

A modified airstair apparatus for studies of allelopathy and other phytotoxic effects

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Abstract. The characteristics and performance of a modified airstair, nutrient solution recycling, apparatus are described.

An experiment in which the allelopathic potential of *Agropyron repens* was examined showed the apparatus to be reliable and accurate in operation. Allelopathic activity by *A. repens* was confirmed.

The apparatus is also suited to studies of phytochemicals produced in stubble retention reduced tillage systems, in green manuring and in cognate areas.

1. Introduction

The presence of phytochemicals and the roles which they play in agricultural systems are receiving increasing attention in the literature. Allelopathic phenomena (RICE 1974, 1979), whilst often difficult to distinguish among the interactions which take place between climate, soil, plants and other organisms (LOVETT 1982), occur commonly and demand better understanding than is presently the case. Phytotoxins are also of interest in reduced cultivation systems, especially where residues of crop and weed plants are retained

and decompose at rates determined by ambient climatic and edaphic conditions, by the activity of micro-organisms and by the method of cultivation (LOVETT et al. 1982).

In the agricultural systems of Finland both allelochemicals and phytotoxins produced during the decomposition of plant residues in cropping systems seem likely to be of importance. Allelopathy has, for example, long been associated with *Agropyron repens* (L.) Beauv. (AAMISEPP 1970) one of Finland's most important weed species. Living *Agropyron* may produce allelochemicals but greater phytotoxicity is usually associated with decomposition of the rhizomes (WELBANK 1963). LYNCH et al. (1980) suggest that short chain aliphatic acids are produced

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where decomposition is under anaerobic conditions. Such conditions also favour the production of similar substances, of proven phytotoxicity, from crop residues (LYNCH 1977).

Edaphic conditions favourable to the activity of facultatively anaerobic micro-organisms frequently occur in the heavy soils of Finland, particularly in autumn and early in spring. The possibility of phytotoxin production from *A. repens*, crop and weed residues is high at that time of year.

WILSON and RICE (1968), BELL and KOEPPE (1972) and WALTERS and GILMORE (1976)

have all applied the staircase apparatus to studies of allelopathy. The technique seeks to distinguish between competition and allelopathy as components of interference (*sensu* HARPER 1977) by providing a nutrient flow through a closed system in which the interfering plants are physically separate but subject to the effects of phytochemicals released into the nutrient flow. In this paper we describe a modified staircase apparatus which had wide application in studies of phytochemicals in agricultural systems.

Table 1. Comparison of allelopathy experiments using the staircase apparatus.

Reference	WILSON and RICE (1968)	BELL and KOEPPE (1972)	WALTERS and GILMORE (1976)
Allelopathy of:	<i>Helianthus annuus</i> L. with nine test species.	<i>Setaria faberii</i> Herrm. with <i>Zea mays</i> L.	<i>Festuca arundinacea</i> Scrib. with <i>Liquidambar styraciflua</i> L.
APPARATUS			
No. of lines		5	4
No. of steps	.. not specified	4	4
No. of replicates		4	3
Method	Gravity flow and recycle 4 h/day. Nutrient solution replenished.	Gravity flow and recycle 4 h/day. Full strength Hoagland's Solution replenished. pH and conductivity of solution monitored.	Gravity flow and recycle 4 h/day. Half strength Hoagland's Solution replenished. pH of solution monitored
Environment	Greenhouse.	Greenhouse. 16 h photoperiod with supplementary light. Temperature: 21°C to 30°C.	Greenhouse. 16 h photoperiod with supplementary light.
Medium	Quartz sand.	Quartz sand.	Quartz sand.
Test substances	Live <i>H. annuus</i> plants.	Live and dead <i>Setaria</i> material.	Live and dead <i>Festuca</i> material. Quantity of tissue related to field conditions.
Parameters monitored	Dry weight of test species.	Fresh weight and dry weight of test species.	Height, dry weight and content of N, P, K, Ca and Mg of test species.
Duration	Four weeks.	Four weeks.	Sixteen weeks.
Observations	Allelopathic effects on five species, including autotoxicity.	Allelopathy by substances from dead <i>Setaria</i> material.	Allelopathy by substances primarily, from dead <i>Festuca</i> tissue.
Allelopathic effect	Dry weight reduced.	Reductions of height, fresh weight and dry weight.	Reductions in dry weight. Possible effects on chlorophyll, N and P content.
Primary mechanism	No information.	No information.	Possible effect on mineral uptake.
Allelochemicals	Chlorogenic and isochlorogenic acids suspected.	No information.	No information.

