Analysis and Characterization of Isopropyl Carbanilate Herbicide and Its Photoproducts

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Abstract

The phototransformation of the herbicide Isopropyl carbanilate (IPC) has been investigated under UV light. Irradiation of the herbicide at room temperature in aqueous and organic solvents such as hexane and methanol afforded new photo-products formed as a consequence of various processes including photo-Fries rearrangement, ring solvolysis, hydrolysis of the amide/carbamoyl and ester bonds, ring coupling and polymerization. The percentage remaining of the herbicide as a function of time was followed periodically starting from zero time up to three hours. Analyses were performed by GC-FID equipped with a semipolar glass column operated at 170°C. The rate of photo disappearance of IPC under controlled lab condition followed 1st order kinetics and found to be solvent dependent in the manner of non polar > polar solvents. The photo-products were successfully separated by GC and preparative TLC (Silica gel F-254) and were identified using either GC-MS and/or MS. Identifications were assigned on the bases of molecular ions, mass fragmentation pattern and whenever possible by comparison with the mass spectra of literature analogues.

Keywords: Phototransformation, Isopropyl carbanilate, Isopropyl-N-Phenylcarbamate, Propham, IPC and GC-MS

1. Introduction

Isopropyl carbanilate or Isopropyl N-phenylcarbamate, also known as propham, Turberite® or IPC, is a xenobiotic urethane derivative with Kow = 2.6 (pH 7 and 20°C), Log P = 2.58 and LD₅₀ ≥ 5000 mg/kg. Propham has the formula of C₁₀H₁₂NO₂ shown in Figure 1.

IPC has been widely used as a soil acting herbicide with selectivity towards weeds and grasses in broad-leaved crop vegetables [1,2]. Once IPC finds its way into the environment, it undergoes biological, chemical and photochemical transformation [3], producing eventually the same or similar degradation products, indicating difficulties in judging the cause for a specific transformation.

Regarding biodegradation of the herbicide, it has been reported that hydrolysis to the respective aniline and isopropanol in a reverse manner to its synthesis constitutes the major metabolic pathway especially in tolerant plants and soil [4-10]. Molecular modification into more polar derivatives through hydroxylation of the aromatic and side alky moieties with subsequent conjugation to endogenous matter constitutes the principal transformation route of IPC in animals [10-12]. Additionally, a minor route for propham metabolism in animals include hydrolysis into aniline with subsequent acylation, hydroxylation and finally conjugation to glucuronide or sulfate [13], N-hydroxylation and ring methoxylation of the intact IPC were also reported to occur in animals [13-15].

Previous studies pertaining to the photolysis of IPC revealed that it was highly resistant and photolysed very slowly even at short UV radiation [16,17]. Other studies in this context showed that IPC photolysed very slowly in a reverse manner to its synthesis giving aniline, phenylisocyanate, isopropanol and diphenylurea, all in accordance to its thermal degradation or pyrolysis products [18,19]. Investigations in relation to IPC persistence in the environment indicated a half-life of 254 days with...
unidentified metabolites in water under simulated sun-
ner sunshine [16]. However, in presence of TiO$_2$/H$_2$O$_2$
and UV, the half-life of IPC was shortened to few hours
[20,21], giving a variety of degradation products including
ortho- and para-hydroxy analogues, benzoquinone,
isopropyl aminobenzoate and aminophenol derivatives
[22-27]. In view of these conflicting reports and inade-
quate data on propham photo-product, herein we report
the IPC photodegradation in various media with its re-
spective photoproducts; however, such data are not fully
available and the demand for it is important both to
chemists and environmentalists.

2. Experimental Studies

2.1. Material and Equipments

Isopropyl- N-penylcarbamate technical grade of purity >
99% from Sigma, solvents and reagents were of analyti-
cal grade or Analar and used as such, water used was
deonised. Thin layer chromatography plates of silica gel
with fluorescent indicator GF$_{254}$ (20 × 20 cm, 2 mm
thickness) were used. The plates were developed with
hexane-toluene- acetone (7:2:1, v/v/v). Irradiation in
solution was performed with UV light from Hanova
mercury lamp (125 W) with two band-passes at 280/254
nm and equipped with internal water cooled quartz im-
mersing jacket. The photolysed solutions were open to
the atmosphere.

