detected in the first and second fractions. This is the characteristic location for synthetic IAA (ref. 8). Relative fluorescence intensity of the solution from the first fraction was 3.4 (equivalent to 2.2  $\mu$ g of IAA in the 10 ml.); that from the second fraction was 2.0 (equivalent to 1.4 µg of IAA).

There seems to be no doubt that the native auxin in the developing banana fruit is indeed IAA. The extracted auxin was active in the Avena curvature test and behaved in a manner identical to synthetic IAA in paper, thinlayer and column chromatography. In addition, the isolated material had the same excitation and fluorescence spectra as IAA and reacted with the classic reagents of Ehrlich and Salkowski. From the fluorometric and biological estimations I conclude that the 300 g of banana fruit tissue contained 18 µg of IAA, a concentration of 0.06 p.p.m.

R. A. KHALIFAH\*

Citrus Research Center, University of California, Riverside, California.

\* Present address: 59 El-Nady Street, Maadi, Cairo, Egypt.

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## Production of Oat Callus and its Susceptibility to a Plant Parasitic Nematode

DICOTYLEDONS grown on White's medium containing 2,4-dichlorophenoxyacetic acid (2,4-D) readily produce callus which is an ideal substrate for plant nematodes<sup>1,2</sup>. Monocotyledons produce callus less readily. Morel and Wetmore<sup>3</sup> initiated callus in species of Amorphophallus with carefully balanced vitamins and added naphthaleneacetic acid (NAA), and maintained the callus by the addition of coconut milk. Carew and Schwarting<sup>4</sup> obtained callus from the embryo of rye by adding yeast extract, casein hydrolysate and 2,4-D to the medium. The production of oat and onion callus tissues is described below. The oat callus was used to culture the chrysanthemum nematode, Aphelenchoides ritzemabosi (Schwartz).

Hulled oat (Avena sativa, variety 'Sun II') seed was surface-sterilized in 20 per cent calcium hypochlorite for 30 min and onion (Allium cepa, variety 'Bedfordshire Champion') seed was sterilized in 327 mg/l. ethyl mercury phosphate for 20 min. The germinated seeds were placed aseptically in tubes containing Heller's<sup>4</sup> nutrient agar medium supplemented in various ways (Table 1). Glucose (20 mg/l.) or the same quantity of sucrose was used as the carbon source, and growth substances were added in the following concentrations: 5 mg/l. of 2,4-D; 2 mg/l. of indolyl-3-acetic acid (IAA); 25 mg/l. of NAA; 100 ml./l.

Table 1. EFFECT OF LIGHT AND ADDITIVES ON THE CALLUS FORMATION OF OATS AND ONIONS ON A STERILE MEDIUM

Additives	Oats		Onions	
	Light	Dark	Light	Dark
(1) S	_	_	-	
(2) $G + coconut$ milk	-		_	_
(3) $S + \text{casein hydrolysate} + \text{yeast}$	-		_	
(4) G + IAA	0	0	-	_
(5) $S + IAA + NAA$	+	-	+	+
(6) $S + 2.4$ -D	+	+	+ +	+ +
(7) $S + 2.4 - D + EDTA$	+ +	+	++	+
(8) $G + 2, 4 - D + IAA$	+ +	+	+	+

0, No test; -, no response; +, poor callus; ++, good callus; S, sucrose; G. glucose.

of coconut milk; 100 mg/l. of casein hydrolysate; 500 mg/l. of yeast extract; and 1 mg/l. of diaminoethanetetraacetic acid (EDTA). The tubes were kept at 24° C, half in the light and half in the darkness.

Callus formed only in the presence of IAA, NAA and 2,4-D (Table 1). Within 6 weeks of inoculation, the medium containing 2,4-D, IAA and glucose produced a firm, golden oat callus, which was maintained through several sub-cultures for 3 yr. Onion produced the best callus on the medium containing 2,4-D, EDTA and sucrose, but this wet, white callus grew slowly and survived only 12 months. Callus formed more readily in the light than in the dark.

Oat callus in tubes was inoculated asoptically with A. ritzemabosi (30 nematodes/tube), and after 6 weeks the number of nematodes increased fifty-fold. A. ritzemabosi is not a recorded parasite of oats, and the fact that oat callus tissue is susceptible parallels the behaviour of rose, apple, potato and red clover callus tissues which also are not hosts of this nematode but give callus that supports its multiplication<sup>2</sup>. When seedlings of resistant oat variety 'Manod' were sprayed with 2,4-D their susceptibility to the stem nematode, Ditylenchus dipsaci, increased<sup>5</sup>.

The fate of excess 2,4-D in monocotyledons is not completely known<sup>6</sup>, but 2,4-D affects oat tissue and dicotyledons similarly in that it releases the natural plant growth substances and increases cell activity, which provide an internal plant environment favouring the multiplication of nematodes.

J. M. WEBSTER

Rothamsted Experimental Station,

Harpenden, Hertfordshire.

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## Cytokinins in Citrus : Isolation of a Cell **Division Factor from Lemon Seeds**

STIMULATORS of plant cell division have been demonstrated in several types of fruits<sup>1-3</sup> and seeds<sup>4,5</sup>. Stimulators of cell division have been reported in the fruitlets of apple, plum, peach, pear, quince and pumpkin<sup>1</sup>. Explants from the secondary phloem of carrot roots were used for bioassay. A highly active kinetin-like preparation has been isolated from peas and has been described as containing at least one active purine derivative which is different from kinetin<sup>5</sup>. Cell division factors from apple fruitlets and coconut milk have also been reported<sup>3</sup>.

The study of naturally occurring hormones of Citrus has revealed the existence in various tissues of auxins<sup>6,7</sup>, gibberellins<sup>8,9</sup> and inhibitors<sup>10</sup>. An important group of hormones, namely, cytokinins, was not investigated in Citrus. The work reported here represents a preliminary investigation of the cytokinins in lemon seeds.

Ten pounds of lemon seeds (Citrus limon, Linn.) were dried in an oven at 60° C and ground up. The powder was extracted with hexane and then with methanol as described previously<sup>11</sup>. The methanol extract was evaporated in a vacuum at 50° C and the residue was dissolved in 200 ml. of distilled water. The pH of the water solution was adjusted to 3.0 with hydrochloric acid and the solution was washed four times with ethyl acetate to remove cell division inhibitors<sup>12</sup>. The aqueous phase was used in the bioassay after being adjusted to pH 5.6 with sodium hydroxide.

We used tobacco callus tissue in the bioassay and the basal medium was a modification of that described by Murashige and Skoog<sup>13</sup>. The organic constituents of the basal medium were sucrose (30.0 g/l.), myo-inositol