Several aspects of the ecology and pathology of Fusarium oxysporum f. sp. cepae in onion soils, G. S. Abawi (Cornell Univ., Ithaca, N. Y.). A direct correlation was found between inoculum density of Fusarium oxysporum f. sp. cepae and damping-off of onion seedlings growing in organic soil artificially infected with the fungus under controlled environmental conditions (26 C-day and 21 C-night; 16 hr fluorescent light/day at 2000 ft-c; 60-70% relative humidity). A population of 5 x 10^6 or more propagules/g oven-dry soil was needed before any significant development of disease could be detected in field soil; 100 propagules caused extensive disease development in sterilized soil. Seedling damping-off increased from 10 to 32 C. Conidia added to field soil exhibited a low percentage of germination, and germ tubes formed were either lysed or converted to chlamydospores. Population of the fungus decreased in the absence of onion and increased in its presence. Roots of Oxalis corniculata were heavily infested by C. cepae when the plant was grown in artificially infested organic soils.

Some effects of pH and nitrogen source on growth of Pythium obursa and P. ultimum in culture. J. J. Albert & C. R. Drake (Va. Polytech. Inst., Blacksburg). Pythium obursa was cultured in liquid synthetic media containing KNO3, NH4NO3, (NH4)2SO4, and L-aspartic acid as nitrogen sources, and sucrose as the carbon source, at initial pH values of 5.5, 5.0, 7.0, and 9.0. The medium also contained KH2PO4, 1.0 g; MgSO4·7 H2O, 0.5 g; Fe(NO3)3·9 H2O, 0.2 mg; MnSO4·4 H2O, 0.1 mg; thiamine, 0.1 mg; and biotin, 0.005 mg. Standing cultures were incubated at 28 C and harvested at 4-day intervals for 24 days, and the mycelium was dried and weighed. At 16 days, dry weights in mg at initial pH values of 5.5, 5.0, 7.0, and 9.0 were: 25, 24, 22, and 26 mg, respectively: KNO3, 207, 238, 232, and 249; NH4NO3, 82, 157, and 225; (NH4)2SO4, 30, 51, 108, and 193; aspartic acid, 153, 153, 151, and 152. There was some evidence that pH of the medium affected growth, independent of the nitrogen source. Nitrate nitrogen statistically supported better growth than organic (aspartic acid) or ammonium nitrogen at each harvest date. Analysis was determined using Duncan's multiple range test for significance at the 5% level. The optimum initial pH value for growth was 9.0 for all nitrogen sources except aspartic acid, where growth was similar at initial values of pH 5.5, 7.0, and 9.0. Growth was extremely sparse when nitrogen was omitted from the medium.

Pathogenicity of Cylindrocladium theae and C. scoparium in roots and leaves of azalea. S. A. Albert, Jr., R. G. Linderman, R. H. Moore, & E. K. Sorensen (Fl. Dep. Agr., Gainesville, ARS, USDA, Beltsville, Md., Univ. Minn., St. Paul, Univ. Ga., Coastal Plain Sta., Tifton). Cylindrocladium theae (Calonectria theae) was isolated from roots, stems, and leaves of several cultivars of greenhouse azalea. It was pathogenic when compared with C. scoparium to roots and leaves of three greenhouse and three hardy cultivars of azalea. No mortality occurred among plants grown for 2 months in soil artificially infested with comparable amounts of blended cultures of each species. However, significant root discoloration was noted in two of six cultivars grown in soil infected with C. theae, and the fungus was reisolated from all but one cultivar. Two of six cultivars grown in soil infested with C. scoparium showed reduced and discolored root systems; three cultivars exhibited yellow discolored roots only. The cultivar had apparently healthy roots from which the fungus could not be isolated. The leaves of all cultivars showed some degree of susceptibility to both species. Symptoms varied from a few irregularly circular, purplish-black lesions in C. theae that covered a large portion of the leaf surface, with slight-to-heavy defoliation.

In general, C. scoparium was more virulent to leaves and roots than was C. theae.

Nuclear phenomena preceding basidiospore formation by monobasidiospore isolates of Lentinus taevea. T. L. Amorese (USDA Forest Serv., Gulfport, Miss.). Some monobasidiospore isolates of Lentinus taevea have the capacity to form fruit bodies and viable basidiospores in culture. This study was initiated to examine this phenomenon cytologically. Most of the probasidia produced by monobasidiospore isolates do not form sterigmata in the usual sense and spores, but continue to grow by either apical or lateral proliferations. In mature basidia produced by monobasidiospore isolates, the nuclear cycle is the same as that in dikaryotic isolates. A fusion nucleus is formed which then divides to form haploid nuclei. These then migrate through the sterigmata and into the developing spores. The nuclei usually divide mitotically as the spores mature. In basidia produced by dikaryotic isolates, two nuclei frequently enter a sterigma and presumably both enter the developing spore. If the two nuclei entering a sterigma contain the same sexual incompatibility factor (i.e., A1 and A2), the spore subsequently containing them, although heterokaryotic, would give rise to simple-septate cytologism (assuming there is a dissimilar sexual incompatibility factor is necessary for clamp-connection formation). Isolates derived from such spores may be the monobasidiospore isolates which have the capacity to fruit in culture.

Ultrastructural determination of the infection process of Macrospora phaseoli in Glycine max. V. D. Ammon, T. D. Wyllie, & M. F. Brown (Univ. Mo., Columbia). Surface-sterilized seed of the soybean cultivars Amsony and Adelphia were grown in steam-sterilized vermiculite for 2-4 days. In addition, seed from the cultivars Amsony and Calland were germinated in petri dish cultures of Macrospora phaseoli. The root systems of the plants grown in pots were exposed to the charcoal bed fungus by injecting into the rhizosphere 50 ml of an actively growing shake culture. After incubating for 1-10 days, selected roots were fixed and embedded in paraffin and in plastic for light and electron microscopy, respectively. Roots of all cultivars of soybean seedlings tested became infected within ca. 3 days as determined by direct observation. Fungal invasion of root tissue occurred intercellularly and intracellularly. Penetration of soybean root tissue was accomplished by changes in the organization of the cytoplasm, disruption of the membranes, and hyphae potassium cell walls. The disruption of cellular and tissue integrity is believed to occur by the production of pectolytic enzymes in association with mechanical pressure.

Heterokaryon formation in Thanatophorus cucumeris ("Pratola"). N. A. Anderson, H. M. Stretton, & N. T. Elsted (Univ. Minn., St. Paul, Waite Inst., Glen Osmond, S. Australia). Field isolates of Thanatophorus cucumeris ("Pratola") type produce single basidiospore progeny which segregate as if a single factor (H factor) controlled heterokaryon formation. When monokaryons of different H factors are paired, a tuft of hyphae forms where the colonies join. Thiamin and nitrate auxotrophs were paired with wild-type monokaryons of different H factors. Hyphal tip colonies established from the tuft mycelium produced basidiospores containing the parental fac- tory, indicating that the tuft hyphae were heterokaryotic. Five single-sporal isolates which failed to form tufts with either parental H factor were found upon sporulation to contain both H factors. Intrafactor recombination producing nonparental H factors occurred at a rate of 1.8% (sample 1274) of progeny from a British isolate and 2.1% (sample 420) from a USA isolate. H factors were obtained.

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from 13 field isolates from Australia, Britain, Canada, and the USA. Heterokaryons were synthesized between mono-
karyons from the four countries.

**Races of Fusarium oxysporum f. sp. pisi.** G. M. Armstrong & Joanne K. Armstrong (Ga. Exp. Sta., Experiment). During the past 5 years, 25 cultivars (CVS) of *Psilium sativum* were inoculated with 19 isolates of *f. sp. pisi* from five countries, but not all CVS were inoculated with all isolates. Three CVS were uniformly susceptible; the others were resistant or susceptible to different isolates. There were four groups of isolates based on previous identifications by the donors; 8 of race 1, 9 of race 2, one of race 4, and one of race 5. Differences in patho-
genicity of the isolates showed that races 1 and 2 are not well defined. Six of eight isolates in the race 1 group are retained in race 1 due to similarities in pathogenicity on six CVS. Two isolates differed from these and from each other and are recognized as two new races. Four of the nine isolates in the race 2 group, as well as the race 4 isolate, are similar on seven CVS and are considered as race 2. Two somewhat divergent isolates were retained with these in race 2. Of the three remaining isolates of the race 2 group, one is the same as the recently described race 5. The two races constitute a new race. Thus, five races are recognized in the collection. Races 3 and 3A were unobtainable.

**Efficacy of triarimol (EL-273) against Venturia inaequalis and Podosphaera leucotricha.** W. R. Arnold, H. B. Ladd, & C. D. Christensen (Ell Lilly and Co., Newark, Del., Plainwell, Mich., & Cazenovia, N.Y.). Triarimol (a-(2,4-dichlorophenol)-a-phenyl-5-pyrimidinemethanol) was evaluated for the control of apple scab and powdery mildew on Rome Beauty, McIntosh, Red Delicious, and Jonathan apple trees in experiments conducted in Pennsylvania, New York, and Michigan. Triarimol was applied every 7, 10, or 14 days through petal-fall, followed by a 14-day schedule for the remainder of the season with high-pressure hand guns and commercial sprayers. Triarimol was also applied on a 7-day schedule for the first three applications, followed by a 14-day schedule for the remainder of the season. Application rates ranged from 30-80 ppm. In areas with less than 10% scab inci-
dence on fruit of control trees, 40 ppm applied on the 14-day schedule for the entire season provided good-to-
extcellent powdery mildew, rust, and excellent control. In all experiments, good-to-excellent mildew control and excellent scab control were provided by 40 ppm applied on the 7- and 10-day spray schedules through petal-fall, followed by applications every 14 days. Similarly, excellent control of these diseases was obtained by applying 40 ppm of the agent, followed by applications made every 14 days. Fruit quality was not adversely affected by triarimol.

**Aggressiveness in Puccinia recondita tritici.** M. Aslam & L. E. Browder (ARS, USDA, Kansas State Univ., Manhattan). The relationship of aggressiveness to pathogenicity in *Puccinia recondita* was studied in three cultures: 66-763 (virulent at 8 out of 12 loci studied), UNO1-68A (virulent only at the P10 locus), and UNO1-68B (avirulent at all loci studied). Survival in mixtures, relative infectivity, and uredospore production were used as criteria for ag-
gressiveness. Three composites of two cultures each were grown through several generations on Biono wheat, which has no known genes for resistance. Frequency of each of these cultures in the composite set of conditions was determined by ascospore frequency on a differential set of near-isogenic lines. UNO1-68B in com-
posite with UNO1-68A and with 66-763 increased from 43% and 54% to 70% and 94%, respectively, after 6 genera-
tions on Biono; UNO1-68A in mixture with 66-763 in-
creased from 54% to 90%. A high positive correlation was found between the number of generations on Biono and the frequency of the cultures with less virulence. Predomi-
nance among cultures was unaffected by temperature or photoperiod during growth. UNO1-668B increased signifi-
cantly more uredospores per pustule than either of the other cultures. The cultures also differed in infectivity on Biono, with the most avirulent culture being significantly more infective.

The effect of oxyxcarboxin on the bean rust disease caused by *Uromyces phaseoli*. J. J. Bates & B. G. Tweder (Univ., Mo., Columbia). Healthy and diseased plants and the fungus, *Uromyces phaseoli*, were treated with various concentrations of oxyxcarboxin. Germination and respiration studies of uredospores were carried out using 50 µg and 100 µg/ml of oxyxcarboxin with samples taken at hourly intervals. 14C-glucose was used to study the effect of oxy-

carboxin on the diseased and healthy plant. Radioauto-

tography was used to study the movement of the fungicide in the diseased plant. Results indicated that the fungicide

did not stimulate spore germination but does inhibit it. Germ

tube extension and respiration were greatly inhibited. Oxyx-
carboxin had no effect upon growth or respiration rate of the bean plant at concentrations which inhibited disease development. Translocation from the roots to the leaves was found to occur in 24 hr. The concentration increased rapidly around the site of infection. After an incubation period of 72 hr, the concentration in the pustule was ca. 50 times that found in the noninfected parts of the plant.

**Proteins from cultured cells of Xanthi-nc tobacco inoculated with tobacco mosaic virus.** R. N. Beachy & H. H. Murakoshi (Mich. State Univ., E. Lansing). Cell suspensions of *Nicotiana tabacum* "Xanthi-nc" tissue culture were inoculated with 100 µg tobacco mosaic virus (TMV) ml-1, washed with fresh medium to remove excess inoculum, and cultured on agar medium under diffuse light. Local lesions were observed on inoculated cell cultures after 40 hr. At 24-hr intervals, inoculated and noninoculated control cells were harvested, and cell extracts were prepared by homogenization and centrifugation at low (15,000 g) and high (144,000 g) speeds. The supernatant from the high-

speed centrifugation was subjected to electrophoresis on 7 and 10% polyacrylamide disc gels, stained for protein in amido black, and scanned in a Gilford densitometer. On a fresh wt basis, more proteins were extracted from lesion-

bearing inoculated tissue than from control tissue. One protein band, characterized as TMV coat protein, was detected from extracts of infected cells. Intensities of sev-

eral normal host protein bands diminished, whereas others increased following infection. Data indicate that protein changes can be detected in virus-infected cells from tissue cultures more easily than in virus-infected leaves.

**Effects of fluometuron and prometryne on Rhizoctonia solani in soil.** H. Wayne Beam & E. A. Curl (Auburn Univ., Auburn, Ala.). Technical grade fluometuron and prometryne [2,4-dis(isopropylamino)-6-methylmercapto-a-

triazine] were tested in tanks with previously autoclaved sandy loam soil for effect on growth response of *Rhizo-

cotonia solani*. The herbicides were applied in a nutrient solution to provide concentrations ranging from 1 to 40 
µg/g of soil. The cultures were incubated at 27°C, and analyses were made at intervals of 3, 5, 9, and 6 days to determine soil-enzyme activity and nutrient uptake. In herbicide-free soil, there was a general increase in β-galac-
tosidase activity during the first 3 days after inoculation with the pathogen; activity changed little thereafter. En-

zyme activity for the 40-µg fluometuron treatment in-

creased rapidly, but was far less at all sampling times than for the control or other treatments. Enzyme activity was higher in the 1-µg fluometuron treatment than in the control. Values for other treatments (3 and 10 µg) were inter-

mediate. In prometryne-treated soil, the 40-µg treatment increased β-galactosidase activity during the period between 2 and 4 days, whereas treatments of 1, 5, and 10 µg en-

hanced activity. Little change in soil pH occurred in any
treatments, but ion uptake was suppressed by the higher concentrations of both herbicides.

Production of cellulase, phosphatase, polygalacturonase, and pectate lyase by prototrophic avirulent and partially virulent mutants of Erwinia carotovora. L. BERRA, B. A. BILLIET, & E. D. GABER (ARS, USDA, Univ. Chicago, Chicago, Ill.). Avirulent and partially virulent mutants of Erwinia carotovora were produced after 5- to 12-min exposure of a 10/5 ml shaken suspension of 24-h-old cells in 250 mg/1 N-methyl-N-nitro-N-nitrosoquazoline in PO, buffer at pH 7.0. The enzyme profiles of five prototrophic, independently obtained, avirulent mutants (AV) were compared with those of four partially virulent (PV) and the fully virulent (RV) and wild-type (WT) strains. In supplemented medium containing the inducers polygalacturonic acid, carboxymethylcellulose, purified soya bean lecithin, or lima bean extract, all AV strains produced low levels of cellulase, variable amounts of polygalacturonase and pectate lyase, and no phosphatase. When compared to AV, those PV strains producing the most cellulase and polygalacturonase were intermediate in virulence. Like the AV strains, all PV strains tested failed to produce phosphatase. By comparison, the virulence of the strains obtained from the AV had consistently higher unit values of polygalacturonase, pectate lyase, cellulase, and phosphatase. The manipulation of enzyme production and the resulting effects on virulence implicate these enzymes directly in the production of soft rot.