Analysis of the remaining IPC was made by GC-FID
(Pye Unicam) fitted with semipolar packed OV-17 col-
umn (1.8 m, 4 mm i.d) operated at 170°C. Mass spectra
for identification were conducted on GC-MS (Hewlett
Packard/5890) and MS. UV spectra were performed on
UV-VIS spectrophotometer (Perkin Elmer).

2.2. UV Spectra

In an attempt to evaluate photolabilaty of IPC and deter-
mine its characteristics absorption maximum, a prelimi-
nary investigation on UV spectra of IPC in methanol and
hexane were taken.

2.2.1. Photoirradiation in Aqueous Solution

One litre solution of 200 mg/L IPC in water was intro-
duced into a UV reactor system and exposed to UV at 254 nm
for 3 hours at room temperature. The reactor was shield-
ed with aluminium foil for protection. Aliquots of 25 mL
were taken periodically every 30 min starting from zero
time, followed by extraction with methylene chloride.
The combined organic extract was washed with deion-
ized water, dried over anhydrous Na$_2$SO$_4$, filtered and
finally evaporated under reduced pressure using rotary
evaporator. The remaining brown red viscous residue
was reconstituted in a small amount of methanol and
completed to 5 mL volume. The collected samples were
analysed for the disappearance of IPC using GC-FID
which was checked periodically for linearity using stan-
dards of the herbicide in methylene chloride. Portions of
the collected samples were also subjected for qualitative
analysis by GC-MS. To examine the effect of acetone as
a co-solvent and photosensitizer, the above experiment
was repeated in presence of 3% acetone. The change of
propham amount with time of irradiation is shown in
Figure 2.

2.2.2. Photoirradiation in Organic Solvents

One litre solution of 200 mg/L IPC in hexane, methanol
was similarly irradiated at room temperature for 3 hours.
Aliquots of 5 mL were withdrawn periodically and ana-
lysed for disappearance of the herbicide. After the ter-
mination period the remaining solution was evaporated
to dryness. Eventually the red viscous residue was redis-
solved in methanol and qualitatively analysed using
GC-MS instrumentation.

Propham in methanol was similarly irradiated and
analysed. Controlled blank experiments were reserved in
a dark room.

2.2.3. Separation and Identification of Photoproducts

The photolysate from different solvents used were care-
fully chromatogrammed on preparative TLC plates with
fluorescent indicator. Plates were developed in a mixture
of hexane-toulene-acetone (7:2:1, v/v/v) respectively.
Following development, plates were visualised under UV
light, the localised bands were scrapped, taken in
methanol and finally analysed by MS spectrometer.

Figure 1. Photodegradation profile of ln IPC (y) with time
value (x) in various media with regression equations for

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\begin{align*}
\text{water: } & y = 0.002x + 4.95, \quad r^2 = 0.990, \\
\text{water/Acetone: } & y = 0.003x + 5.88, \quad r^2 = 0.977, \\
\text{MeOH: } & y = 0.009x + 4.52, \quad r^2 = 0.974, \\
\text{Hexane: } & y = 0.012x + 4.58, \quad r^2 = 0.997.
\end{align*}
\]
Identifications were made on the basis of parent molecular ions (M⁺), mass fragmentation pattern and whenever possible by comparison with mass spectra of literature analogues.

3. Results and Discussion

From the spectral data of propanil in hexane and methanol it was expected that the herbicide was not a good candidate for direct photolysis. As it showed negligible absorbance above solar cut off wavelength and down to the maximum output of UV mercury lamp used. This is in accordance to the fact that only absorbed radiation is effective in producing chemical changes (Grothuss-Drapper law). However, IPC in hexane and methanol showed two intense maxima at λ (ε × 10⁶) 203(18.0), 233(16.5) and 205(14.2), 236(16.4), respectively and were assigned to π – π* transition of the carbonyl bond.