2. Material and Methods

Development of the stairstep technique

MARTIN and RADEMACHER (1960) reported an improvement to a system devised by BÖRNER (1958) where in nutrient solution circulated through a series of pots in which plants of two species were planted, alternately. Various combinations of plants were tested by MARTIN and RADEMACHER (1960) who met »the most important technical requirement of the experiment«, the prevention of depletion of nutrients in the system, by regular supplementation. There were some, apparently, allelopathic effects but no allelochemicals were identified.

The MARTIN and RADEMACHER apparatus does not appear to have followed the stair-step principle and we assume that this is the modification of the nutrient flow system to which WILSON and RICE (1968) refer. The salient features of their technique and those of BELL and KOEPPE (1972) and WALTERS and GILMORE (1976) are summarised in Table 1.

The Viikki apparatus

A modified stairstep apparatus was designed and built at the Department of Plant Husbandry, University of Helsinki. Each nutrient solution recycling unit within the system (Figure 1) comprised five steps. From a storage tank at the top of the system (step one) solution flowed, by gravity, through a sequence of pots to a collector tank (step five) from which it was automatically recycled by an electric pump (JOHNSON L 100 Bilge Pump.).

On step two of the apparatus was an incubation tray (55 × 37 × 18 cm) from which the circulating solution was channelled to test pots on each of the third and fourth steps. The test pots (15 cm diameter, 14 cm height) were replicated four times and were individually supplied with solution from 3 mm internal diameter tubing inserted into collecting pipes below the second and third steps.

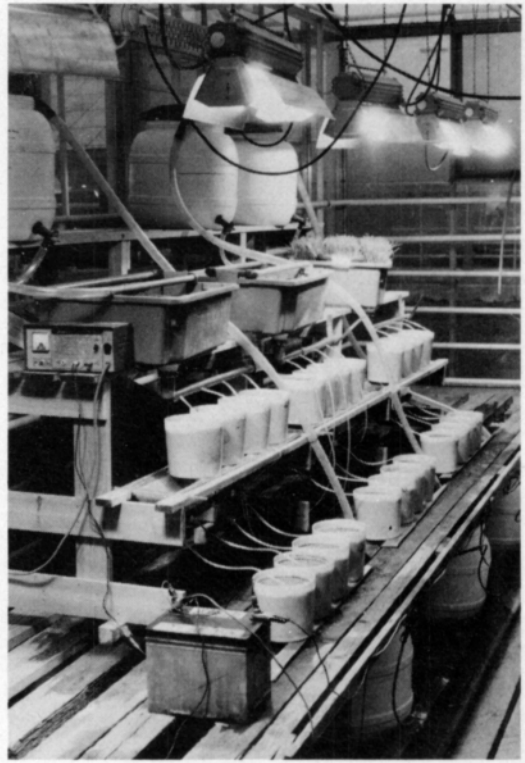


Fig. 1. The Viikki apparatus.

The incubation trays were lined with nylon mesh (0.6 mm aperture) on which was placed a layer of gravel (4.60–5.70 mm particle size, 15.4 kg dry weight). A second mesh, above the gravel base, was covered with sand (0.50–1.20 mm particle size, 18.0 kg dry weight) and the entire system thoroughly flushed with tap water. Individual test pots were filled in similar fashion and contained 770 g gravel and 2.3 kg sand.

Three recycling units were set up in a greenhouse (mean temperature $21^{\circ}\text{C} \pm 2^{\circ}\text{C}$). A 16 h photoperiod was used, supplementary illumination being provided throughout by A/S Bergen RS 400, Lucalox U 400/40 lamps. The lamps did not affect temperature at the different step levels but there was a drop in light intensity from $370 \mu\text{Em}^{-2}\text{s}^{-1}$ to $200 \mu\text{Em}^{-2}\text{s}^{-1}$ between the third and fourth steps ('Top' and 'Base' sets of test pots).