Numbers of trapped Cercospora apii spores determined by length of temperature, favorable moisture periods, and disease incidence. R. D. BERGER (Univ. Fla., Belle Glade). Sporulation laws governing in large Florida celery fields for over 1250 “trap-days” revealed that numbers of trapped Cercospora apii spores were greatly reduced when the following conditions were not met: 12 hr/day of humidities near 100% above 15 C (blight favorable hours). The numbers of spores trapped when the BFH exceeded 12 hr/day were directly associated with the amount of disease observed in the field; the highest numbers (400-700 spores/day) occurred when disease expression was maximal. Low disease incidence, small leaf areas (young plants), morning cloudiness, and high daytime humidities were the most important factors giving low spore numbers (0-100 spores/day) when seemingly blight favorable conditions existed. Except for those occasions when mechanical operations (harvesting, cultivating, weeding, etc.) detached spores, significant spore numbers (over 50 spores/day) did not occur when weather unfavorable for blight prevailed. The numbers of trapped spores provided a valid estimate of daily “blight pressure,” and to some extent estimated the amount of disease in the field. The spore numbers also provided the growers with a basis to establish satisfactory spray schedules and a means to determine the effectiveness of particular spray programs.

Nutrition and metabolism of the haustorial mycoparasite Tichocromyces parasiticus in axenic culture. F. L. BINDER & H. L. BARNE (W. Va. Univ., Morgantown). Tichocromyces parasiticus, a haustorial mycoparasite, makes no growth on hexas, penes, or trichocroacic acid intermediates as carbon sources. Excellent growth occurs in liquid media with glycerol. Cell-free extracts of glycerol-grown cultures contain all enzymes of the Embden-Meyerhof and hexose monophosphate pathways. In the presence of Tween 80, glucose is used as a carbon source. Parasite growth on C basal medium level of casein hydrolysate is high (10-40 g/liter). Fractionation of casein hydrolysate showed that the parasite had an absolute requirement for cysteine or methionine, and is highly dependent upon valine and leucine for axenic growth. Parasite growth was proportional to the available amino acids, those derived from the aspartate family, and histidine did not affect axenic growth. The parasite requires thiamine and biotin for growth. The inability of certain haustorial parasites to grow in axenic culture may be the result of multiple deficiencies of specific amino acids and vitamins, as well as problems associated with the uptake of specific nutrients.

Hosts and pathogenesis of a new Heterodera sp. on grasses. W. BIRCHFIELD (USDA, ARS, La. State Univ., Baton Rouge). A new species of Heterodera induced pathological symptoms on grasses different from those previously described for other members of this genus. Rice, Johnson grass, and barnyard grass were found to be hosts of this nematode, but no other grasses tested, including sugarcane, ryegrass, winter wheat, nor any of several dicotyledonous plants, were parasitized. Larvae penetrated the epidermis and traveled intercellularly to the vascular cylinder, where they positioned themselves with the head toward the main root turned from the root tip. They penetrated the peri- and root in the phloem between the xylem vessels. No giant cells were formed as reported for other species of Heterodera, but enlargement of phloem near the rootlet was evident. Cell walls near the feeding site were thickened, as evidenced by heavier stain absorption. Galling of roots and top symptoms were not observed on host plants. Scutellar secretions were not evident in egg layer. Swollen, females engorged and epidermis and deposited a few eggs outside, but most were retained. Males did not penetrate to the vascular area, but remained in the cortex near the epidermis.

Residue monitoring: a system for timing spray schedules. C. H. BLAIREZ (Univ. Fla., Immokalee). A system for timing spray schedules was developed by relating the decay rate of fungicide leaf residues with their ability to inhibit spore germination in a bioassay test. Foliar residues of Difolatan 45% (1,2,4-tetrachloroethyl thiocarbamide) sprayed at the rate of 3 pints/100 gal of water were washed with benzene from tomato leaves beginning 3 hr after spraying and every 24 hr thereafter until complete breakdown occurred. Difolatan residue analysis was done by thin-layer chromatography. Nelly's bioassay test was used to determine the per cent germination of Alternaria solani spores with daily concentrations of fungicide residues. Difolatan residues varied from 400 to 0 ppm at 3 hr and 7 days, respectively. Spore germination in the bioassay test increased 6 to 100% from 3 to 7 days after spraying. Concentrations above 100 ppm were fungicidal, while those below 75 ppm were few stastic. It was possible to estimate the number of days of effective protection by knowing the Difolatan decay rate and foliar residue. These results suggest that this analytical bioassay method may be used to accurately determine the proper timing of fungicidal sprays for disease control.

Winter wheat as a reservoir for maize dwarf mosaic virus. C. W. BOTTLEY & C. P. ROMAIN (Cornell Univ., Ithaca, N.Y.). Winter wheat (Yorkstar) with mosaic was found near a corn field (Pa 290) in which a few plants showed symptoms of maize dwarf mosaic (MDM). Corn seedlings (NY3 X DS0) mechanically inoculated with leaf juice of diseased corn and of diseased wheat from this field showed mosaic symptoms. The isolates from corn and from wheat were transferred mechanically to corn (NY3 X DS0) and to sorghum (Atlas, but not to wheat (Yorkstar). Each of the two isolates was identified serologically as MDMV-A. When aphids (Rhopalosiphum maidis) were placed on corn leaves mechanically inoculated with the wheat isolate, and subsequently on healthy corn seedlings, severe mosaic developed in the corn but only a mild mottle in the wheat. Mechanical inoculation of corn with leaf samples from this corn and wheat leaves, however, resulted in an expression of pronounced mosaic symptoms. In another test, Macrosiphum avenae, Myzus persicae, R. maidis, R. padi, and Schizaphis graminum were placed on corn leaves infected with the wheat isolate, then on corn and wheat seedlings. In 3
weeks, no symptoms developed on wheat, but mosaic symptoms appeared in corn exposed to all aphid species. The most severe leaf mosaic symptoms occurred in seedlings exposed to Schizaphis graminum, the least severe in those exposed to M. avenae.

The experimental infection of Solanum tuberosum by tomato ringspot virus, J. S. Boyle (Pa. State Univ., University Park). Ten 6-week-old greenhouse-grown potato seedlings were mechanically inoculated with an isolate of tomato ringspot virus (TomRSV). Initial symptoms of infection, characterized by inconspicuous and sparse chlorotic and necrotic spotting, were observed on double-leaved seedlings 6 days after inoculation. The virus was later recovered from terminal tissue from two of the plants, indicating that systemic infection had occurred. Tubers from these plants were saved and stored at 40°F until they sprouted. Three tuber generations have been produced from these plants with several unexpected results. The symptoms of chronic TomRSV infection in potato are strikingly different from any previously encountered syndrome in this plant. Many tubers formed on TomRSV-infected plants do not sprout, thus eliminating the need for future generations. Plants that do sprout from tubers are characterized by a striking chlorosis restricted to the tips and margins of leaflets. As the plants grow, symptoms fade from the old leaves but continue to develop in the young expanding leaflets. Frequently, plants will continue to grow indeterminately with large numbers of aerial tubers.

Purification and properties of the viruslike particles of Penicillium chrysogenum, R. F. Bozart & H. A. Wood (Boyle Thompson Inst., Yonkers, N.Y.). The viruslike particles (VLP's) of Penicillium chrysogenum (ATCC No. 9480) were purified by extraction with chloroform and 0.1 M, pH 7.0 potassium phosphate buffer, differential centrifugation, sucrose density-gradient centrifugation, and sucrose density-gradient electrophoresis. In addition to the major component which sediments at 150 S, there were additional sedimenting components at 81, 101, and 212 S. All components had spherical VLP's 40 nm in diameter and ultraviolet spectra typical of nucleoproteins. Nucleic acid extracts of purified VLP's made by the single-phase phenol-SDS procedure and assayed by polyacrylamide gel electrophoreses contained three classes of double-stranded RNA with molecular wt of 2.18, 1.99, and 1.89 X 10^6 daltons. Antiserum to the VLP's of P. chrysogenum reacted specifically with its homologous antigen and also with the VLP's of P. brevi-compactum. On the basis of serology and similar morphology, the VLP's of P. chrysogenum are considered to be related to those of P. brevi-compactum.

Screening red clover introductions for resistance to Staphylococcus aureus. S. W. Braverman (ARS, USDA, N.Y. State Agr. Exp. Sta., Geneva). At the Northeast Regional Plant Introduction Station, Geneva, N.Y., 550 red clover (Trifolium pratense) introductions and 23 standard cultivars were grown in the greenhouse and evaluated for resistance to Staphylococcus aureus. Seedlings were inoculated using a mixture of two cultures isolated from highly susceptible red clover accessions, increased on potato-dextrose agar, and cultured in a blender, and subsequently applied under 70 psi on 3- to 4-inch-tall 5-week-old seedlings. Inoculated plants were incubated under 100% RH at 20-24°C for 48 hr, then removed to a greenhouse bench. Plants were rated for amount and prevalence of infection 30 days after inoculation. Plant introductions 238545, 238581, and 238588 were rated as highly resistant, 50 were rated as moderately resistant, while 64 varied from slightly resistant to slightly susceptible. Wisconsin Mildew Resistant, N1-17-1-64 (FC 39493), Common Indiana Medium (FC 39370), and Medium Red (FC 40362) were slightly resistant. The remaining 28 standard cultivars were moderately to highly susceptible to the pathogen.

The use of the computer in analysis of Puccinia recondita pathogenicity data, L. E. Brown (ARS, USDA, Kansas State Univ., Manhattan). The basic model of the gene-for-gene relationship permits four experimental designs; one of these is to hold host-genotype for resistance and environment constant and study parasite pathogenicity. This design, used on a parasite sample, can be used to study pathogenicity of parasite populations in time and space. The infection-type data obtained from such experiments usually are so extensive that only the simplest analyses can be made by manual methods. The capacity of the computer was used to make detailed analyses of survey data of Puccinia recondita on Triticum aestivum. A parasite-culture X host-line infection-type matrix was put into core-storage along with appropriate indices of information about host and parasite entities used. Specific arrays of data could be associated with specific host-lines or parasite-cultures by subscripting techniques. The arrays could be compared in any combination. Programs have been developed to produce race summaries, to tabulate virulence frequencies to each host line used, to tabulate associated pathogenities to two or four host lines in all combinations, to summarize pathogenicity according to similarity to "type" cultures, and to sort data on collections by collectors' names.

Studies on the activity of triarimol (EL-273) against certain powdery mildew fungi, L. F. Brown, Jr., & H. R. Hall (El Lilly & Co., Greenfield, Ind.). Greenhouse studies have demonstrated the curative activity of triarimol against Erysiphe graminis f. sp. tritici and E. polygoni. When conidia of E. polygoni were placed on bean leaves sprayed with 1, 10, or 50 ppm of triarimol, no difference in germination or germ tube development was observed. Bean or wheat plants show no macroscopic signs or symptoms of disease when sprayed with 5-10 ppm. Triarimol at 1 ppm was applied to foliage of bean and wheat plants 0-4 days before or after inoculation. Regardless of the time of application prior to inoculation, or for 1 hr thereafter, the conidia of both fungi germinate and haustoria develop. However, no additional fungal growth occurs after the formation of haustoria in treated plants. However, application of triarimol 24, 48, 72, or 96 hr after inoculation arrests further development of these fungi. No additional haustoria develop from superficial mycelium. When triarimol is applied to soil at 0.5 g/kg soil, no macroscopic symptoms of disease occur on wheat or bean plants. Microscopic examination of treated plants shows that haustoria form, but no further fungal development occurs.

Germination of Fomes ignarius var. populinus basidiospores on aspen phšk., T. S. Brown, Jr., & W. Merrick (Pa. State Univ., Univ. Park). Controversy exists as to whether Fomes ignarius var. populinus (F) penetrates through branch stubs. Some workers have reported that germination of F occurred only after 7 days' incubation in fresh wounds, and that F occurred in branch stubs only as a lateral outgrowth from an internal decay column. Following our observations that sometimes the pith of branch stubs appeared decayed whereas the remaining wood appeared only discolored, we placed spore suspensions of F onto microtome sections freshly cut from several Populus tremuloides branch stubs. Percentage germination (based on 200 spores/rePLICATE section) remained high on pith and adjacent annual rings. Greatest germination appeared to be directly on the pith itself, but the small size of the pith made it impossible to accurately separate germination on the pith from that on the adjacent annual ring. Germination was never greater than 2.5% on any other material, and usually was 0.0% on the surfaces of the stub and fresh sapwood. Germination usually was greatest on pith cut from the distal end of the branch stub, and least on pith cut from close to the zone of discoloration usually present in the bases of the stubs. This
suggests that the pith of branch stubs may be an avenue of entrance for *P. ignarius var. populinus*.

Influence of gamma radiation on vessel development and spore movement in American elm. C. O. N. Cardoso & M. O. Garraway (Ohio State Univ., Columbus). Hypocotyls of *Phaselus vulgaris* incubated with *F. solani f. phaseoli* (FSP) and of healthy plants were extracted with ethanol at various intervals. The phenol concentration (Folin Denison m/1000 equivalents) of extracts from diseased hypocotyls increased with time to a maximum of 1.65 mg/plant 15 days after infection. Three percent of the total radioactivity of the petroleum ether fraction (PEF) of the extracts reached 25.5 μg/plant at 12 days and remained constant thereafter. Healthy hypocotyls contained no detectable PEF phenols. Growth of *F. solani f. cucurbitae* (FSC), but not of FSP, was inhibited by PDA containing 5.8 μg/ml PEF phenols. Similarly, spore germination and germ tube elongation of FSC, but not of FSP, were inhibited in water agar containing PEF. Separation of PEF on silica-gel plates with pentane-ether-acetic acid revealed that two of the phenols produced red colors with FeCl₃. The one at Rf 0.35 differed from phenol red by having a UV absorptions maximum at only 280 nm. The other, with Rf 0.65, has a UV spectrum similar to phaseolin (max at 315 and 280, 315:280 ratio of 0.237). Both phenols selectively inhibited the nonpathogenic FSC. Data suggest that at least two phenolic antimicrobial substances, probably phytotoxins, accumulate in FSP-infected bean hypocotyls.

Environmental contamination by airborne fluorides in Montana. C. E. Carlson & J. E. Dewey (U.S. Forest Serv., Missoula, Mt.). Gaseous and particulate fluoride emissions from an aluminum reduction plant in western Montana have caused considerable environmental damage over a large geographical area. Herbaceous plants, shrubs, and trees showed foliar burn correlated with excessive fluoride accumulations. Conifer needles showed definite histological responses to fluoride levels, including proplastidic and nuclear hypertrophy of phloem parenchyma. High levels of fluoride were found in cambial feeding insects, indicating that fluorides may be translocated within the vascular system of conifers. The occurrence of elevated fluoride levels was indicated throughout the food chain of the ecosystem. Foliar necrosis due to fluorides was found in Glacier National Park, representing an unwanted intrusion by technology of man into one of the few remaining truly pristine habitats of the world.

