal tissue of Cyprinus carpio (common carp). An in vivo fish acute toxicity test was conducted according to OECD Test Guideline No. 203 using Aphyocypris chinensis (Chinese bleak), Oryzias latipes (Japanese medaka), and C. carpio. The results showed that the sensitivity of the cell viability assay for the pesticides was similar to the fish acute test in ranking order despite having approximately 10 times less absolute sensitivity. The $r^2$ correlation values were calculated as 0.38 ($p = 0.26$), 0.76 ($p = 0.05$) and 0.90 ($p = 0.01$) for A. chinensis, O. latipes, and C. carpio, respectively. These results suggested that the potential of EPC cell viability assay as an alternative to the fish acute toxicity test due to their good correlation and NRU assay is expected to serve as a useful tool for predicting acute fish lethality for pesticides if further studies with a large set of pesticides are conducted.

Key words cytotoxicity, EPC cells, fish acute test, NRU assay, pesticides

Introduction

As the use of pesticides has made a stable food supply and an alternative to labor shortages possible, they are regarded as the most economic and effective agricultural materials in modern agriculture development (Rasmussen et al., 1998). According to the Rural Development Administration (RDA) in South Korea, 24,000 tons of pesticides have been steadily consumed in South Korea since 2004. In addition, RDA also showed the number of pesticide had increased twice from 1152 in 2004 to 2265 in 2009 (RDA, 2013). In the near future, as demands for the development of more effective and safer pesticides continue, the number of pesticides in use is expected to grow.

At the same time, the fish acute toxicity test to obtain information about the toxic potential of these pesticides is globally required in the registration process of pesticides, including in Korea. However, under the new European chemicals policy, Registration, Evaluation and Authorisation of Chemicals (REACH), the fish acute toxicity test (as a vertebrate animal test with mortality) will be reduced or even replaced by alternative methods. Because conducting this test is costly and time-consuming and requires a considerable
Materials and Methods

Chemicals

The 5 pesticides (an herbicide, germicide A, germicide B, insecticide A, and insecticide B) used in this study were purchased from pesticide companies. These pesticides are chosen randomly among relatively recently developed pesticides which are expected to be used more gradually. The active ingredients for each pesticide were shown in Table 1. Neutral red was purchased from Sigma (St. Louis, MO, USA), and the cell culture media and all of the supplements were purchased from Gibco (Paisley, Scotland, UK). Trypsin powder was purchased from Amresco (Solon, Ohio, USA).

Cell culture

Epithelioma papulosum cyprini (EPC) cells from carp (C. carpio) epithelium, which is an established monolayer-type cultured fish cell line, were used. The EPC cells were distributed by Chonnam National University (Department of Aqualife Medicine). The cells were maintained at 20°C in Dulbecco’s Modified Eagle’s Medium (DMEM) with 10% fetal bovine serum (FBS) and 1% penicillin-streptomycin. When needed, the cells were trypsinized with a solution containing 8 g of NaCl, 0.2 g of KH₂PO₄, 0.2 g of KCl, 1.15 g of NaHPO₄, 0.2 g of EDTA, and 1.2 g of trypsin powder in 1 L of deionized, distilled water.

Neutral red uptake assay

We seeded 1 × 10⁴ cells in 100 uL of DMEM in 96-well tissue culture microtiter plates. After incubating for 24 hrs, the medium was removed, and the cells were treated with varied concentrations (0.1, 1, 10, 100 mg/L) of the test chemicals, including a control. After another 24 hrs of incubation, the medium was removed and replaced with 200 uL of DMEM and 3 uL of neutral red dye and incubated for 2 hrs. The cells were washed once with PBS, and then 100 uL of 1% acetic acid in 50% ethanol was added to the wells. The plate was incubated for 10 min at 20°C, and then the absorbance was read at 540 and 690 nm. The viability percentage (%) was calculated as follows:

% Viability =

\[
\frac{(\text{Absorbance}_{540\text{nm}} \text{test material}) - (\text{Absorbance}_{540\text{nm}} \text{control})}{(\text{Absorbance}_{690\text{nm}} \text{control}) - (\text{Absorbance}_{690\text{nm}} \text{test material})} \times 100
\]

Fish

The A. chinensis, O. latipes, and C. carpio used in this study were obtained from the Korea Institute of Toxicology (Daejeon, Korea). The fish were maintained in dechlorinated tap water at 22 ± 1°C with a photoperiod of 16:8 hrs (light:dark). All of the fish were fed twice a day with flake food (TetraMin, Tetra Corp., Melle, Germany) for A. chinensis and O. latipes and top meal (Tabia Corp., Korea) for C. carpio.