The volume of 30 l 0.2 % 'Kemira Neste-mäinen Y-lannos' complete nutrient solution (pH 7.2, N 140 mg/l, P 20 mg/l, K 120 mg/l, Na 16 mg/l, S 14 mg/l, B 54 µg/l, Mo 12 µg/l, Fe 18 µg/l, Mn 6 µg/l, Ca 4 µg/l, Mg 1.2 µg/l, Cu 0.6 µg/l, Zn 0.6 µg/l, Co 0.6 µg/l), made up with tap water, in the storage tank was replenished on alternate days. Flow rate of solution through the system was adjusted to 23 l/h and the solution was cycled for 6 h/day.

Application of the apparatus

The modified staircase apparatus was used in an assessment of the allelopathic potential of *Agropyron repens* (L.) Beauv. (couchgrass).

Couchgrass was freshly harvested from the field at Viikki. Top growth was cut back to 3 cm height and the rhizomes thoroughly washed in tap water. Into the incubation tray of one unit was placed 1.2 kg (fresh weight) rhizomes with top growth, an amount which corresponded to that recovered from an equivalent volume of soil in a badly contaminated field. A similar weight of material was finely chopped and incorporated into the sand at 10 cm depth in a second incubation tray, where it decayed. The third, control, tray received only nutrient solution.

Each experiment in a series ran for one week. The height couchgrass foliage in the 'Live' treatment was maintained at, or near, 10 cm height. Any shoots appearing from the 'Decay' treatment were removed.

The test species was *Hordeum vulgare* L. (barley) cv. Kustaa. Into each test pot were sown 14 graded seeds (2.5 to 2.8 mm diameter). Emergence was monitored from its commencement on Day 3 of each experimental run. On day 7 the plants were carefully washed from the test pots. Root loss was minimal. The height of the first leaf and length of the longest seminal root were determined. The plants were divided into leaf, seed and root portions which were dried at 100°C for 48 h and weighed.

It was possible to thoroughly clean and refill the test pots with fresh gravel and sand; to clean the tubes in the recycling system, and to commence the next run on the day of harvest.

Data were analysed using a statistical package available through the Helsinki School of Economics Computer Centre. Analysis of Variance was applied to untransformed data. Variability was small and there was, for example, no necessity to re-randomise pots within steps as the flow rate of nutrient solution and the ambient conditions were uniform.

3. Results and Discussion

Previous examples of the use of the staircase apparatus (Table 1) provide little information on performance. The Viikki apparatus proved simple and reliable in operation. pH was monitored daily and did not vary from the initial value of 7.2. The NO₃ concentration of the solution in the recycling units varied from 133 to 146 mg/l as compared with 151 mg/l in unused solution. Germination tests, using solution pipetted from the recycling units whilst in operation, indicated no contamination by pathogens which might affect growth of the test species. Some algal growth was, however, observed.

Developments from earlier versions (Table 1) include the provision of large incubation trays, as compared with pots of 12 cm diameter (BELL and KOEPPÉ 1972) or 19 cm diameter (WALTERS and GILMORE 1976). The relative homogeneity of phytotoxin-producing material, the amount of which was related to field conditions, and the possibility of maintaining the material for indefinite periods in large containers are significant advantages. They permit, for example, monitoring of phytotoxin production over time, and sequential harvesting as compared with single harvests in earlier experiments, Table 1.

A difference in light intensity between steps was observed by WALTERS and GILMORE (1976). This occurred also in our experiments

but there was no difference in ambient temperature. Statistically significant effects of position were recorded for shoot height at harvest (Table 2) and for shoot and total dry weight.

Whilst data are here presented, primarily, in validation of the technique, they confirm the allelopathic potential of *A. repens*. Thus the presence of decaying or live couchgrass material affected the test species from the commencement of growth and was evidenced in morphological and physiological characteristics at harvest, Tables 3 and 4. The apparently progressive decline in total dry weight (Table 4) is a consequence of delayed sowing in Run 2 whilst the system was cleaned and of the removal of the second leaf of all plants in Run 3 for the determination of chlorophyll content.

The primary causes of the effects and the possibility that different chemicals are produced by live (GABOR and VEATCH 1981) and decaying (LYNCH 1977) material of *A. repens* are the subject of further investigations.

In earlier experiments, alternate pots of phytotoxin-producing and test species have been employed (BÖRNER 1958; MARTIN and RADEMACHER 1960; WILSON and RICE 1968; BELL and KOEPPE 1972; MARTIN and GILMORE 1976). Only WILSON and RICE (1968) used species which demonstrated that autotoxicity could occur in a staircase apparatus. The type and amount of phytochemicals produced by autotoxified plants may, of course, differ from that of non-toxified material.