Induced resistance in alfalfa to *Corynebacterium insidiosum* by prior treatment with avirulent cells. R. B. Carrol & F. L. Lukezic (Pa. State Univ., Univ. Park). Resistance to virulent isolates of *Corynebacterium insidio-

A soluble antigen related to infection with apple chlorotic leaf spot virus. R. Chavez & R. M. Light (Purdue Univ., Lafayette, Ind.). Virus-specific antigen reacting in ring tests to titers of 1/32-1/128 by using monoclonal antibody for apple chlorotic leaf spot virus (CLSV) purified from *Chloropodium quinoa* by a method previously shown to give highly purified virus preparations. Used in ring tests at dilutions of up to 1/64, these antiserum also detected a virus-related soluble antigen in supernatant fluids from infected *C. quinoa* extracts subjected to prolonged ultracentrifugation (15 hr at 27,000 g in a Spincido No. 30 rotor). In gel-diffusion tests, this soluble antigen was detected in crude extracts from infected *C. quinoa*. Furthermore, gel diffusion tests showed that the antigen was also present in virus preparations, but was distinguishable from another virus-specific antigen that was not present in preparations of the soluble antigen. The soluble antigen was concentrated by precipitating in 50% ammonium sulfate solution and resuspending in buffer. It remained antigenic after storage at 4°C for 1 week and 3°C for at least 2 weeks. In gel diffusion and ring tests, no such antigen was detected in crude extracts from healthy *C. quinoa*, or in control preparations concentrated from healthy *C. quinoa* or from *C. quinoa* plants infected with several other viruses.

Germination of uredospores of *Puccinia recidivans* inhibited by blue, red, and far-red light. Ho-Shi Chiang & L. Calfoutos (Univ. Minn., St. Paul). A crude action spectrum was obtained for light inhibition of uredospore ger-
mination in the wheat leaf-rust pathogen (race UN-2). The illumination system consisted of a slide projector, water-cooling bath, interference filters (half-band width, 4 or 10 mm), and a sharp cut-off filter when testing the red and far-red wavelengths. Spores were exposed to 100% relative humidity overnight, then placed on water agar and illuminated for 2 hr at 8,000 erg/cm² per sec, and 100 spores were observed for germination. The agar plate was kept at 20-21 C. Inhibition of 97% or greater (as compared to germination of spores in darkness) occurred at wavelengths of 400, 419, 651, 699, 710, 720, and 750 nm. Moderate inhibition occurred at 390, 425, and 603 nm. Little or no inhibition occurred at 452, 493, and 552 nm. A response to far-red radiation is unusual in fungi.

Graminella nigrifrons as a vector of corn stunt agent. M. M. Choudhury & E. Rosenkrantz (Miss. State Univ., ARS, USDA, State College, Miss.). Graminella nigrifrons was established as a vector of both the Ohio corn stunt agent (CSA-OH) and the Mississippi corn stunt agent (CSA-MS). CSA-OH-infected CSA-MS was inoculated with CSA-MS by the more efficient vector, Dalbulbus maidis. The latent period of CSA-MS in G. nigrifrons was 15-18 days. The shortest incubation periods for CSA-OH and CSA-MS in corn infected by viruliferous G. nigrifrons were 11 and 15 days, respectively. After a 14 day access period, no infected plants were observed. The efficiency of transmission of CSA-OH by G. nigrifrons was 34%, that of CSA-MS, 3%. Female leafhoppers appeared to transmit CSA-OH more efficiently than their male counterparts. Some individual G. nigrifrons were found to transmit both CSA-OH and CSA-MS to the same test plants after feeding on source plants that exhibited symptoms of only Ohio corn stunt (CS-OH) or only Mississippi corn stunt (CS-MS). Such doubly infected test plants consistently developed symptoms first of CS-OH and then of CS-MS. These studies also revealed the natural occurrence in field corn of both CSA-OH and CSA-MS at State College, Miss. Moreover, noninfected G. nigrifrons acquired both CSA-MS and CSA-OH upon feeding on the same naturally infected corn plants.

Influence of fertilizer treatments and cropping sequence on populations of spiral nematodes. R. J. Collins & R. Rodriguez-Kabana (Auburn Univ., Auburn, Ala.). A 2-year study on the effect of fertilizer treatments on populations of spiral nematodes was made in plots within a 10-year-old fertility experiment. Plots studied were under the following sequence: winter wheat, corn, soybeans; fallow; cotton; and, in some plots, winter legume as green manure. Fertilizer treatments varied from a complete formulation (N, P, K, lime, minor elements) to treatments deficient in one or more components. Populations of spiral nematodes (predominantly Helicotylenchus dikystera) were highest in corn, followed by cotton and soybeans in decreasing order. Spiral nematodes occurred in high numbers only in soil and roots from plots receiving all major elements, lime, and a winter legume; the addition of minor elements did not affect spiral populations. In all crops, application of N in the form of a winter legume resulted in higher spiral populations than when inorganic N was used. With the exception of cotton, numbers of spiral nematodes were higher in plots deficient in P than those lacking K; difference between those treatments was not evident in cotton. Omission of lime restricted spiral populations in all crops.

Exchange of materials between corn tissues and Helminthosporium carbonum as determined by electron microscopy and conductivity techniques. J. C. Cosmetro & R. P. Schepfer (Mich. State Univ., E. Lansing). Many fungal spores lose nutrients when germination starts; external supplies are required. The microprobe was used to monitor content of elements in Helminthosporium carbonum (HC)-condina on corn leaves. Magnesium in condina at time of inoculation gave 1,200 cpm under our conditions. Conidia lost Mg (—30 to —50%) on susceptible leaves for 16-24 hr; the original Mg level was then regained by 36-48 hr. Conidia lost more Mg on resistant than on susceptible leaves, but no recovery was evident. Conidia containing Rb lost equal amounts (80%) to both resistant and susceptible Rb-free leaves. Conidia lost equal amounts of S on resistant and susceptible leaves during 36 hr. Helminthosporium victoriae (oat pathogen) conidia on corn lost Mg, Rb, and S in amounts comparable to the resistant corn. Susceptible leaves inoculated with Hc-condina (10-20/mm² leaf surface) lost more electrolytes (23 µmhos) than did control leaves (4-5 µmhos) by 24 hr. Such materials are available to the fungus. Resistant leaves had no increase in electrolyte loss 32 hr after inoculation (10 µmhos). Helminthosporium victoriae did not induce electrolyte loss from corn leaves. Exchanges between host and pathogen occur early, and differ in resistant and susceptible tissue.

Interaction of Verticillium albo-atrum and Pratylenchus penetrans under controlled inoculum densities. J. J. Conboy, R. J. Green, Jr., & J. M. Farris (Purdue Univ., Lafayette, Ind.). The influence of Pratylenchus penetrans on incidence of infection of Verticillium albo-atrum on tomato under controlled inoculum densities was considered. Inoculum densities by V. albo-atrum on tomato were established with controlled inoculum densities. Infection incidence of 100% occurred at the inoculum density of 200 propagules/g soil, and infection levels were progressively lower at inoculum densities of 100, 75, 50, and 25 propagules/g. Consistent increases in infection were found at all inoculum densities in the presence of the nematode. The number of nematodes extracted from roots of tomato plants infected with V. albo-atrum was significantly lower than the number extracted from roots with the nematode alone. The role of the nematode in increasing susceptibility of tomato to V. albo-atrum was considered, using a split root technique. Slightly higher levels of infection were found when the fungus and the nematode were placed on opposite sides of the split root system as compared to the fungus alone on one side, suggesting a host physiological response to the nematode which makes the plant more susceptible to infection by V. albo-atrum.

Influence of nutrition and total nonstructural carbohydrate content on Helminthosporium sativum-initiated leaf spot of Kentucky bluegrasses. H. B. Couch & L. D. Moore (Auburn State Tech. Inst., Auburn, Ala.). Thirteen Kentucky bluegrass cultivars were grown in expanded shale and supplied with three levels of nitrogen nutrition (N), respectively; standard Hoagland's solution (10 n) and Hoagland's solution modified to contain 0.1×N or 3×N. The plants were inoculated with an aqueous spore suspension of Helminthosporium sativum. Analyses were made of foliage collected prior to inoculation for total carbohydrate and fructose. Anheuser, Pelly, Pennstar, and Bellfert were moderately to highly resistant, and Delta, Merion, Fylkin, and Newport were moderately to highly susceptible at the three nitrogen levels. The highest levels of susceptibility occurred at the 3×N nutritional regime. Cougar, Winsor, Park, BA6124, and Kenblue were highly resistant only at the 0.1×N nutritional regime. There was no correlation between susceptibility and the concentration of either total and reducing carbohydrates or fructose. These studies indicate that while susceptibility of Kentucky bluegrasses to H. sativum is generally increased under high nitrogen nutrition, the total range of susceptibility of a given cultivar is more fully described when grown under a broad spectrum of nutritional regimes, and is independent of the total nonstructural carbohydrate content of the tissue.

Development of multiple disease-resistant fresh market varieties adapted for machine harvest. J. P. Crall, T. S. Byrnes, & R. W. Strode (Univ. Fla., Bradenton, Gainesville). In Florida it has been desirable to exploit
genetically controlled disease resistances, as the limiting factor in tomato (Lycopersicon esculentum) production is plant disease. Tomato pathogens in Florida controlled by the use of resistant or tolerant cultivars include races 1, 2, 3, 4, and 6 of Cladosporium fulvum (leaf mold), Alternaria solani (bacterial leaf blight), Verticillium albo-atrum (Verticillium wilt), Fusarium oxysporum lycopersici (races 1 and 2 of Fusarium wilt), Stemphylium solani (gray leaf spot), and five strains of tobacco mosaic virus. Resistance to fruit pox, gold fleck, blossom-end rot, crease-stem, graywall, and yellowtop diseases is also genetically controlled.

Genes for resistance to these diseases, along with the horticultural characters necessary for a machine-harvest freshmarket tomato, have been incorporated into a single genotype. Release of this variety should reduce production costs considerably and revolutionize the fresh-market tomato industry, as it can be harvested mechanically and possesses more disease resistance genes than does any previous cultivar. Estimated minimum number of disease resistance genes in this variety is 21. Genes for resistance to 12 more fungal, bacterial, viral, insect, and nematode pests have been identified and are being incorporated into this genotype.

The role of the basidial stage in the life cycle of Typhula idahoensis, B. M. Confer (Wash. State Univ., Pullman). Fertile spores of Typhula idahoensis, the major snow mold pathogen of winter wheat and barley in Washington, are usually found from late October until snowfall under field conditions. Sclerotia stored dry during the summer were placed outdoors at intervals during fall. Mature sporophores began to produce basidiospores in early November, and continued to produce spores until snowfall. Basidiospores were placed on 2% water agar on glass slides for 24, 56, and 72 hr at 1, 5, 10, 15, and 20°C in the dark. Germination occurred at all temperatures, but 10°C was most favorable for germ tube elongation. At 20°C, germination was lowest and germ tubes were short and abnormal. Winter wheat and barley were inoculated with basidiospores by suspending sporophores over plants in pots enclosed in polyethylene bags. Pots were kept at 5°C in the dark, or in a greenhouse at 10°C for 3-20 days. After inoculation, plants were covered with moist cotton and incubated at 1°C for at least 90 days. Typhula idahoensis was reisolated from about 8% of the inoculated plants. Disease symptoms and typical sclerotia developed on the plants. Failure to obtain a high level of infection may be due to the inoculation technique or to an inherent low-inoculum potential of basidiospores.

Mechanics of infection of soybean seedlings by overwintered and secondary Pseudomonas glycinea. G. C. Day & C. Lezen (Ohio Agr. Res. Dev. Cent., Wooster). Seeds were planted with abrasives (sand or Carborundum) and diseased leaf litter in growth room tests. Abrasives increased the number of cotyledon lesions on the emerged seedlings. In three tests, cotyledons bore 1.8-2.3 times more lesions than control cotyledons. Lesion area was also increased. Most lesions were on the proximal cotyledon end, which is usually the advancing end, as the cotyledon penetrates the soil. The proximal area most diseased was the "tough", the depression in which the radicle rests during dormancy. Apparently the trough was injured by abrasives caught by the upper radical before it broke through the cotyledon. Lesions appeared rarely on uninfected leaves; none occurred on trifoliate leaves. Tests were made in which seedlings (1/5 pot) with lesioned cotyledons, uninfected, or first trifoliate leaves were subjected to simulated windstorm. New lesions were initiated on leaves above the organ discussed prior to the storm. These findings suggest that pre-emergence infection of cotyledons is the initial event in the increase of overwintered inoculum, and that secondary infection on seedling leaves occurs during wind-spring storms.

Germination of Penicillium digitatum spores as affected by solutions of volatile components of citrus fruits. P. L. Davis & J. J. Smoot (ARS, USDA, Orlando, Fla.). Volatile compounds known to occur in mature citrus fruits were evaluated for their effects on spore germination of Penicillium digitatum, a major citrus fruit-rotting fungus. The compounds tested were obtained from commercial sources and included aldehydes, alcohols, esters, and terpenes. Spores were grown on 2-week-old culture dishes in liquid synthetic medium at about pH 4 containing solutions of the test materials. The aldehydes inhibited germination, whereas the esters, terpenes, and alcohols had no effect on germination. In quantitative tests with short-chain aldehydes, the inhibitory effect increased with chain length from C3 to C10. The minimum concentrations in milliliter producing 50% or more inhibition were 1.0 mg for valeric aldehyde, 0.2 mg for hexanal, 0.1 mg for heptanal, and 0.06 mg for octanal.

Surface stain on sapwood of partially wetted longleaf pine logs. R. C. De Groof & F. J. Cazayoux (USDA Forest Serv., Gulfport, Miss.). A continuous spray of water over logs in storage can protect wood from decay and staining fungi. The determination of the proper volume of water needed is critical to the economical and efficient use of spraying systems. Three patterns of biologic activity were observed under the bark of longleaf pine logs that were partially immersed under a continuous spray of water for 6 weeks. In portions of logs in which the bark was continuous, reddish-orange stain developed on the surface of the wood. The part of the log which remained dry except for occasional exposure to rain was rapidly colonized by dermatocystes, blue stain fungi, decay fungi, and bark-inhabiting insects, but the orange-red stain did not develop there. The red stain spread over the surface of the xylem tissues, but did not penetrate the sapwood. Several bacteria and yeasts were associated with the stain, but their relationship to the discoloration has not been defined. It is suggested that freshly cut longleaf pine logs can be used to monitor the effectiveness of water spraying systems.

Resistance to rust in red clover. S. Diachun & L. Hensson (Univ. Ky., Lexington). Rust, caused by Uromyces trifolii var. fallens, is common on red clover under natural conditions in late summer in Kentucky. Several hundred seedlings and detached leaves of several hundred plants were inoculated with suspensions of uredospores of an isolate of U. trifolii var. fallens from red clover. Inoculated leaves and plants were incubated 14 days in growth chambers. A rust-resistant plant in a population of Lake red clover was selected in 1965 and is maintained as a clone, designated KyCL75. In F1 populations from crosses between KyCL75 and several rust susceptible clones, resistant and susceptible plants occurred in approximately equal numbers. In F2 populations from crosses between susceptible clones, all plants were susceptible. These results indicate that resistance in KyCL75 is inherited as a single dominant factor, and that clone KyCL75 is heterozygous for this factor.

Evidence for possible genomic masking between two unrelated plant viruses. J. A. Doode & R. I. Hamilton (McGill Univ., Montreal, Quebec, Can.). Mixtures of purified barley stripe mosaic virus (BSMV) and tobacco mosaic virus (TMV) were prepared in two ways. The first involved purifying the nucleoprotein from barley plants doubly infected with the two viruses. The second involved mixing the two viruses in vitro so that the product resembled the mixture purified from the doubly infected plants. Both mixtures were repeatedly incubated with TMV antibody until no precipitate was obtained. The BSMV which remained from the natural mixture contained TMV infectivity, most of which was removed by incubation with BSMV antibody. The mixture which was prepared in vitro contained a much smaller amount of TMV infectivity which was not removed by BSMV antibody, but which was removed by TMV antibody. The produc-
tion of genomically masked virus particles with the coat protein of BSMV and the nucleic acid of TMV in doubly infected plants would explain these results.

