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PESTICIDE FORMULATIONS

Modification of Liquid Chromatographic Method for Analysis of Rotenone Formulations

RODNEY J. BUSHWAY

University of Maine, Department of Food Science, Orono, ME 04469

Modifications have been made on the official first action liquid chromatographic method for analysis of rotenone formulations. Methanolwater was replaced with acetonitrile-water as the eluant and methanol was substituted for dioxane as the extracting solvent. The extraction procedure has been changed from 1½ h of rotary shaking to 5 min of sonication. The rest of the procedure is identical to the published method. For 9 different products analyzed 6 times each, the percent coefficients of variation were all below 2.40. Five of these 9 samples were previously used in the rotenone collaborative study. A comparison of results from both studies showed that the modifications did not affect the percent rotenone found. It is recommended that these changes be tested in a mini-collaborative study.

In 1982, a reverse phase liquid chromatographic analysis of rotenone formulations was successfully collaborated and adopted official first action (1). Once completed, several collaborators suggested possible ways to improve the method. Also, interferences from naphthalene and sulfur occurred when a pesticide manufacturer tried to use the collaborated method on a formulation mixed with these 2 chemicals. Naphthalene was not resolved from rotenone, and sulfur eluted late, affecting other injections.

Because of the suggested improvements and the formulations problem, an investigation was undertaken to determine if the official LC method for rotenone would be changed to make it better and more versatile. This paper describes these improvements.

METHOD

Apparatus and Reagents

(a) Liquid chromatograph.—Waters Associates (Milford, MA) 6000A pump, U6K septumless injector, Schoeffel variable wavelength UV detector (Westwood, NJ), and Omniscribe recorder (Houston Instrument, Austin, TX). Operating conditions: injection volume, 5 μ L; flow rate, 1.1 mL/min; wavelength, 280 nm; absorbance range, 0.4 AUFS; recorder setting 10 mV; chart speed, 1.0 cm/min.

(b) Chromatographic column.—Partisil 5 ODS-3, 5 μ m particle size, stainless steel, 25 cm \times 4.6 mm id (Whatman Inc., Clifton, NJ).

(c) *Mobile phase.*—Acetonitrile-water (70 + 30). All solvents LC grade (Fisher Scientific, Fair Lawn, NJ).

(d) Sample extraction solvent.—Purified grade methanol (Fisher Scientific).

(e) Rotenone standard solution.—Weigh 20 mg 99% rotenone (Penick Co., Lyndhurst, NJ) into actinic 50 mL volumetric flask and dilute to volume with purified grade methanol. Actinic glassware is necessary because rotenone degrades readily in most types of light.

Preparation of Sample

(a) Dust.—Weigh sample equivalent to 20 mg rotenone into glass-stopper Erlenmeyer flask. Pipet in 50 mL methanol and sonicate in sonic bath (Bransonic 32, Fisher Scientific) 5 min while swirling gently. Let settle, and filter 5–10 mL aliquot through 0.45 μ m filter (Millipore Corp., Bedford, MA). Inject 5 μ L into liquid chromatograph.

(b) Liquid.—Weigh sample equivalent to 20 mg rotenone into glass-stopper Erlenmeyer flask. Pipet in 50 mL methanol and shake to dissolve. Filter through 0.45 μ m filter (Millipore Corp.). Inject 5 μ L into LC system.

Determination

Inject standard, followed by 2 injections of sample. Finally inject another standard. Measure peak heights, then average, and substitute into formula below:

% Rotenone = $(H/H') \times (W'/W) \times \%$ purity of std

where H and H' = average peak heights of sample and standard, respectively; W' = g rotenone standard/50 ml and W = g sample extracted.

Results and Discussion

After the rotenone collaborative study, participating we entists made numerous suggestions as to possible improvements for the procedure. Many of these suggested changes could be incorporated into the collaborative report without further study. However, 2 possible modifications required extensive investigation before they could be adopted into the procedure.