Table 2. Effect of position of test pots on height or barley shoots (means of four replicates).

Position	Mean shoot height (cm)		
	Run 1	Run 2	Run 3
Top	18.58	19.37	18.05
Base	17.58	18.78	17.15
p	< 0.001	< 0.01	< 0.001
LSD (t) p < 0.05	0.52	0.34	0.52
p < 0.01	0.71	0.47	0.71
Coefficient of Variation (%)	3.36	2.08	3.43

Table 3. Effect of couchgrass on length of barley roots (means of four replicates).

Treatment	Mean root length (cm)		
	Run 1	Run 2	Run 3
Control	11.11	13.91	10.75
Decay	9.97	12.48	11.25
Live	9.71	11.25	12.42
p	< 0.05	< 0.001	< 0.001
LSD (t) p < 0.05	1.01	0.84	0.72
p < 0.01	1.38	1.15	0.98
Coefficient of Variation (%)	9.32	6.35	5.95

Table 4. Effect of couchgrass on total dry weight of barley (means of four replicates).

Treatment	Mean total dry weight (mg)		
	Run 1	Run 2	Run 3
Control	39.69	32.78	29.36
Decay	39.94	30.86	29.02
Live	39.34	30.64	27.38
p	N.S.	< 0.05	< 0.01
LSD (t) p < 0.05	—	1.66	1.21
p < 0.01	—	2.27	1.66
Coefficient of Variation (%)	5.58	5.02	4.02

The Viikki apparatus does not exclude the possibility of autotoxicity, however, output from the incubation trays is filtered through two series of test pots before recycling, reducing the possibility of its occurring. This proposition is supported by the interaction between position and treatments (Figure 2),

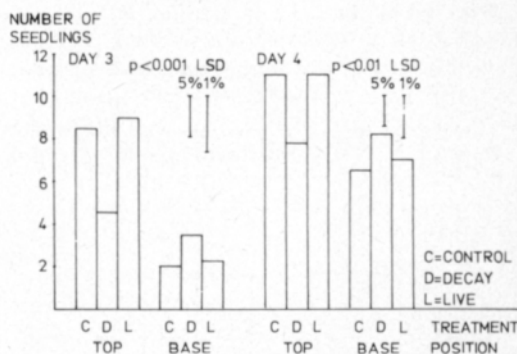


Fig. 2. Position by treatment interaction in emergence of seedlings. Run 2.

in which an apparent inhibition by decaying couchgrass at the top position becomes a stimulation at the base. This effect implies a reduced concentration of phytochemicals, and the type of response noted by LOVETT (1982).

More detailed information on allelopathy by *A. repens* as manifest in the staircase apparatus will be published elsewhere. The apparatus is equally suited to studies of phytochemical production during decomposition of crop and weed residues in reduced cultivation systems or where green manuring is practised.

Given that the growing season in Finland

is short it is desirable to minimise the stresses to which crop and pasture plants are subject, particularly during early growth and development. Understanding phytochemical effects is of potential significance in reducing stress and improving productivity in many agricultural systems.

Acknowledgements: The Finnish Academy of Sciences generously funded the visit, organised by Professor Eero Varis, which made possible the participation of one of us (J. V. L.) in this programme. The equipment was skilfully constructed by kenttämestari Reino Hakala.

Our thanks for these several sources of support are gratefully recorded.

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Ms received December 1, 1984

SELOSTUS

Muunneltu uuttojärjestelmä allelopaattisten ja muiden fytotoksisten vaikutusten tutkimiseksi

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Tässä työssä kuvataan kiertävään liuosjärjestelmään perustuvan laitteiston ominaisuuksia ja käyttöä. Koe, jossa tutkittiin juolavehnan mahdollista allelopaattista vaikutusta, osoitti, että kehitetty laitteisto on käytössä luotettava ja tarkka. Koe vahvisti myös käsityksiä, joiden mukaan juolavehnä on allelopaattisten yhdisteiden tuottaja.

Laite on sovellettavissa myös tutkimukseen, jossa selvitetään eri viljelymenetelmissä mahdollisesti esiintyvien fytokeemikaalien vaikutusta kasvien alkukehitykseen.