Implications of inoculum dispersal pattern, meteorological data, and serological comparisons to epidemiology of *Sclerotodes lagerbergii*. C. D. Dvorak & R. B. Mattheson (Can. Forestry Serv., Ontario Region Insect Pathol. Res. Inst., Sault Ste. Marie, Ontario). *Sclerotodes lagerbergii* is extremely damaging to pine on cool moist sites in Europe, and in plantations on hot sandy plains in central Ontario and adjacent areas east and west. Epiphytotics in North America intensify most rapidly in geographical depressions and on plains, this effect usually being attributed to pre-positioning of the suscepts to infection by frost damage. Temperature differences in a red pine plantation, however, were too slight to account for a 100% reduction in survival in the depressions as compared with adjacent areas. Depressions received 1 to 4 times as much inoculum as adjacent areas, depending upon their capability to serve as air drainage troughs or basins. Serological comparisons among five isolates of *S. lagerbergii* from central and northern Ontario indicated that all were closely related and four were identical. *Sclerotodes lagerbergii* in Ontario is interpreted as a relatively uniform clone which escaped from the basin area of Lake Superior, where it was largely contained in small areas where conditions were optimum for infection, and subsequent local spread was minimal. It is now as far north as Lat. 52 degrees, where cool moist weather and continuous high-suscept density exist, and there the pattern of the epiphytotic should more closely resemble that described from Europe.

Toxic Fusaria isolated from moldy sweet potatoes involved in an episodic of atypical systemic necrosis in cattle. B. D. Drucker, Jr., O. H. Jones, Jr., & J. C. Peckratis (University of Georgia, Coastal Plain Exp. Sta. and Diagnostic and Research Lab., Tifton). Several isolates of *Fusarium moniliforme*, *F. roseum*, and *F. solani* were obtained from moldy sweet potatoes involved in an episodic of atypical systemic necrosis (AIP) and screened for toxicity to chicks. The isolates were separately grown on moist, autoclaved corn for 2 weeks, dried, ground, and mixed with protein supplement (6:4 w/w), then fed to day-old Bobwhite cockerels for 2 weeks. Of 15 isolates screened, one of *F. moniliforme* and two of *F. solani* caused weight reduction of 50% or more compared to control chicks and one *F. solani* isolate was lethal. Clinical signs and lesions of AIP were reproduced in a steer and a cow by giving orally a homogenate of viable sweet potatoes which had been sliced, inoculated with the lethal isolate of *F. solani* and incubated for 10 days. An ether extract of a similar homogenate produced AIP when given orally to a cow. Although this disease has been known for over 40 years, this is the first reported success of its reproduction in cattle under controlled conditions using a known organism.

The periderm, a factor limiting apple scab development. C. R. Drake & G. M. Shear (Va. Polytechnic Institute, Va. State Univ., Blacksburg). A periderm was found to be associated with apple scab (Venturia inaequalis) infection of apple fruit, apple pedicels, and leaf petioles, but not leaf lamina. The periderm usually was initiated in a row of cells immediately beneath the epidermis, where infection had occurred. Periderm development was more extensive in fruit and leaf pedicels than in leaf petioles. The formation of the periderm seemed to be related to the ranidity of hyphal proliferation. During early stages of infection and when the environment was favorable for rapid hyphal mat formation between the epidermal cells and the cuticle, the periderm was initiated, but there was only a minor change in the host tissue. The most extensively developed periderms were associated with the older areas of infection. In the older infections, cell division in the periderm frequently continued until 12 to 14 rows of additional host cells were produced. In the case of fruit pedicels, the extra rows of phellem cells to the outside frequently turned the pedicel 15 to 30 degrees from its long axis, resulting in a deformed, weakened structure. The epidermis of the suberized phellem cells of the periderm inhibited radial movement of the pathogen, but had no effect on lateral proliferation of the advancing hypha.

Ultrastructure of soybean leaves affected by systemic toxemia. J. Duex & R. J. Zeeck (Univ. Minn., St. Paul). Systemic toxemia in young trifoliate leaves of soybean (*Glycine max*) is caused by toxin(s) produced in older leaves infected by the bacterial pathogen, *Pseudomonas glycinea*. Affected leaves can recover from symptoms. Ultrastructural changes in leaf mesophyll affected by toxemia, and of chlorotic areas adjacent to lesions, were observed in young leaves 8 days after inoculation. Recovered and comparable healthy tissue were sampled 18 days after inoculation. Mesophyll cells in young healthy tissue had large vacuoles and chloroplasts with well-developed grana and abundant starch grains. Systemically affected leaves had small and thin cells with dense cytoplasm, small vacuoles, and chloroplasts with few lamellae and little starch accumulation. Cells of recovered leaves had large vacuoles, normal chloroplasts with well-developed grana, and extreme starch accumulation similar to comparable healthy cells. The toxin(s) did not cause a lesion of cell organelles, but delayed mesophyll cell maturation.

Water relations of tomato plants infected with *Fusarium oxysporum* f. sp. lycopersici. J. M. Dunaway (Univ. Cal., Davis). Determinations of leaf water and solute potentials showed that wilting of infected plants was due to water stress. When the diffusive resistance of leaves to water vapor loss was measured as a function of decreasing leaf water potential, the diffusive resistance in infected plants was as high as, or higher than, the resistance in healthy plants at a given leaf water potential. A leaf disc experiment showed that the disease caused an inhibition of stomatal opening which was independent of disease effects on water potential. Measurements of water flow through excised root systems indicated that infection did not increase the resistance of roots to water flow. When water was forced through stem segments, the resistance of infected segments decreased markedly as water flow occurred, and accurate estimation of xylem resistance in infected plants was impossible by this technique. Steady-state measurements of transpiration and leaf water potential indicated that the resistance to water flow in noninfected plants approached infinity as wilting occurred, and it was concluded that a high resistance to water flow in xylem is the sole cause of the wilting which characterizes the *Fusarium*-wilt disease of tomato.

Survival of *Corynebacterium michiganense* in soil as free cells and in infected tomato tissue. E. Eichland (N. C. State Univ., Raleigh). *Corynebacterium michiganense* could not be recovered by planting tomato seedlings in soil collected from around the base of diseased tomato plants at the end of the growing season. However, tomato seedlings transplanted immediately into soil artificially infected with 2 × 10⁷ cells/g of soil developed canker, while those transplanted 2 weeks later remained healthy. Moreover, *C. michiganense* was not recovered from centrifuged suspensions of soil artificially infected with 2 × 10⁷ cells/g after 1 week at 25°C, nor from dialyzing bags containing the organism and buried in the soil under the same conditions. However, *C. michiganense* was recovered from naturally infected tomato stems, roots, leaves, fruit skins, and seeds which overwintered on the soil surface, and from undecomposed woody tissue transplanted in the soil. Tomato seedlings transplanted immediately into soil infected with surface plant debris became infected, but seedlings transplanted 2 weeks later remained healthy. When small pieces (ca. 0.5 × 10 mm) of infected, overwintered woody tissue from buried plants were placed on healthy tomato leaflets, 85% of the inoculated leaflets became infected. *Corynebacterium*
michiganense was short-lived as free cells in the soil, and only survived overwintering in undecomposed plant tissues.

Evaluation of imines of sec-butylamine for control of Penicillium decay of oranges, J. W. Eckert, M. L. Rahm, & R. J. Stewart (Univ. Calif., Riverside). Imines synthesized by reaction of sec-butylamine (SBA) with 2-penta-
none, 2-heptanone, 2-undecanone, benzaldehyde, furfural, and vanillin, respectively, were evaluated as vapor-phase treatments to prevent decay of Valencia oranges. At doses equivalent to 0.1 g SBA/m³ air, the imines were more effective than SBA, but the fruit were imines and the imines were in contact with branch bark. Benzaldehyde, furfural, or their imines. The imines of 2-
pentanone and 2-undecanone (aliphatic imines) hydrolyzed rapidly in the gaseous state in moist air, whereas the imines of furfural and benzaldehyde (aromatic imines) were stable under these conditions. The latter imines hydrolyzed in moist paper, and the rate of hydrolysis was greater below pH 7. Hydrolysis of the aromatic imines increased the pH of unbuffered paper from 5 to 9.1 within 2 hr, and 80-
100% of the SBA from the imines was recovered from the atmosphere over a 5-day period. The aldehydes and ketones at concentrations equivalent to the imines had no effect upon fruit decay. The superiority of the aliphatic imines over SBA in reducing decay must be due to the sustained evolution of gaseous SBA, but for the aromatic imines, the relative contribution of imine and SBA to disease control is difficult to assess.

The association of Fusarium moniliforme with the crown rot complexes of asparagus, R. M. Endo & E. C. Burke (Univ. Calif., Riverside). Crown rot of asparagus plants in southern California are frequently unproductive and short-lived. Inland plantings and fields cut for 3 or 4 months are more severely affected than coastal plantings and fields cut for 1 or 2 months. Vegetative stems of affected plants decrease in number and size; reproductive stalks are yellow, stunted, and reduced in vigor; and fern stalks die in various stages of elongation without vascular discoloration. A dry, brown crown rot was observed on all plants in the field. Severity of crown rot appeared inversely related to the vigor and size of affected plants. Fusarium moniliforme was isolated from 93% of the crowns, F. oxysporum from 22%, and Penicillium marneffii from 6%. Out of 80 single-
spore cultures of F. moniliforme isolated from affected crowns 5 counties were pathogenic to Fusarium wilt-resistant asparagus seedlings, UC 72 and 66; reddish-brown lesions were visible on the crown base, stems, leaves, and feeder roots. Removal of a portion of the stems or storage roots significantly increased the severity of crown rot. The fungus was also isolated from commercial lots of seeds, from asparagus debris, and from volunteer asparagus seedlings. Fusarium moniliforme is considered an important pathogenic component of the crown rot complex.

The pathogenicity of Helminthosporium maydis and Helminthosporium spp. on carnation, snapdragon, and geranium, A. W. Engelsman (Univ. Fla., Bradenton). Helmin-
thosporium maydis race T and race O from corn, and Helminthosporium spp. isolates obtained from a petal and a leaf of Chrysanthemum morifolium and from a leaf of Maranta leuconeura, were pathogenic on geranium leaves and on the flowers of carnation, snapdragon, and geranium but not on the flowers of the azalea, Rhododendron 'Erie'. Infection on the flowers of each host was evident after incubation for 24 hr at high humidity. All five isolates induced similar symptoms on the same host. Tan necrotic spots up to 2 mm in diameter developed on snapdragon flowers within 10 days, and areas of the spadix turned brown. General collapse of the snapdragon flowers occurred within 3 days. The symptoms were similar to those induced by Botrytis cinerea in the same experiment. Irregular, tann-colored lesions, varying from the size of a pin point to 4 mm in length, developed on the petals of carnation flowers. Initially, lesions 1-2 mm in diameter developed on geranium flowers, followed rapidly by necrosis of large areas of the petal tissue. Irregularly shaped, sunken lesions up to 8 mm in diameter, with smooth gray centers, developed on geranium leaves.

Stalk rot in corn influenced by Texas male sterile cyto-
plasm. A. H. Epstein, J. E. Reynolds, & W. P. Dietz (Iowa State Univ., Ames). Isogenic lines of hybrid corn (AG19 X AG32) containing normal (N) cytoplasm and Texas male sterile (Tms) cytoplasm were compared for incidence and severity of foliar disease and stalk rot. Stalk rot severity was measured in terms of stalk-breaking strength, or "moment". This value was determined by exerting a lateral force at the fifth node sufficient to break the stalk, and measuring the distance from the 5th to the 6th node to the break. The product of the force in kg by the distance in cm is the "stalk-breaking moment". Foliar diseases such as yellow leaf blight (Phlyosphaera zeae) and southern corn leaf blight (Helminthosporium maydis race T) were more severe on plants with Tms cytoplasm. The first five leaves of Tms plants were dead due to Phyllosticta sp. in-
fec tion by 18 June; little further development occurred. Helminthosporium maydis infections were numerous on all leaves of Tms plants by 26 August; plants appeared dead by 1 September. Stalk rot in terms of stalk breaking moment was more severe in Tms plants. The average breaking moment was 102.5 for Tms plants and 278.2 for N plants.

Vertical spore concentrations of three wheat pathogens above a wheat field. M. G. Eversmeyer, C. L. Kramer, & J. R. Burleigh (Kans, State Univ., ARS, USDA, Manhattan). Kramer-Collins samplers were used to measure concentrations of Erysiphe graminis conidia and Puccinia graminis and P. recondita, spores in the wheat canopy, 1, 3, and 6 m above ground level. Although considerable daily and hourly variations were observed in spore concentrations, epidemic patterns with distinct peaks occurred. Time of maximum spore concentrations varied with meteorological factors present during a particular hour or day. Changes in wind velocity or turbulence accounted for most of the peaks observed in hourly spore concentrations. Higher wind velocities were required to dislodge spores from wet plants than from dry plants. Higher wind velocities were required to disperse spores over to leaves and stems in the lower 3 m in spore concentrations generally occurred after plant surfaces dried in midmorning; they occurred later on days when plant surfaces remained wet longer. Peaks in spore concentration occurred as wind velocities and turbulence increased prior to a rain shower or passage of a cold front, and not as a result of the impact of rain drops on spore-bearing tissue. Profiles of spore concentrations at different heights distinguished between endogenous, exogenous, and mixed inoculum sources.

A selective medium for assay of Colletotrichum coccodes in soil. J. D. Farley (Ohio Agr. Res. Devol, Cen., Wooster). A selective medium was developed which successfully assayed Colletotrichum coccodes in nonsterilized soil artificially infected with conidia or sclerotia at populations as low as 100 propagules/g. The medium consisted of the following ingredients/liter: 10 g polygalacturonate acid; 1.5 g KH₂PO₄; 4 g K₂HPO₄; 25 ml soil extract; 17 g Dical agar; and the following antimicrobial agents added after autoclaving: 0.1 g pentaachlorothiophene; 0.1 g benomyl; 50 mg streptomycin sulfate; and 50 mg chloramphenicol. The pH was adjusted to 5.0 prior to auto-

claving, and plates were incubated at 26 C. Colletotrichum coccodes colonies produced abundant sclerotia and brown mycelium; the medium should be aerated. Controls cannot be differentiated from other fungi when examined on a white background. In addition to restricting spread and development of other fungi on the medium, PCNB also enhanced sclerotial production of C. coccodes. The medium was used to isolate C. coccodes from overwintered tomato fruit, and shows promise of direct isolation of the pathogen from field soil.
Antibiotic production by Pseudomonas morsprunorum. D. M. Filienger & D. Powell (Univ. Ill., Urbana). Pseudomonas morsprunorum, a pathogen of stone fruit trees, produces an antibiotic in peptone-containing liquid or solid media. This constitutes the first report of antibiotic activity by this bacterium. The antibiotic is readily produced in King A & King B media, but not in asparagin-dextrose agar nutrient agar, and synthetic media. The production of the antibiotic is greatly influenced by peptone source, amount, and lot. Modified King B medium (modifications: no agar, pH 7.0, and 12 g/liter of Neopeptone (Difco)) gives excellent results. The antibiotic is soluble in ethyl alcohol, benzene, petroleum ether, ethyl acetate, and chloroform. Part of this information is based on results using crude solutions or lyophilized extracts. The antibiotic appears to be primarily antifungal. Geotrichum candidum is very sensitive to it.