The undertaken preliminary lab investigation in this context suggested that IPC is a photo-susceptible compound as its solution in aqueous and organic solvents became increasingly yellow and turbid with some adhered to the wall of the photoreaction vessel, in contrast to its intact solution.

Initially the use of aqueous solvents was used because water is the most universal polar solvent available and might come into contact with the applied herbicides in the environment. The addition of acetone to aqueous solution of IPC was chosen to act as a co-solvent and a triplet photo-sensitizer that mimics the sensitising effect of humic matter in natural water [28,29]. However, the use of organic solvents was conducted to enhance the solubility of IPC and facilitate its direct injection into the chromatographic system.

The mode of degradation was carried out by monitoring its remaining concentration as a function of time compared to its initial concentration. Figure 2 demonstrates the photolysis of IPC in aqueous and organic solvents at a concentration of 200 mg/L. From the representative graph it is clear that the disappearance of IPC follows 1st order kinetic in agreement to what has been reported for most herbicides [30,31]. The photolysis rate of IPC in different solvents demonstrated similar behaviour with an order of hexane > aqueous acetone > methanol > water. This trend is due to differences in the routes followed for transformation [32]. Homolytic cleavage with subsequent hydrogen or solvent abstraction was a general and dominant mechanism in organic solvents [32]. The slower rate in polar solvents as compared to that in hexane could be assigned to the association of the herbicide with protic solvent via hydrogen bond in agreement with literature report [22], or due to stabilization of the excited state of its bi-radical cage [ArNH ČOOH] [19,33,34]. The higher rate of disappearance in hexane is attributed to its hydrogen radical donating ability as compared to that of methanol and water. The GC-MS and mass spectral analysis of the photolysates together with the separated photo-products revealed the formation of various compounds. The major photo-products in hexane were two isomers of IPC with molecular ions, m/z⁺, 197 with different Rf values and were assigned as isopropyl ortho- and para-aminobenzoate, additionally two solvolysis isomers with molecular ion m/z⁺, 263 were identified as hexylpropanil.

Those products were expected to proceed via homolytic cleavage of the carbamoyl bond giving either a bi-radical cage [ArNH ČOOH] or normal free radicals ArNH & ČOOH [33,34]. In both cases the radicals were able to abstract hydrogen atom and undergo concerted photo-Fries intramolecular rearrangements to ortho or para positions or else the radical may delocalise over the entire ring and abstract solvent molecules.

The photoirradiation of IPC in methanol provided variety of products including two isomers of ortho- and para-methoxy IPC with m/z⁺, 209, two coupled dimers with m/z⁺, 356 in contrast to the situation in hexane, hydroxymethylphenol m/z⁺, 123, aminocinnoline derivative with m/z⁺, 193 and isopropyl aminobenzoate isomers, m/z⁺, 179.

Eventually, the photolysis of IPC in aqueous media afforded two isomers of ortho- and para-hydroxy IPC, m/z⁺, 195 with different Rf and tᵣ values, two coupled derivatives with m/z⁺, 356, two isomers of IPC m/z⁺, 179 and benzazolone, m/z⁺, 135, together with an unidentified polymeric matter. The formation of benzazolone derivative is a result from internal thermal cyclisation of the respective hydroxy-IPC [16].

In this context it is worth noting that the demonstrated modes of IPC phototransformation i.e. solvolysis, photo-Fries intramolecular rearrangement and dimerization were not reported in any previous study. However, these have similarities with other photolysed herbicides [21, 22,33-35] or some have been reported to occur for IPC in animals [13-15]. Additionally phenylisocyanate, isopropanol, di-isopropyl ether and amino phenols were also identified from the MS spectra and could be attributed to the hydrolysis of the carbamoyl and ester bond as previously reported [18].

4. Conclusions

The result of this investigation revealed the photolability of IPC under the influence of short UV radiation in polar and non polar media. The rate and route of IPC photodegradation were affected by the nature and polarity of the photolysed media. The principal routes of IPC disap-
pearance were solvolysis, hydrolysis, photo-Fries concerted rearrangement and dimarization. These finding indicate the photolabilaty of IPC under UV exposure and provides valuable information both to chemists and environmentalists.

5. References


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