The first suggested change was replacing the solvent system (methanol-water) with acctonitrile-water; such a modification would reduce the back-pressure on the column. High pressure can shorten the normal life span of a column, injuctor, and pump. Back-pressure decreased from 2300 psi (methanol-water) to 800 psi (acctonitrile-water).

Modifying the extraction procedure was the other major change, resulting, in part, from a mistake by one of the collaborators in which methanol was used for extracting instead of dioxane and, in part, from a radial compression method (2) which was developed using sonication for the extraction technique. These led to an extraction method involving a 4 min sonication step with methanol in place of the 1½ h diox ane procedure.

The effects of these changes on rotenone analysis were evaluated. Preliminary work was performed using a sample previously analyzed by collaborators and found to be the most difficult because of interfering compounds. When the formulation was analyzed using acetonitrile-water, rotenone eluted in 6.7 min while it eluted in 10.2 min with methanol water. Other compounds were also affected by the solvent changes. Rotenolone, deguelin, and tephrosin are resolved from each other in methanol-water, but tephrosin co-clum matographed with a large peak believed to be an aromatik hydrocarbon which elutes before rotenone. Analysis of this rotenone sample with acetonitrile-water only partially suparated rotenolone and tephrosin. The aromatic hydrocarbon and deguelin co-chromatographed, but the hydrocarbon clutoit after rotenone. A comparison of percent rotenone values iii this formulation agreed well between the solvent systems.

Because of the correlated results obtained from the preliminary investigation with acetonitrile and methanol, a full scale study was performed using 5 samples from the 1982 collaborative study along with 4 others. These 9 samples were

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Table 1. Analysis of rotenone formulations
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Sample	Formulated, %	Modified method		Collaborative method	
		Found, %	CV, %	Found, %	CV, %
1. Liquid	5.0	4.55*	1.65	4.76*	2.47
2. Dust	1.0	1.00	1.58	0.99	3.19
3. Dust	5.0	4.86	1.96	4.87	2.54
4. Dust	20.0	17.57*	1.16	18.14*	2.62
5. Dust	0.75	0.88	1.68	0.89	3.19
6. Liquid	0.75	0.83	1.98		
7. Dust	0.20	0.18	2.28		
8. Dust	1.0	1.08	1.52		
9. Dust	34.0	38.84	2.39		

analysis.

Rotenone is sometimes formulated with other pesticides

such as carbaryl (1-naphthol may be present as a degradation

product), folpet, captan, difolatan, methoxychlor, piperonyl

butoxide, pyrethrins, naphthalene, sulfur, and other roten-

oids (deguelin, tephrosin, rotenolone, and isorotenone). None

of these compounds interfered with this modified rotenone

LC method for rotenone formulations result in a better method

because of less pressure on the system, faster analysis time,

elimination of peroxides, and ability to analyze formulations

containing sulfur and naphthalene. It is suggested that this

REFERENCES

(1) Bushway, R. J. (1983) J. Assoc. Off. Anal. Chem. 66, 796-800

(2) Bushway, R. J. (1983) J. Assoc. Off. Anal. Chem. 66, 793-795

modified method be tested in a mini-collaborative study.

In conclusion, these modifications of the official first action

Mouns are significantly different at the 0.01 level by the t-test.

analyzed 6 times each on 3 different days (2 samples per day).

The results are given in Table 1. As can be seen, the repro-

ducibility was excellent with this modified procedure. The

precent coefficients of variation ranged from 1.16 to 2.39. The

sumples analyzed using both the official and modified official

methods were quite close in agreement. However, 2 samples,

and 4, had significantly different means. A mini-collabora-

Sample 7, a louse powder formulated with sulfur and

implithalene, could not be analyzed by the conditions set forth

in the collaborative study (Figure 1). Naphthalene was not

isolved from rotenone, while sulfur was extracted into the

Johnne and eluted 1 h later to cause interferences. However,

when the acetonitrile-water system was used in conjunction

with methanol for extraction, the naphthalene peak was sep-

mated from rotenone, and the sulfur was barely extractable

In methanol, causing no problems. There is a small sulfur

live would point out any possible discrepancies.

mak (0.4 cm) that elutes at 31 min.

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