Possible xylem translocation of pea streak virus. R. E. Ford & S. W. Rosso (Iowa State Univ., Ames). Pea streak virus (PSV), Oregon isolate, produces in peas necrotic local lesions, necrosis of veins and veins, and savoy of youngest leaves. Infected Perfected Genes peas were fixed in FAA, dehydrated in acetone, and embedded in Epon, sectioned, stained with uranyl acetate and lead citrate, and observed with a Hitachi HU 11C electron microscope. Large aggregates of rod-shaped particles were found in parenchyma cells. These aggregates were also found in immature vessels that were developing secondary wall thickenings. Particles of PSV were seen after cytoplasmic contents had disintegrated. This evidence suggests that PSV is quite stable in vivo and that PSV may move via the xylem or the phloem.

Rhiococnia blight of water hyacinth. T. E. Freeman & F. W. Zettler (Univ. Fla., Gainesville). During 1970, surveys were conducted in search of phytopathogens exhibiting potential for biological control of noxious aquatic weeds. During one of these surveys in the Canals Zone of Panama, a blight of anchoring hyacinth (Eichhornia azurea) incited by Rhizocconia solani was noted along the Rio Chagres. An isolate of the fungus from this source proved to be extremely pathogenic on water hyacinth (E. crassipes). Inoculation of water hyacinth with mycelial fragments of the fungus caused the mortality of the plant which frequently resulted in death of the entire plant. Disease development was favored by temperatures of 22-27°C with a marked decrease in severity at 32°C. This decrease was not correlated either with growth of the parasite in vitro or with its pathogenicity on cucumber. Symptoms incited by this fungus and its host range were similar to that of R. solani isolated from kenaf in Florida; however, the Panama isolate was more pathogenic on water hyacinth. Sclerotia of this fungus maintained their viability without loss of virulence after being submerged for 9 months in lake water.

Germination of ureidoepoxides of wheat stem rust pre-treated with volatile stimulators. R. C. French & M. D. Balamore (Plant Sci. Lab., Ft. Detrick, Frederick, Md.). Many diverse types of chemicals, including nonanol isolated from ureidoepoxides, have previously been reported to stimulate the germination of ureidoepoxides floated on water. Pretreatment in Conway diffusion cells of dry spores with high concentrations of supporting vapors of polar stimulators (10 µl per liter n-nonanol or n-nonanol/4.0 ml water in the annulus) in the presence of water vapor, reduced germination to zero and spore damage was apparent. In the absence of water vapor, pretreatment with stimulator vapor had no effect or pretreatment with supporting vapors of nonanol (0.1 or 0.01 µl nonanol/4.0 ml water) stimulated germination above that of the controls. Nonane, one of several nonpolar hydrocarbon stimulators, neither reduced germination nor damaged the spores at the rate of 10 µl/4.0 ml water. Nonanol at 0.5 to 10 µl/4.0 ml water stimulated germination when applied to spores floating on water. When the same concentrations were applied as 2-h pretreatments to dry spores, subsequent germination on water was virtually zero.

Stimulation of Phytophthora cactorum by fatty acids and metabolism of oleic acid-1-14C. R. E. Gain, F. L. Brown, & C. D. Briscoe (Marshall Univ., Huntington, W. Va., & Va. A&M Univ., Morgantown). Oleic, linoleic, and linolenic acids (5 mg/liter) stimulated growth of Phytophthora cactorum when added individually to a basal glucose-asparagine medium. Five days' incubation with oleic acid in 25 ml medium at 20°C yielded a 5- to 7-fold increase in dry weight of the controls. Similar increases were observed with linoleic and linolenic acids. Saturated fatty acids such as lauric, myristic, palmitic, and stearic were also stimulatory, but to a lesser degree. When cholesterol (10 mg/liter) was added to the basal medium, the individual saturated fatty acids caused a slight growth increase over the cholesterol controls, whereas the unsaturated fatty acids resulted in an approximate doubling of the dry weight after 5 days. This additive stimulation of growth by cholesterol and the unsaturated fatty acids lessened with incubation time until maximum dry weights of test cultures were not much larger than the cholesterol controls after 10 to 12 days. Phytol was incubated with oleic acid-1-14C (2 µc, 5 mg/liter) in a basal medium without cholesterol for 8 days in flasks equipped with CO2 traps, then extracted with chloroform and analyzed for transformation products. Less than 5% of the oleic acid was metabolized to CO2, ca. 27% of the recovered activity was unchanged oleic acid, and ca. 70% was esterified to unidentified compounds.

Changes in soil and the death of woody ornamentals associated with leaking natural gas. J. H. Garner (N.C. State Univ., Raleigh). This study was made to identify changes in the biotic and mineral composition of the soil associated with leakage of natural gas and the deterioration and death of trees and woody ornamentals. In soil contaminated with natural gas, (i) the soil atmosphere was anaerobic and large numbers of hydrocarbon-utilizing and sulfur-reducing bacteria were present; (ii) the pH range was 5.5-8.5; (iii) there was an increase in manganese and organic matter; (iv) sulfides of hydrogen and formic acid and the associated 'sour-gas' smell were present; (v) there was a decrease in numbers of actinomycetes, and fungi when compared to control soils. The changes indicated by the study are consistent with those noted by petroleum microbiologists when studying the effects of leaking natural gas in the soil. The changes occur due to the adaptation of the microbial ecosystem to a new environment. Sulfides and excesses of manganese are detrimental to plant growth.

Chemical evidence that the short particle of tobacco rattle virus species coat protein. S. A. Ghahrial & R. M. Lister (Purdue Univ., Lafayette, Ind.). Biological evidence strongly suggests that the essential genome of tobacco rattle virus (TRV) is divided between the ribonucleic acids of "long" and "short" tubular nucleoprotein particles, which have been termed short- and long-particle RNAs. Short-particle RNA replication, and serological and surface charge studies suggest that the coat protein is specified by short-particle RNA. More rigorous chemical evidence has now been obtained to confirm this conclusion. Amino acid analyses were performed on the protein component of two biologically distinctive strains of TRV, TRV-Y and TRV-Z, and also on the protein of an artificial hybrid, "TRV-Y/Z", produced in mixed infections using long particles of TRV-Z and short particles of TRV-Y. Protein preparations were obtained by dialyzing purified virus from 0.04 M calcium chloride. Results showed that the protein of TRV-Y and that of TRV-Z differed markedly in amino acid
composition, especially in the content of alanine, aspartic acid, glutamic acid, glycine, and serine. The amino acid composition of the hybrid TRV-V/Y was the same as that of TRV-Y, the strain from which the short particles were used in all cases. In the work, the protein subunit had a molar weight of ca. 20,000.

A single high-rate application of Difolatan was taken for the control of apple scab, J. D. GILPATRICK, M. S. ZUSKIN, & S. D. G fuss (N.Y. State Agr. Exp. Sta., Geneva). A single sprayer of Difolatan 4 P (capitol) at the high rate of 5 quarts/100 gal (350 gal/acre), to large, bearing trees at the green tip stage of leaf development, gave excellent control of apple scab beyond petal fall. This spray replaced and gave a better performance than 5 weekly precrop sprays of captan SOWP, 2 lb/100 gal. Control throughout the rest of the season was obtained when Difolatan was followed by cover sprays of captan; but in 1970 these were not required. Gas-spectrophotometric analyses of residues on the surface of trees sprayed with Difolatan in 1969 and 1970 revealed levels of 500-700 ppm on the bark of branches until after bloom, declining to 3-40 ppm by harvest time. Residues of 300-700 ppm were found on clusters, 60-100 ppm on flowers, and 35-75 ppm on terminal leaves during bloom, even though only partially exposed, if at all, to Difolatan at the time of application. Much lower residues were found on mature green fruit and none on ripe fruit. The success of this promising new method of scab control is due mostly to the persistence of Difolatan on the trees and its redistribution by rains so that sufficient protective levels of the chemical are present on susceptible tissues during bloom, a critical stage for infections.

Ribonucleic acid polymerase from Rhizopus stolonifer. C. GONG & J. L. VAN ETTER (Univ. of Nebraska, Lincoln). Extracts of germinated spores and filaments of Rhizopus stolonifer were treated with RNA polymerase and a diethylaminoethyl cellulose column. Similar preparations from ungerminated spores contained only two active fractions corresponding to fractions I and III. Fraction III from both spore states exhibited identical characteristics, whereas fraction I from dormant spores required conditions for optimal activity slightly different from those for fraction I of germinated spores. All fractions required four ribonucleoside triphosphates, a divalent cation, and template DNA for activity, and were inhibited by actinomycin D but not by rifamycin. The individual fractions differed in the optimal Mn++ or Mg++ ion concentrations. However, optimal activities were obtained with Mn++ and denatured DNA for all fractions.

Re-evaluation of the role of NH₃ as the cause of the hypersensitive reaction. R. N. GOODMAN (Univ. of Missouri, Columbia). Contrary to published reports from our laboratory, recent experiments have disclosed that NH₃ accumulation and increased pH in tissues undergoing biologically induced (10⁸ cells/ml) hypersensitive reaction (HR) are not causally related to the resultant permeability changes and membrane damage. In previously published experiments, we studied HR induced by incompatible bacteria (Pseudomonas chlororaphis and Pseudomonas viridiflava) and it was observed that the cells were kept in moist chambers. In this system, desiccation of tissue and other symptoms of HR were delayed more than 48 hr, which provided more time to study the otherwise rapid death of inoculated tissue. Data are presented from experiments with the same system that required less than a 10-fold increase in electron micrographs disclose dense membrane damage 4 hr after inoculation. These occur 18 hr before the observation of either a pH rise or an accumulation of NH₃.

Transpiration of bur oak during oak wilt pathology. G. F. GREGORY (USDA Forest Serv., Delaware, Ohio). Transpiration (in a growth chamber) was determined for pairs of 2-year-old bur oak ramets after inoculation of one member of each pair with Ceratocystis fagacearum. The other as a control. All plants were weighed daily to determine transpiration loss. These pairs of ramets were harvested periodically, and the two members were compared for flow rate of sterile water through the stem sections of each plant, water content of leaves and stem (by fresh and dry weight measurements), and ash content of the leaves and stem. Transpiration held steady, then declined rather rapidly at varying time intervals after inoculation. The period from inoculation to start of transpiration decline was dependent on the period from inoculation to first leaf symptoms. When averaged for plants showing this decline, this was 57%. Flow rate of water through stem sections was nearly always markedly reduced by the time leaf symptoms appeared. Symptomless inoculated plants that showed transpiration decline also had reduced water flow capacity. Water content of the leaves of symptomless plants was not significantly different from their corresponding noninoculated ramets, whereas that of plants with leaves showing symptoms was reduced. No significant difference occurred in ash content.

Induction and photoconversion of peroxidase in leaves of Phaseolus vulgaris 'Pinto III'. H. HANEB & C. R. CURTIS (Univ. of Maine, College Park). Detached Pinto bean leaves 9-10 days old were placed in a moist chamber 6 inches from two germicidal UV radiators (G15T8, 2537A), irradiated for 0, 15, 30, 60, and 120 min, then incubated in darkness at 25 °C for 24 hr. Two leaf discs (12-mm diam) from each of four leaves were homogenized in 10 ml of cold 0.1 M phosphate buffer, pH 5.8. The homogenate was filtered and the filtrate centrifuged for 30 min at 30,000 g at 2-4 °C. The supernatant was assayed for peroxidase activity using a spectrophotometric guaiacol-H₂O₂ method. Peroxidase activity was increased in extracts from all far UV (2537A)-irradiated leaves. The peroxidase stimulation was confined only to the irradiated areas, and did not occur in nonirradiated half-leaves or other unexposed portions of leaves. Peroxidase stimulation due to far UV irradiation of leaves was reversed by subsequent exposure to near UV (GE 2537A-FLL, principal wavelength 3650 Å); the degree of reversal was related to the length of exposure to near UV irradiation. Near UV irradiation alone failed to stimulate peroxidase activity. Since considerable increases in oxidative enzymes are known to occur during infection in certain plant diseases, these results indicate a means of changing peroxidase activity in host cells without infection.

The nature of polypeptide, polyanine, and protein molecules which promote phytoalexin production. L. A. HAUWIESE (Wash. State Univ., Pullman). Many components released by plant pathogens and other microorganisms stimulate phytoalexin production in plants. Some of these components are basic polypeptides. The synthetic polypeptide poly-L-lysine (1 mg/ml) enhanced phenylalanine ammonia lyase (PAL) activity > 6-fold, and initiated the de novo synthesis of > 100 μg/g per 24 hr in pea pod tissue within 24 hr after application. Synthetic polypeptides (containing lysine, ornithine, or arginine) effective in inducing PAL and pisatin contained 14 to 100 amino acid residues. Optimal induction was obtained with peptides containing 14-50 residues, which will account for less than four or more than 700 residues and 20 other synthetic polypeptides not containing basic amino acids were ineffective as PAL inducers. Sporine, putrescine, cadaverine, protamine, and several other basic polypeptides also induced PAL and pisatin formation in peas. These and other phytoalexin-inducing compounds have in common an affinity for DNA.

Some characteristics of pectic hydrolases produced by Pseudomonas fluorescens, a causal agent of "pink eye" disease of potato tubers. S. S. HAGAR & G. A. McINTYRE (Univ. of Maine, Orono). Potato tuber slices were inoculated with Pseudomonas fluorescens. After 120 hr incubation at 20 °C, the tissue was extracted. The extract was centrifuged, filtered,
dialyzed, and assayed for pectic enzyme activity. No pectin methylesterase activity was observed in extracts of control plants. However, there was lyase activity. The extract was placed on a pH 9.4 DEAE-cellulose column and eluted by a gradient of pH 7.6 Tris buffer (0.01-0.1 M). Two pectic lyase systems (peaks 1 and 2) were eluted. Peak 1 was more active than peak 2. Optimum pH range of activity was 8.5-9.5. Activity of both peaks was enhanced by Ca2+ (0.001 M). Both peaks were inactivated when held above 50°C for 1 hr. Substrate preference was Na-polysaccharide > pectin N.F. > polygalacturonic acid. Thin-layer chromatography of hydrolytic products showed that unsaturated uronides and pectic acid fragments were present after 2-hr hydrolysis, but 96-hr hydrolysis, only unsaturated uronides were observed. Molecular weight determinations by Sephacryl G-200 filtration were ca. 18,000 (peak 1) and 22,500 (peak 2). Both enzyme systems appear to be endopeptidases.

Egg hatching of Meloidogyne incognita in the neutral carbohydrate fraction of root exudates of gnotobiotically grown Dupuits alfalfa. R. A. Hamlen, J. R. Bloom, & F. L. Lukesic (Pra State Univ., Unv. Park). Due to the implication by previous works of carbohydrates in egg hatching of Meloidogyne incognita, we studied egg-hatching success in soil sterilized by gamma irradiation and wetted with root exudates from plants in different stages of development and subjected to various levels of clipping. Carbohydrate components of the exudates were identified by thin-layer chromatography-mass spectrometry. Our results suggest the involvement of glucose with hatch; however, subsequent testing of glucose alone did not confirm this association. Percentage of hatch also was positively correlated with concentrations of inositol, sucrose, and three unidentified compounds in the exudates. Greater hatch occurred in exudates from seedling alfalfa than from mature, nonflowering plants, and more occurred in exudates from flowering plants than from nonflowering plants. Hatch was significantly higher in samples from lightly clipped alfalfa than from severely clipped plants. However, hatch in root exudates was never significantly different than in distilled water. The results indicate that the total neutral carbohydrate fraction and component carbohydrates considered in this study were not effective hatching stimulants of M. incognita under these experimental conditions.