Fish acute toxicity test

The fish acute toxicity tests were conducted according to the OECD Test Guideline No. 203 (OECD, 1992). Briefly, fish which were corresponded with length suggested in OECD test guideline for O. latipes and C. carpio and 5-6 cm for A. chinensis were selected for the test. The tests lasted for 96 hrs in static systems without medium changes. Each...
test unit contained 7 fish, both male and female, in 5 liters of test media, and the test was performed with no replicates.

Statistics and data analysis

All of the statistical analyses were conducted using the Statistical Package for the Social Sciences (SPSS) 20 (IBM, United States), and all of the data were plotted using SigmaPlot 10.0 (Systat Software Inc., Germany). Probit analysis was used for calculation the IC$_{50}$ and LC$_{50}$ values.

Results and Discussion

The apparent difference in the loss of the cell monolayer integrity and changes in the cell shape could be observed for all of the pesticides. Figs. 1B, C, and D show the changes in cell density, breaks in the cell monolayer (arrows), and alterations in cell shape and cell detachment, respectively, while no morphological changes were observed in control cells in Fig. 1A. Cell growth, division, death and changes in cell shape are important for tissue morphogenesis during development. Cell shape is controlled by the regulation of intra-cellular mechanisms and the cell’s physical interaction with its environment. Cell shape changes generate disturbances in cell migration and contribute to heterologies of tissue morphogenesis such as ventral furrow formation, dorsal closure, and convergent extension at the development level (Paluch and Heigenberg, 2009). In the present study, the 5 pesticides all showed changes in cell shape and density similar to Fig. 1. The light microscopy appearance of germicide B is shown as a representative.

The in vitro IC$_{50}$ values for the different pesticides with the EPC cells were rank ordered from the most toxic to the least toxic as follows: Insecticide B (0.10 mg/L) > Germicide B (0.58 mg/L) > Herbicide (1.25 mg/L) > Insecticide A (7.27 mg/L) > Germicide A (42.4 mg/L). The in vivo LC$_{50}$ values for the 3 species of fish had a similar rank order as the IC$_{50}$ values in $A$. chinensis and $O$. latipes: Insecticide B (0.01 and 0.09 mg/L, respectively) > Germicide B (0.07 and 0.13 mg/L, respectively) > Insecticide A (0.16 and 1.05 mg/L, respectively) > Herbicide (3.54 and 1.27 mg/L, respectively) > Germicide A (3.81 and 2.43 mg/L, respectively). In $C$. carpio was shown as Germicide B (0.08 mg/L) > Insecticide B (0.12 mg/L) > insecticide A (0.11 mg/L) > Herbicide (0.8 mg/L) > Germicide A (3.07 mg/L). Although there were some differences in order, more than 2 steps were not exceeded. Previous study demonstrated cell line tests could be used as

Fig. 1. Representative light microscopy appearance of EPC cells following 24 hrs of exposure to pesticides. (A) Control cells maintained in DMEM displaying an intact, fully confluent monolayer. (B) Cells exposed to 0.1 mg/L germicide B showing a discernible change in the density of the monolayer. (C) Cells exposed to 1 mg/L germicide B displaying a clear loss of cell monolayer integrity. (D) Cells exposed to 10 mg/L germicide B illustrating clear changes in cell shape and density. Magnification: 100 X. Arrows indicate the loss of cell monolayer integrity.
screening tools because of their different sensitivities (Ni Shuilleabhain et al., 2004). We also show the potential of EPC cell line as a screening tool in comparison ranking.

Linear correlations between the \textit{in vitro} \(IC_{50}\) values and each \textit{in vivo} \(LC_{50}\) value were also determined. The results revealed that a linear correlation between the \textit{in vitro} \(IC_{50}\) for the EPC cells and the \textit{in vivo} \(LC_{50}\) for \textit{C. carpio} were the most highly significant \((p = 0.01\) with \(r^2 = 0.90)\) followed by \textit{O. latipes} \((p = 0.05\) with \(r^2 = 0.76)\). For \textit{A. chinensis}, no linear correlation was observed \((p = 0.26\) with \(r^2 = 0.38)\) (Fig. 2). The respective \(IC_{50}\) values for the EPC cells and the \(LC_{50}\) values for \textit{A. chinensis}, \textit{O. latipes}, and \textit{C. carpio} are shown in Table 1. The results showed that the \(IC_{50}\) values were closely correlated with the whole fish \(LC_{50}\) values and those highly significant linear correlations between the \textit{in vitro} and \textit{in vivo} values were found for \textit{C. carpio} and \textit{O. latipes}. However, \textit{A. chinensis} had no correlation for the pesticides comparing the \textit{in vitro} cytotoxicity to the \textit{in vivo} lethality and showed approximately 10 times less sensitivity.