Cell permeability changes in sunflower caused by infection by Sclerotinia sclerotiorum. J. G. Hancock (Univ. Cal., Berkeley). Influx and efflux of water (bulk flow), ions, and protons were determined by electrical measurements. Sunflower (Helianthus annuus) hypocotyl sections from above lesions caused by Sclerotinia sclerotiorum than for those from healthy plants. Urea uptake by sections from above lesions is reduced (squash, tomato) or unchanged (bean) in other hosts after Sclerotinia infection, and it is reduced in diseased sunflower tissues even during brief incubation periods. Efflux of urea from sunflower hypocotyl sections is biphasic, suggesting diffusion from two compartments (cytoplasm and vacuoles). Urea efflux from cytoplasm in diseased tissues (t1/2 = 59 min) is faster than from vacuoles of cells in healthy or diseased hypocotyl tissues (t1/2 = 250 min), but slower than that from cytoplasm of healthy ones (t1/2 = 58 min). Increased resistance to diffusion of urea in host cells above lesions apparently resides in the plasmalemma. In contrast to intact plants, electrolyte leakage is greater from tissues adjacent to lesions than from comparable noninoculated plant materials when excised sunflower shoots or celery petioles are used. Increased permeability stimulated by infection of excised organs is more reminiscent of senescent processes than disease or wound responses.

Toxicity of aqueous extracts of peas infected with Aspergillus ruber to healthy excised pea embryonic axes. G. J. Schliemann (N.Y. State Agr. Exp. Sta., Geneva). Aspergillus ruber (NRRL 52) is highly pathogenic to pea seed. Germination of surface-sterilized inoculated seed stored at 30°C and 87% relative humidity for 14 weeks was reduced from 98 to 30-35%, whereas germination of noninoculated peas was about 90%. Peas infected with A. ruber for 14 weeks, similar peas which had been placed under conditions favoring germination for 6 days, moistened, autoclaved peas infected with A. ruber for 7 days, or uninfected controls were comminuted in distilled water. The resulting mixture were centrifuged, and the supernatant liquids passed through sterile HA Millipore filters. The sterile liquids were mixed with a nutrient salts-sucrose agar medium and excised, healthy, surface-sterilized, embryonic pea axes placed on the medium. Extracts from infected autoclaved peas caused a characteristic mycosis in root axes, in fresh weight and elongation of the radicle, prevented the formation of root hairs, and caused extreme brittleness of the embryonic axes. Extracts from infected, living peas caused similar, though much less severe, effects. Extracts from uninfected peas had little effect on excised embryonic axes. Toxicity of the extracts was not lost on autoclaving for 15 min.

Electron microscopy of host cell and stunted diseased hibiscus blueberry. J. S. Hartmann, G. R. Hooper, & J. E. Bath (Mich. State Univ., East Lansing). Viral particles 28 to 30 nm diam were found in ultrathin sections of leaf and root tissues from hibiscus blueberry affected by host cell disease. Leaf epidermis, palisade, spongy mesophyll cells, and xylem tissue contained characteristic virus particles. Crystalline arrays and larger masses were particularly evident in root xylem tissue. Particles hexagonal in outline, 26-28 nm diam, were partially purified from diseased leaves. No such particles were detected in healthy blueberry tissues. Blueberry stunt, another disease of unknown etiology (Chen in 1971 associated a mycoplasmalike organism with the disease) was studied concurrently with host cell disease. In ultrathin sections of phloem sieve-tube elements from diseased Concord and Coville cultivars, mycoplasmalike organisms were found. Spherical, elongate, and irregular, forms as evidenced by serial section methods, were present in leaf and fruit pedicel tissue. A distinct external unit membrane, presumed ribosomes, and fibrillary nucleic acid were characteristic of these structures. No mycoplasmalike structures were observed in healthy blueberry plants of the same cultivars.

Polar lipids of fungi. J. W. Hendrix & G. Rousser (Univ. Ky., Lexington, City of Hope Med. Center, Duarte, Cal.). Polar lipids of eight fungi from all three fungal classes were determined, and phospholipids were determined quantitatively. Viable lipid extracts are tested for phosphatidyl...
present in seeds of some breeding lines. Seed-borne *Phoma*
must be eliminated for gnotobiotic host-pathogen black
root investigations. *Phoma*-contaminated seeds were treated in
a water bath at 60 or 65°C for varying time intervals
on the same day or on 2 consecutive days (seed dried at
28°C overnight) and at 20°C and germinated at 60-100%,
depending on length of treatment. Seeds treated
on a single day, whether for one or two intervals, com-
monly were contaminated with spore-forming bacteria.
Commercial monogram seed was more sensitive to length
of treatment at 60°C than was seed from breeding lines
(25% germination vs. 80%, respectively) and in the treat-
ment of three breeding line seed lots (monogram and multigerm)
for 10 min at 60°C on consecutive days failed to
eliminate all *Phoma*; whereas a 15-min treatment
interval did so with a 20% or lower reduction in germina-
tion. Variation in heat tolerance was not correlated with
differences in average seed wt. The heat tolerance of dif-
cerent cultivars and breeding lines must be ascertained by
testing.

Callus deposition and abnormal secondary wall thick-
ening as possible restrictive factors for virus spread. C.
Hiruki & J. C. Tu (Univ. Alberta, Edmonton, Can.). Po-
tato virus M (PVM)-incited local lesions on leaves of Red
Kidney bean were investigated at the three developmental
infection phases represented by nongenve, seminecrotic,
and necrotic cells from the periphery to the center of
a lesion. Thickening, due to callous deposition, was
detected on the inner wall of nonneocrotic cells immediately
adjacent to seminocrotic ones by fluorescent microscopy 3
to 4 days after inoculation. The ultrastructure of such
thickening was simpler than that of "abnormal secondary
wall thickening", due to deposition of paramural bodies,
in seminocrotic cells. The callos deposition was not
detected in necrotic cells. PVM particles were observed in
seminocrotic cells but not in necrotic cells. Temporary and
partial thickening of PVM spread by callos deposition may
occur at the early stage of individual cell infection. How-
ever, callose alone may not be the determining factor in
restricting virus spread, since PVM lesions continued to
expand for a few days even after a band of callos was
deposited around seminocrotic cells. The movement of
PVM appeared to be ultimately restricted by more com-
plex "abnormal secondary wall thickening" which prevailed
in the seminocrotic zone.

The relationship of light to growth and sporulation of
Botrytis cinerea on potato-dextrose agar. R. E. Hite (ARS,
USDA, Pa. State Univ., University Park). Certain isolates of
Botrytis cinerea sporulate profusely when grown on
potato-dextrose agar at 21°C and exposed to continuous
near-UV radiation. The same isolates grown under con-
tinuous darkness produce only sclerotia. General Electric
cool-white fluorescent lamps emit sufficient near-UV ir-
radiation to stimulate sporulation. Sporulation is accelerated by exposure to General Electric BLB fluorescent black
light lamps which emit a continuous spectrum between
300 and 400 nm with 97% of the radiation in this region.
When near-UV is filtered from the lamp source, colony
growth consists of a fluffy, aerial, vegetative mycelium
with no sclerotia and essentially no sporulation. Sporula-
tion varied in relation to the near-UV transmission of
sporulation was simpler when exposed to either cool-white fluorescent or BLB fluorescent black
light lamps, the latter being the more effective. When near-
UV is filtered from the source, the sclerotal resume growth through the production of vegetative mycelium.
The sporulation response requires very low levels of near-UV ir-
radiation under long-term exposure.

The ultrastructure of Aphanomyces euteiches during dif-
f erentiation of primary asexual spores. H. E. Hoch & J. E.
Mitchell (Univ. Wis., Madison). Ultrastructure changes in
vegetative hyphae of Aphanomyces euteiches just prior to
differentiation of primary asexual spores included the
segmentation of the central vacuole by cytoplasmic strands,
the enlargement and sitation of electron-dense vesicular
inclusions, and the movement of nuclei from a peripheral
position to one near the longitudinal axis of the hyphae.
Differentiation of primary spores occurred within minutes,
and began with the migration of cytoplasm toward each
nucleus and the withdrawal of the plasmalemma from the
hyphal wall. Simultaneously, the central vacuole evaginated
and emptied its contents into the space developed be-
tween the plasmalemma and the hyphal wall. The tono-
plast and the plasmalemma jointly formed the plasma
dec line enveloping the primary spores. Primary spores
somes, initially consisting of tightly packed tubules,
became increasingly disorganized during differentiation.
The irregularly shaped spores, generally connected in suc-
cession by strands of membrane-bound protophelim, were
entirely passive as the hyphal contents were rapidly ex-
truded through a rupture in the sporangium apex. No flagella were observed in any spores at this stage. Within
minutes after extrusion, wall material was excreted to
complete the formation of the primary spore cyst.

Effect of different soil constituents on uptake of two
benzimidazole compounds by American elm seedlings. W. K.
Hock & L. R. Schreiber (USDA, ARS, Delaware, Ohio).
Benzylam and thiabendazole (TBZ)-fungicides, applied as
drenches to 4-month-old container-grown American elm seedlings, were absorbed by the periderm and transported
to the roots of young seedlings. Leaf, wood, and bark sections were placed in petri dishes containing potato-dextrose agar seeded with either Ceratocystis ulmi (for benzyam bioassey) or Penicillium
sp. (for TBZ bioassey). The diameter of the zone of in-
hibition around each tissue section was measured to de-
termin the relative concentration of fungictoxic present
in each seedling. The zones of inhibition around tissues
from plants grown in sand and treated with benzyam were
about 2 times larger than from plants grown in treated
soil, and about 3-4 times larger than from plants grown
in a potting mixture of soil:perlite (1:2:2). When
seedlings were grown in their respective media for 90 days,
ells grown in treated sand still contained more fungictoxic
than those in either treated soil or the mixture. When-
ever peat was included in various media, uptake of both
benzyam and TBZ was inhibited significantly. Plants grown
in perlite and sand always contained the most fungictoxic.
Uptake of benzyam by seedings was greater from
heat-sterilized soil than from nonsterile soil.

Water solubility of a volatile factor inducing soil fung-
stasis. T. S. Hora, Maureen Romine, & R. Baker (Colo.
State Univ., Ft. Collins). Earlier workers have experienced
difficulty in extracting a fungistic principle from soil.
Extraction of this inhibitor was attempted, taking into
account the volatile nature of the factor as previously re-
ported. Germination of conidia of common soil fungi was
significantly inhibited by air which had passed through a
column of soil. This inhibitory influence of air, however,
was lost upon passage through water, indicating water
solubility of the volatile inhibitor. To demonstrate the
activity of the fungistic principle in aqueous solution,
air was washed in concentrated sulfuric acid was passed through
1 kg of clay loam soil (pH 8.5) packed in a 1,000-cc
burette at 25°C, and bubbled into 2 ml of water
at 2°C for 24 hr with a flow rate of 6-8 cc/min. Conidia
of eight of ten soil fungi showed a 20-65% decrease in
germination when suspended in 0.1 ml aliquots of the
water extract and placed on water agar discs for 24 hr.
These observations suggest that the volatile fungistic
factor is soluble and extractable in water.

The effect of temperature on symptom expression of
chrysanthemum inoculated with chlorotic mottle virus. R. K.
Horst & S. P. Krycyznski (Cornell Univ., Ithaca, N.Y.). Symptoms expressed by Chrysanthemum morifolium 'Duchess's
Delight', subsequent to inoculations made by Chrysanthemum chlorotic mottle virus (ChCMV), are influenced by temperature. Experiments
were performed under controlled environmental conditions so that light and moisture were nonvariables and temperature was constant at 18, 24, and 30°C. Symptoms were pronounced by determining numbers of leaves and lateral shoots developing symptoms after inoculation, and by length of time for symptoms to appear. Symptom expression, as determined by all three means of evaluation, was optimum at 24°C. Delay in symptom development was most pronounced at 18 than at 30°C. Mean temperatures resulting from a diurnal temperature program yielded results similar to those from corresponding constant temperatures. Recovery of ChCMV (by tissue implantation procedure) from the experimental plants was independent of temperature treatments.

The effect of triarimol on oxidation of glucose and acetate to carbon dioxide and on spore germination of Aspergillus niger and Cladosporium cucumerinum. L. D. Houseworth, E. W. Brumton, & B. G. Tweedy (Univ. Mo., Columbia). C-1, C-3, C-4, C-6, and uniformly labeled glucose and C-6 labeled gluconate were used to determine the effect of triarimol on the respiratory pathways of germinating spores of Aspergillus niger and Cladosporium cucumerinum. Triarimol did not affect the quantity of CO2 evolved when the organisms were incubated with any of the labeled substrates, indicating that the fungicide has no effect on the rate or participation of pathways for glucose catabolism. To determine the effect of triarimol on Kreb's cycle, C-1 and C-2 labeled acetate were administered, and triarimol did not alter the rate of [14CO2] evolution from labeled acetate. Fractionation of A. niger spores previously incubated with C-2 acetate showed that triarimol had no effect on the quantity of the methyl moiety incorporated into various organic fractions. These studies suggest that triarimol has no effect upon respiratory pathways during the initial stages of spore germination. Spore germination studies showed that triarimol had little or no effect on the initial process of both organisms, but germ tube elongation of C. cucumerinum and A. niger was arrested at 0.5 μg/ml and 20.0 μg/ml, respectively.

Chemical limitation of discoloration and decay associated with wounds in red maple and yellow birch. D. R. Houseworth (USA, Northeastern Forest Exp. Sta., Hamden, Conn.). Chemicals, including fungicides, antibiosis, and paraformaldehyde, were placed immediately into increment borer wounds made in 65 red maple and 35 yellow birch trees in New Hampshire. Three years later, the trees were dissected, the vertical extent of the discoloration and decay associated with the wounds was measured, and increments in thickness were measured. Discoloration in both species was reduced significantly by chemicals. Red maple wounds treated with glue, and yellow birch wounds treated with glue or Methocel, had the least discoloration. In red maple, the most extensive discoloration and decay, except for controls, occurred in trees treated with paraformaldehyde. Decay in this species was least when wounds were treated with Actidione. Microorganisms isolated from discolored and decayed tissues were typical inhabitants of wound-associated defects, and included bacteria, new-Hymenomycetes, and Hymenomycetes. Graphium sp. was common in both tree species.

Cassia obtusifolia, a possible reservoir for inoculum of Colletotrichum fragariae. C. M. Howard (Univ. Fla. Agr. Res. Ctr., Center, DeLand). Strawberries anthracnose caused by Colletotrichum fragariae has been a serious disease in Florida strawberry nurseries for many years. Since 1968, anthracnose has severely limited production in Florida of the popular but extremely susceptible cultivar, 'Tonga'. Colletotrichum fragariae has not previously been reported attacking plants of any genus other than Fragaria and Duchesnea. In 1970, 3- to 6-inch chance seedlings of Cassia obtusifolia in strawberry nursery flats were found to be infected by a Colletotrichum species. Infection by a Colletotrichum species of C. obtusifolia seedlings up to 18 inches high in and around a strawberry nursery were found later. Typical anthracnose lesions developed on runners and petioles of potted strawberry plants inoculated with spores isolated from isolates obtained from both Cassia sources. Aervula with black setae developed on C. obtusifolia seedlings inoculated with isolates from the weed or from strawberry plants. Cassia obtusifolia is widely distributed in west central Florida throughout the year, and may be the principal source of primary inoculum of C. fragariae.

The role of bacteria in inoculated seedlings of sugar maple, Acer saccharum. R. Howland (Univ. N.H., Durham). Sugar maple seedlings were inoculated with several selected species of bacteria representing the genera Bacillus and Pseudomonas. Subsequent inoculation of the inoculated species proved successful in the majority of cases. Naturally introduced gram-negative organisms have also been isolated frequently from the same inoculated seedlings. Wood sections taken from the nonwounded control trees were free of bacteria. Several species representative of both the Aerobacter-Klebsiella and Pseudomonas groups have been identified. Cultural and histological studies indicate that some of the gram-negative bacteria that develop naturally in wounded sugar maple trees may be of importance in predisposing the tissues to further colonization by numerous other organisms higher than bacteria.

Peak concentrations of potato yellow dwarf viruses in Nicotiana rustica revealed by infectivity assay on vector cell monolayers. H. T. Hsu & L. M. Black (Univ. Ill., Urbana). The concentrations of two varieties of potato yellow dwarf virus (PYDV) in the stem portion bearing the acutely infected leaves of Nicotiana rustica plants, grown at 30°C, were determined by infectivity assay on their respective vector cell monolayers. The variant SYDV (sanguinella yellow dwarf virus) was assayed at pH 7.93 on AS2 cells, whereas variant CYDV (constricta yellow dwarf virus) was assayed at pH 5.25 on AC20 cells. Both viruses attained a peak concentration 10 days after the first appearance of systemic symptoms. The relative virus concentrations of each variety were calculated as percentages of its peak concentration. About 30% of SYDV infectivity was lost by the 12th day and 40% was lost by the 14th day; thereafter the infectivity remained almost constant until the last measurement on the 24th day. About 70% of the peak infectivity of CYDV was lost by the 14th day, and more than 50% was lost by the 14th and 16th days. Purification to obtain highly infectious PYDV free from microorganisms was best achieved by harvesting infected plant materials at the peak virus concentration. Purified viral inocula can be stored at -80°C for more than 1 year without detectable loss of infectivity.

58S transfer from host to parasite during primary infection of barley by Erysiphe graminis f. sp. hordei. S. C. Hsu & A. H. Elingboe (Mich. State Univ., E. Lansing). Six-day-old barley plants with different MI genes (Mla, Mlg, Mlb, or mle) were inoculated with two isolates of Erysiphe graminis f. sp. hordei with Ps Pp Pk Ph and Pa pg Pk Ph. The percentage of the applied conidia that infected the host cells and produced secondary hyphae at least 10 μ in length was recorded at 2-hr intervals between 16 and 30 hr after inoculation. At various hours after inoculation, leaves carrying the several different MI genes were allowed to take up 58S, for 5 hr. The portion of the parasite on the surface of the leaf was removed from the leaf with parlodion, and the 58S in the parlodion strips was determined by scintillation counting. The percentage of high efficiency of infection (Pa Pp Pk Pp/mla mlg mlk mlp) and Pa pg Pk Pp/mla mlg mlk mlp had also high rates of 58S transfer from host to parasite. The incompatible parasite/host genotypes (Ps Pm, Pp/mlb, or Pp/mlb) had lower efficiency of infection and a reduced rate of 58S transfer from host to parasite, as compared to compatible combination.
Properties of chloroplast structural proteins isolated from tobacco leaves undergoing bacterially induced hypersensitive reaction. J. S. Huang & R. N. Goodman (Univ. Mo., Columbia). Chloroplasts were isolated from tobacco leaves harvested by homogenization and differential centrifugation, 20 minutes, 3 hr., and 6 hr after infiltration with 10^8 cells/ml of Erwinia amylovora. The hypersensitive reaction (HR) is normally visible at 6-7 hr after infiltration. Isolated chloroplasts were sonicated and centrifuged to obtain membrane and lamellar fragments. Structural proteins (SP) were isolated from the fragments, using acetyl. The resultant acetylene powder was dissolved in 0.002 M Tris buffer (pH 8.5) containing sodium dodecyl sulfate and urea. The protein fraction collected was that precipitated with 12-16% (NH_4)_2SO_4. All SP thus prepared showed UV absorption spectrum maxima at 280 nm and minima near 250 nm. They were heterogeneous, however, and were resolved as several protein bands by electrophoresis. The SP prepared from leaves 6 hr after infiltration (during the "developing" stage of HR) when HR symptoms were present, had a lower O.D.ratio, reduced phospholipid-binding capacity, and lower solubility than SP isolated from the leaves 3 hr after infiltration (during the "incubation" period). These differences in criteria of SP were even greater between the 3-hr sample and that taken 20 min after infiltration (during the "induction" period). Therefore, it seems, therefore, that SP denatures progressively as a consequence of HR.

Factors affecting production of microsclerotia by species of Cylindrocladium, B. B. Hunter & H. L. Barnett (West Va. Univ., Morgantown). Cylindrocladium scoparium, C. floridanum, and C. fimbriatum were tested for production of microsclerotia. Effects of temperature and light (40 w General Electric cool-white fluorescent lamps, 125 ft-c) were tested on glucose-casein hydrolysate agar. A range of 24-28°C was optimum, and light had no apparent effect on numbers of microsclerotia. Effects of concentration and ratios of carbon to nitrogen were tested at 25°C on glucose-potassium nitrate agar medium and on sand supplemented with the same nutrients, minus agar. Similar results in agar and sand cultures were obtained with C. scoparium. The amount of mycelium decreased, but the number of microsclerotia increased (15 mg and 350 mg/culture, respectively) with widening carbon:nitrogen ratios (1:1 to 100:1). Casein hydrolysate and r-tyrosine provided carbon as well as nitrogen for microsclerotial production, but r-glutamic acid did not. Microsclerotia were recovered from nursery soil to which 2 g of glucose were added to 25 g of soil and incubated in the laboratory for 12 days, but were not recovered from soil without added glucose.

Peroxidase activity in wheat leaves infected by Puccinia recondita. L. B. Johnson & B. A. Cunningham (Kansas State Univ., Manhattan). Peroxidase activity was determined in primary leaves of healthy and inoculated near-isogenic wheat lines differing in susceptibility to isolate 85-68B of the leaf rust pathogen, Puccinia recondita. Activity was similar in healthy leaves of both lines, and increased with leaf age. In inoculated Thatcher, which develops a high infection type (4), peroxidase activities at 2-9 days were 20-48% higher than in healthy tissue. In LR10(TC), which developed a low infection type (2) in these experiments, peroxidase had increased up to 100% over healthy control after 9 days. Disease development did not affect buffer-soluble protein in trichloroacetic acid-precipitable in either line. Peroxidases from healthy and inoculated LR10(TC) were separated on a Sephadex G-100 column into two distinct molecular weight groups. Although total peroxidase activity was greater in infected tissues, the ratio of activities in these two molecular weight groups did not differ significantly between 9-day rusted LR10(TC) and healthy tissue. The low molecular weight group is composed of peroxidase isoenzymes with an average molecular wt of near 35,000. The higher molecular weight fraction has at least twice this molecular wt. The physiological significance of these two molecular sizes is unknown.

Telephones as computer terminals for plant disease information. A. L. Jones & S. B. Hard. (Mich. State Univ., E. Lansing). An automated method for handling questions on fungicide compatibility, disease diagnosis, and calibration of spray equipment has been produced using 12-button telephones as computer input/output terminals in combination with an IBM 7777 audio response unit capable of giving a verbal reply to digital input. Codes used frequently for identifying the user are entered using a "Call-a-matic" device or preprinted plastic cards and a card-dialer option. Names of fungicides and disease symptoms are entered in coded form, using the push-button keyboard. Data requests by the program and output from the analysis are returned over an auxiliary speaker or hand receiver. The programs, written in FORTAN IV, allow the operator to alter the input values to obtain custom replies. Field tests indicate that the operating procedures are simple enough for use by Cooperative Extension workers. Because low cost, multipurpose telephones and standard communication lines are used, the voice answerback system is potentially available to a large number of remotely located users of disease information. Push-button telephones connected in parallel with dial telephones can be used in areas without push-button telephones.

Interaction of five soil-borne pathogens of tomato. J. P. Jouny & A. J. Oveeman (Univ. Fla., IFAS Agr. Res. Education Center, Bradenton). Tomato seedlings of a breeding line resistant to Fusarium oxysporum f. sp. lycopersici (Fol), race 1, but susceptible to race 2, were grown in methyl bromide and steam-sterilized, composted, Leach line sand artificially infested with Fol races 1 and 2, Verticillium albo-atrum, Tylecorkhychus capitatus, and Meloidogyne incognita, using all possible combinations of these five pathogens. Race 1 interfered with pathogenesis by race 2 at 22 and 25°C. In 14 of 16 race 2 combinations containing race 1, disease incidence was significantly less when compared to identical combinations without race 1. Tylecorkhychus capitatus did not affect Fusarium or Verticillium wilt development at 22 or 25°C. Meloidogyne incognita did not affect the development of Verticillium wilt at either temperature, but increased the incidence of Fusarium wilt incited by race 2 at 22°C.

Effect of glucose and amino acids on growth and sporulation of Fusarium oxysporum f. sp. lycopersici. Race 2. J. P. Jorns & S. S. Wolf. (Univ. Fla., Bradenton). Race 2 was cultured 7 days in pH 6.0 liquid media containing in addition to a basal inorganic nutrient solution, 0.0, 0.3, and 1.2% glucose alone and in combination with 1,250 ppm of each of the following amino acids: L-asparagine, L-aspartic acid, L-cysteine, L-glutamic acid, L-proline, and L-tyrosine. Growth increased as the glucose concentration increased. All amino acids except cysteine supported some growth in the absence of glucose. Tyrosine increased growth the most at each glucose level, although it furnished less nitrogen than any other amino acid utilized. Few microconidia were found in the basal medium (no glucose or amino acids). Microconidial numbers increased as the glucose concentration increased. Several amino acids increased microconidial sporulation in the absence of glucose, but none did so at the 0.3% glucose level. Proline and glutamic acid increased microconidial production at the highest glucose level. No microconidia formed in the basal medium, but production was increased by increasing glucose concentrations. Asparagine inhibited microconidial formation, whereas proline, aspartic acid, and glutamic acid increased microconidial numbers 2 to 3 times. However, the effect of amino acids on microconidial formation was highly dependent on the glucose concentration.

Inhibition of the polysaccharide-degrading ability of Fusarium oxysporum f. sp. lycopersici culture fluids by a protein isolated from tomato stem cell walls. T. M. Jones,
A. J. Anderson, & P. Albersheim (Univ. Colo., Boulder). When cultured in liquid medium containing 1% cell walls (10%) isolated from tomato stems as the sole carbon source, Fusarium oxysporum secretes a variety of basidio- 
chid-degrading enzymes. These include an endopolygalacturonase, a cellulase, α - and β-1-4-galactosidases, an α-1-4-arabinohydrolase, and a β-1-4-xylosidase. The enzyme activities appear in the culture fluid in a definite order: the polygalacturonase is first to reach maximal activity, followed by the glycosidases, with cellulase activity reaching a peak after the other enzyme activities have diminished. The ability of culture fluid enzyme mixtures to degrade cell walls isolated from tomato stems has been demonstrated by determining the rate of the material solubilized in the medium through the treatment of the walls with enzyme mixtures, and by examining the composition of the cell walls prior to and following enzyme treatment. A protein extracted from tomato stem cell walls by treatment with 0.5 M potassium phosphate buffer, pH 7.1, has been observed to inhibit the action of the polygalacturonase and to prevent cell wall degradation by culture fluid enzyme mixtures. Similar polygalacturonase inhibitors have been obtained from Phaseolus vulgaris stems and from suspension-cultured Acer pseudoplatanus cells.

Production of chlamydosporos by Phytophthora palmariorum. Janice Y. Kadooka & W. H. Ko (Univ. Hawaii, Honolulu). Only thin-walled chlamydosporos were produced by Phytophthora palmariorum, using the submerged and low temperature method described by Tuset & Paraska. However, observations of naturally infected papaya fruits showed that as many as 90% of the chlamydosporos had thick walls. Similar results were obtained by culturing the organisms for 4 weeks at 22°C in a liquid medium containing 30% papaya juice and 0.2% CaCO3. Only thin-walled chlamydosporos were observed in the 1st week of growth in the papaya medium, moderately thick-walled chlamydosporos in the 2nd week, and mostly thick-walled chlamydosporos in the 4th week. The percentage germination of thick- and thin-walled chlamydosporos on water agar was 85 and 92%, respectively. On V-8 juice agar, thick-walled chlamydosporos did not germinate, whereas 80% of the thin-walled chlamydosporos germinated.

Histology of onion leaves infected with Pseudomonas cepacia. S. O. Kawamoto & J. W. Loberger (Cornell Univ., Ithaca, N.Y.). When stably inoculated into the base of onion leaf blades, Pseudomonas cepacia moved into the leaf sheath, invaded the intercellular spaces, and formed bacterial masses. Although large bacterial masses occurred, separate small groups of bacteria were frequently observed in the intercellular spaces of both blade and sheath. Such distribution was observed when the lesions were periodically sprinkled with water. Except for the yelworms and endodermal cells, expanding bacterial masses in the intercellular spaces crushed the surrounding cells. In the blade, the large parenchyma cells were the first to be crushed by the bacterial masses; the smaller, more closely packed parenchymatous cells near the periphery of the blade were crushed later. Tissues composed of these cells eventually were macerated. In the shoot, bacteria were widespread among the loosely organized parenchyma cells adjacent to the adaxial epidermis. Bacterial masses and scattered bacteria in close association with host cell walls were commonly observed in this area. Bacterial masses in the intercellular spaces of the closely packed parenchyma cells beneath the abaxial epidermis were restricted to dense, compact masses.

Nature of aerial strands of Erwinia amylovora. H. L. Keil & T. van der Zut (ARS, USDA, Beltsville, Md.). Erwinia amylovora produced abundant aerial strands on inoculated Bartlett pear trees sprayed with 1% commercial spray oil. Morphology of the strands was studied in a scanning electron microscope. Strands were easily distinguished from natural host tissues. They were brittle and broke when crushed, resulting in irregularly fractured ends which showed proportionately more matrix than bacteria. Strands, when dissolved in water and vacuum-evaporated, formed a network of adhesive material which connected the bacterial cells. When the weebike network was placed in ethyl alcohol, this material was dissolved, exposing individual cells. Smooth strands were most common, had about the same diameter throughout their length, and appeared to have a uniform quantity of matrix. We believe that this matrix was extruded at a constant pressure. Rough strands appeared to be formed by irregular pressure. Oil apparently plugs the strands, opening the plant, thus creating internal pressure. As the bacterial matrix is forced out it solidifies upon exposure to air, and the strand increases in length as more matrix is added basipetally. Since bacteria in these strands consistently proved virulent, aerial strands in nature may play an important role in the epidemiology of fire blight.

The electrostatic response of zoospores of several species of Phytophthora K. L. Kim & G. A. Zentmyer (Univ. Cal., Riverside). Electrotaxis was studied with five species of Phytophthora: P. cinamomoni, P. palmivora, P. capsici, P. cactorum, and P. citrophthora. In deionized H20 at 0.1-0.5 μm d-c (≤ 1.2 V/cm), zoospores of all five species exhibited an excited and increased swimming velocity as they approached the anode. This was usually followed by an active-oriented attraction and accumulation. As intensity of current increased, a repulsion zone appeared below the anode while a progressive immobilization of zoospores occurred at the cathode. The zoospores at the cathode exhibited a decreased swimming velocity, rotation, cessation of motion, and bursting. Studies were also made in solutions of various mono- and disaccharides and metabolically inhibitory in an attempt to interfere with the basic pattern of electrostatic response. None of these completely alleviated or inhibited electrotaxis without also affecting motility. Microelectrophoresis studies of swimming and pre-enveloped zoospores indicated that both were negatively charged. Electrokinesic force could contribute in part to electrotaxis. Both chemical and electrical stimuli can induce a change in the membrane potential of zoospores which might serve as a signal to initiate a respective chemotactic or electrotactic response.