Table 2 shows the correlations between the \(LC_{50}\) values from fish acute toxicity testing and the E/\(IC_{50}\) ratios from cell viability testing in previous studies. Such results demonstrating that the sensitivity to lethal effects in fish may appear differently depending on the type of fish support the lack of a correlation for \textit{A. chinensis} in the present study. In addition, Castano et al. (2003) and Segner (2004) have also pointed out that the sensitivity of cells toward individual chemicals appears to be less than that of whole fish, and therefore, the lower sensitivity of the cell culture assay is a major stumbling block to be overcome in the regulatory testing of chemicals. Nonetheless, because the ranking order of cell viability for the 5 pesticides was similar to whole fish and the correlations between the \textit{in vitro} and \textit{in vivo} test results for \textit{C. carpio} and \textit{O. latipes} were significant, ranking

Table 1. Comparison between \textit{in vitro} (\(IC_{50}\)) and \textit{in vivo} (\(LC_{50}\)) toxicity

<table>
<thead>
<tr>
<th>Pesticides</th>
<th>Active ingredient of pesticide</th>
<th>EPC</th>
<th>(IC_{50}) mg/L (95% C.I.)</th>
<th>(LC_{50}) mg/L (95% C.I.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Herbicide</td>
<td>Oxadiargyl EC</td>
<td>1.25 (0.57-2.14)</td>
<td>3.54 (2.11-5.92)</td>
<td>1.27 (1.06-1.61)</td>
</tr>
<tr>
<td>Germicide A</td>
<td>Tebuconazole + Prochloraz EC</td>
<td>42.4 (25.8-55.2)</td>
<td>3.81 (3.45-4.26)</td>
<td>2.43 (2.02-3.01)</td>
</tr>
<tr>
<td>Germicide B</td>
<td>Cyazofamid + Chlorothalonil SC</td>
<td>0.58 (0.09-2.57)</td>
<td>0.07 (0.07-0.08)</td>
<td>0.13 (0.10-0.16)</td>
</tr>
<tr>
<td>Insecticide A</td>
<td>Imidacloprid + Bifentihrin WP</td>
<td>7.27 (4.16-8.74)</td>
<td>0.16 (0.15-0.18)</td>
<td>1.05 (0.89-1.26)</td>
</tr>
<tr>
<td>Insecticide B</td>
<td>Fenbutatin oxide EC</td>
<td>0.10 (0.09-0.23)</td>
<td>0.01 (0.01-0.09)</td>
<td>0.09 (0.01-0.01)</td>
</tr>
</tbody>
</table>

\(r^2 = 0.38, p = 0.26\) \(r^2 = 0.76, p = 0.05\) \(r^2 = 0.90, p = 0.01\)

\(95\%\) Confidence Interval
pesticides in order of toxic status on cell viability assay could be suggested as a screening tool and alternative to the fish acute toxicity test.

In conclusion, the NRU assay using EPC cell was well correlated with acute toxicity of *C. carpio* and *O. latipes*. Therefore, it will serve a potential tool as an alternative to the fish acute toxicity test. However, further studies with a larger set of pesticides are needed to strengthen the reliability of the assays and to validate the correlation with *in vivo* data.

Acknowledgment

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Literature Cited


어류급성독성시험 대체법으로서 잉어표피세포를 이용한 neutral red uptake 분석법 적용

요약
본 연구는 5가지 제품농약을 이용하여 어류 급성독성시험 결과 (반수저농도)와 잉어의 표피에서 유래된 EPC 세포를 이용한 neutral red uptake 결과 (반수저농도)를 비교함으로써 동물 실험의 대체 가능성을 평가하기 위하여 수행되었다. 어류 급성 독성시험은 왜몰개 (Aphyocypris chinensis)를 포함하여 OECD와 농촌진흥청의 녹색에 대한 독성시험기준에서 추천하는 어종인 종사리 (Oryzias latipes)와 잉어 (Carpinus carpio)를 이용하여 수행하였다. 5가지 제품 농약에 대한 민감도는 어류에 비하여 세포에서 약 10배 더 낮게 확인되었지만, 독성을 서열화 하였을 때 나타나는 순서는 두 가지 방식에서 모두 비슷하게 나타났다. 5가지 제품 농약에 대한 세포와 어류 독성값의 상관성을 분석한 결과는 A. chinensis, O. latipes와 C. carpio에서 각각 \( r^2 = 0.38 \) (\( p = 0.26 \)), \( r^2 = 0.76 \) (\( p = 0.05 \)), \( r^2 = 0.90 \) (\( p = 0.01 \)을 나타내었다. 본 시험의 결과, EPC 세포를 이용한 NRU 시험은 O. latipes와 C. carpio에 대한 어류 독성시험 결과와 상관성이 높으므로 향후 더 많은 약제시험을 통해 어류 급성독성시험의 대체시험법으로서의 가능성이 기대된다.