Pythium stem blight of beans. S. H. Kim & J. G. Kandt (Univ. Md., College Park, Md.). Association of Pythium spp. with bean stem blight was evident in New York and Delaware during 1968, 1969, and 1970. Pythium ultimum was isolated from diseased bean (Phaseolus vulgaris) plants in the field during June, July, and September; P. anamidquatatum was isolated only during the warmer months of July and September. Both species, however, were isolated from the soil at all times of the year. Infection and the development of symptoms required the presence of water droplets containing the organism on susceptible tissues. When water droplets containing zoospores or mycelia evaporated within 8 hr of inoculation, disease development at 23°C was prevented. Maintaining water droplets on buds and leaves for 2 days in a moist chamber, or 5 days of continuous sprinkler irrigation in the field, permitted full disease development on Tenderecroft bean plants inoculated with zoospores of P. anamidquatatum. Laboratory and greenhouse studies on P. anamidquatatum revealed that zoospores from zoospores were as pathogenic as zoospores from sporangia. Zoospores were released from 1-year-old dry zoospores in water. Thus, zoospores may be vital to the development of stem blight of beans caused by P. anamidquatatum.

Resistance of Xanthomonas dierenbachiae isolates to streptomycin. J. F. Knauss (Univ. Fla. Agr. Res. Center, Fort Lauderdale). Bacterial leaf spot and tip burn of Philodendron oxycardium incited by Xanthomonas dierenbachiae is the most important foliar disease of foliage plants grown in Florida. Foliage grower applications of 200 to 400 ppm streptomycin under summer conditions at 4- to 7-day intervals for disease control were found ineffective, and ap-
peared to intensify disease development. Direct isolation of the pathogen from infected leaves taken from streptomycin-sprayed plants, from foliage nurseries, and also from experimentally inoculated but nonsprayed plants onto potato-dextrose-agar resulted in a dramatic increase in 1,000 embryonated eggs. Isolations revealed vigorous growth of the pathogen only from sprayed leaves. Growth occurred at all streptomycin concentrations. In isolations from sprayed leaves, only slight growth occurred at 100 ppm streptomycin with none at 200 ppm concentrations. Inoculations with streptomycin-resistant isolates of *P. oxalacia* sprayed with 600 ppm streptomycin resulted in more severe disease development than did similar inoculations made with streptomycin-sensitive isolates.

A selective medium for determining population of *Rhizoctonia solani* in soil. W. H. Ko & Frances K. Hora (Univ. Hawaii, Honolulu). A medium containing 1 g KIPO₃, 0.5 g MgSO₄·7H₂O, 0.5 g KCl, 10 mg FeSO₄·7H₂O, 0.2 g NaNO₃, 0.4 g gallic acid, 90 mg Dexam [sodium P- (dimethylamino)benzamidazolosulfonate], 50 mg chloramphenicol, 50 mg streptomycin sulfate, and 20 g agar in 1 liter distilled water enhanced the growth of *Rhizoctonia solani* from soil and suppressed the development of undesired molds. Recovery of the pathogen from natural soil inoculated with sclerotia was 90-100% with this selective medium. For determining *R. solani* population in naturally infested soil, 1 g of soil was evenly distributed in 10 clumps on a plate of selective medium and observed microscopically for *R. solani* mycelium after 24- and 48-hr incubation. The population of *R. solani* in natural soil ranges from 1 to 9 propagules/10 g dry soil. Soils inoculated with the same amounts of *R. solani* sclerotia were capable of causing pre-emergence damping-off of beet. Low populations of *R. solani* in soil may account for the unexpected high moisture of the dilution plate method used for this fungus. The combination of carbon and nitrogen sources selective for *R. solani* and inhibitors of undesired microorganisms is the basis for the selective medium.

The effect of H₂O₂ under acid conditions on strength, swelling, alkali solubility, reducing capacity, and weight loss of cellulose. J. W. Koenig (Forestry Sci. Lab., Research Triangle Park, N.C.). In earlier studies, brown and white rot basidiomycetes produced extracellular H₂O₂. Formation of H₂O₂ by both groups suggested that H₂O₂ might be involved in the decomposition of cellulose. In vitro, H₂O₂ swollen cotton-fiber cellulose and increased slightly its alkali solubility and reducing capacity. H₂O₂ affected neither strength nor the reducing capacity or weight loss of cellulose. A rot-infected treatment was not affected by the presence of iron. Weight loss increased with H₂O₂ concentration, but strength loss increased more rapidly, especially at low concentrations of H₂O₂; swelling, alkali solubility, and reducing capacity increased in a more linear relation to H₂O₂ concentration than strength loss. These results suggest that H₂O₂ produced by microbial enzymes may swell cellulose, and that previously unconsidered mechanisms could be involved in cellulolysis. H₂O₂-swollen cellulose is a suitable substrate for assessing the role of certain cellulase components.

Effect of low temperature on germination of teliospores of *Cronartium ribicola*. R. G. Kreeb (USDA Forest Serv., Logan, Utah). Along the southern limits of its known range in the Rocky Mountains, blister rust is most frequently encountered in white pines at elevations below 8,000 ft. This study was made to determine if low temperatures such as might occur in pine stands at higher elevations, inhibit the spread of the rust by depressing basidial sporulation. Tela from inoculated *Riberia fascnare* kept in growth chambers at 18°C days and 13°C nights were removed from leaves and subjected in petri plate moist chambers to constant temperature at 3°C intervals from -3 to 15°C for periods of 3, 6, 12, 24, 48, and 72 hr. Basidiospores were cast from telia by 12 hr at 15°C, by 24 hr at 6°C, and by 48 hr at 3°C. No basidiospores were cast at 0 and -3°C. Casting rates were reduced at low temperatures; after 72 hr, total basidiospores cast per mm of telium averaged from a few hundred at 3°C to several thousand at 15°C. These results suggest that at low temperatures the telium can function efficiently at the low temperatures; at 3°C, the fusiform rust has a low activity for long, moist periods, and greatly reduce the inoculum potential of *Cronartium ribicola* in high-elevation white pine forests of the central Rocky Mountains.

Some properties of *chrysanthemum chlorotic mottle* virus. S. P. Kryczynski, R. K. Horst, & A. W. Dlmoek (Cornell Univ., Ithaca, N.Y.). Current studies confirm that *chrysanthemum chlorotic mottle* virus is a virus. Since the causal virus has previously occurred in the *Chrysanthemum* virus, and no evidence of relationship to other common viruses has been found, the name *chrysanthemum chlorotic mottle* virus (ChCMV) is proposed. A technique which assures a good, consistent infection level by mechanical (sap) inoculation has been developed. The technique was examined on any of 15 other plant species, representing 6 families, inoculated by the abovementioned method. The most commonly used virus indicator plants, some of them Composite. ChCMV never recovered either from inoculated or from newly developed leaves of any of these plants.

**Triarimol (EL-273), a new fungicide for tart cherry disease control.** D. H. Laid & C. D. Christensen (Eli Lilly & Co., Plainwell, Mich., Cazenovia, N.Y.). Triarimol [α-(2,4-dichlorophenyl)-α-phenyl-5-pyrimidinemethanol] was applied to foliage of tart cherries with high-pressure hand guns on commercial schedules of 20-30 ppm in several replicated experiments in Michigan and New York. Harvested fruit was subjected to standard fruit quality determinations. Dodine at 300 ppm and Difolatan [N-(1,1,2,2-Tetrachloroethoxy) sulfinyl-ethyl-4-cyclohexene-1,2-dicarboximide] at 500 ppm were reference compounds. At harvest, Dodine at 30 ppm provided 98-99% control of cherry leaf spot (Coccomyces hiliensis) and 97% powdery mildew (Podosphaera oxyacantha) control. Ninety to 100% brown rot (Monilia fructicola) control was provided by 30-40 ppm. Dodine and Difolatan provided leaf spot and brown rot control, but were ineffective for the control of powdery mildew. Triarimol ranked between dodine and Difolatan for the prevention of defoliation from late season leaf spot invasion. Yield, ripening rate, terminal growth, fruit, and lateral bud set were not affected by triarimol. Fruit quality as determined by color, character, freedom from defects, freedom from surface pits, grade, U.S. No. 1, size, pH, and percent sugar were also unaffected by triarimol.

**Genetic evidence for pleiotropic factors involved in the tumor-inducing ability of Agrobacterium tumefaciens.** R. A. Langley & C. I. Kado (Univ. Cal., Davis). N'-methyl-N'-nitro-N-nitrosoguanidine was used to mutagenize cells of *A. tumefaciens* LBI35. Mutants auxotrophic for various amino acids, resistant to neomycin, kanamycin, or Ampicillin, were tested for tumor-inducing ability. A number of histidine, leucine, and tryptophan auxotrophs were avirulent on sunflower. Addition of the nutritional requirement to the inoculum (1 mg/ml of the amino acid) did not restore virulence. Spongiform tumors in revertants of histidine-avirulent auxotrophs simultaneously regained virulence. Neomycin- and kanamycin-resistant mutants were highly virulent, indicating that alterations in the 305 ribosomal proteins has no direct effect on virulence. Rifampicin-resistant mutants were also as virulent as the parental wild-type strain, suggesting that changes in the structural gene of
DNA-dependent RNA polymerase also are not directly involved in promoting virulence. These results indicate that the tumor-inducing ability is not due to a single gene product, but requires an alteration of the physiology of the bacterium.

**Enhancement of diagnostic symptoms of potato spindle tuber virus by manganese.** C. R. Lee & R. P. Singh (Can. Dep. Agr., Fredericton, New Brunswick). The tomato cultivar, Allerfruhe-Hemeland, has been used successfully as an indicator host for the mild strain of the potato spindle tuber virus (MSTV). Recently, some inconsistency in symptom expression was encountered. Manganese nutrition was suspected. Tomato plants were grown in sand cultures containing various levels of Mn, Fe, or Zn. Increasing Mn from 0 to 9 µg/ml drastically increased the characteristic necrosis symptoms with MSTV, as well as with severe strain. At 9 µg/ml Mn, increasing Fe and Zn in media to 20 µg/ml decreased the necrosis. Zinc had no effect on the necrosis. Other tomato cultivars (Bonny Best, Michigan-Ohio, Rutgers, and Sheyenne) previously reported symptomless for MSTV developed necrosis when grown in sand cultures receiving 9 µg/ml Mn.

**Study on interaction of soybean mosaic and bean pod mottle viruses in soybean.** Yee-Siong Lee & J. P. Ross (N.C. State Univ., Raleigh, N.C.). Inoculation of soybeans with soybean mosaic virus (SMV), followed 1 week later by bean pod mottle virus (BPMV), often produced top necrosis. Factors affecting this reaction and associated cellular changes were studied. Conditions favoring top necrosis included diurnally fluctuating (27°C-21°C) rather than constant temperature, and structures indicating high SMV titers, and inoculation of young rather than old plants. SMV titers in young leaves and stem apices were highest 3 weeks after SMV inoculation when determined by serological and infectivity dilution end points and by virus particle counts. By the last harvest, SMV titers were twice as high in doubly as in singly infected plants. Neither light microscope observations for inclusion bodies nor electron-microscope observations for pinwheel, circular, and bundle inclusions revealed any consistent differences in number or size of the structures between SMV and SMV-BPMV-infected plants. BPMV-like particles and SMV inclusions were found in the same cells. Starch granules were more numerous and larger in healthy than in singly or doubly infected plants. Lipid globules in chloroplasts adjacent to necrotic tissue in doubly infected plants were larger in young, remote tissues; lipid globules were rarely observed in healthy plants.

**Inhibition of F-2 (zealenone) biosynthesis and perithecia production in Fusarium roseum 'Granimeraeum.'** J. R. Lieberman, J. C. Wolf, H. R. G. Rao, & P. K. Harelln (Univ. Minn., St. Paul). Dichlorovos (2,4-dichlorovinyl dimethyl phosphate) was applied at 10²-10³ µg to 1 cm discs of Coons' medium agar seeded with spores of *F. roseum 'Granimeraeum.'* The carrier solvent was 10 parts diethyl ether-dimethyl sulfoxide (97:3, v/v). Ten days after seeding, control discs had perithecia, whereas discs treated with 10 µg dichlorovos had none. Cultures of the fungus on moist, autoclaved rice grown at 25°C for 1 week and held at 12°C for 3 weeks yielded 150 ppm F-2 (zealenone), whereas if the medium contained 20 ppm dichlorovos, no F-2 could be detected. When 5-g slices of a culture of *Fusarium* on moist, autoclaved rice were sprayed with 2.5 ml diethyl ether containing 1.8 mg dichlorvos, 1-1°C- acetate was incorporated into F-2 only 50% as rapidly as in slices sprayed with ether alone. Vegetative growth of the fungus was not affected by the toxicant. These results suggest that production of perithecia may depend on F-2 in strains of *F. roseum 'Granimeraeum'* which normally produce the sexual stage.

**Influence of transmissible diseases on toxin production in Helminthosporium maydis.** G. D. Lindberg (La. State Univ., Baton Rouge). Almost all isolates of *Helminthosporium maydis* obtained from corn in Louisiana in 1970 were diseased. The agent of disease was readily transmitted to apparently healthy *H. maydis* (HIM-T) by
hymal contact inoculation with diseased *H. maydis* (DHM-T). Symptoms of diseased colonies were lyed cells and arrested growth. HM-T frequently changed to diseased (DHM-T), as if spontaneously. The toxin produced by HM-T and DHM-T was > 10 times more inhibitory to the roots of F1a-200A corn (T-cytoplasma) than was the toxin produced by a disease isolate of Hooker's race O of *H. maydis*. The diseased isolate of Hooker's race O produced a toxin 5 to 10 times more active against roots of Ls-2111 corn (regular cytoplasma) than the HM-T and DHM-T toxin. The agent of disease of DHM-T was not transmitted to Hooker's race O. Isolate BslpHM was transmitted to a variant set of seeds in a colony of HM-T, and was obtained also from Hooker's race O. BslpHM was immune to infection by the agent of disease of DHM-T, and produced no toxin. The agent of a diseased isolate of BslpHM (C) BslpHM was transmitted to HM-T, but not to HM-T. C BslpHM produced toxin, and toxin activity was induced in BslpHM by transmission of the agent of disease. A close relationship was observed, therefore, in strains of *H. maydis*, to susceptibility to a given agent of disease in the fungus and to specific toxin activity.

Presence of an antifungal compound in peanut cotyledons. D. L. Lindsey & R. B. Turner (New Mexico State Univ., Las Cruces). Surface-sterilized seeds without tests from freshly harvested peanuts were found to inhibit the growth of *Aspergillus flavus* and *Trichoderma viride* on peanut agar medium. Inhibition of these fungi by seeds with intact tests, by tests from freshly harvested peanuts, or by seeds without tests from dried peanuts was observed. A material inhibitory to *A. flavus* was extracted from acetone from the cotyledons of freshly harvested peanut seeds kept for 3 days in sterile petri dishes containing 10 ml sterile distilled water. The inhibitor was further purified by extraction of the dried acetone extract with peroxide-free dry diethyl ether. The ether-soluble fraction was chromatographed on silica Gel G thin-layer plates. An active zone was detected at Rs 0.4 using benzene:1,4-dioxane (8:2) as the developing solvent. Partition extraction has indicated the active material to be an acidic.

Chemical composition of the oospore-oogonium wall complex of Phytophthora megasperma var. sojae. Eleanor Lippman, D. C. Erwin, & S. Bartnick-Garcia (Univ. Cal., Riverside). A homobamboo strain of *Phytophthora megasperma* var. sojae (PsO45) produces abundant oospores on clarified V-8 juice agar in the dark at 25°C. These oospores (enclosed in their oogonium walls) were separated from the mycelium by homogenizing the entire culture and washed by centrifugation. The cells were broken in a Teflon Potter-Elvehjem tissue grinder, and the walls separated by differential centrifugation. The final wall fraction was free of cytoplasmic contamination and contained oospore walls, oogonial walls, and probably some unrecognized fragments of antheridial walls. The bulk of these walls was composed of alkali-insoluble glucan polymers (78%), the balance consisting of protein (10.6%), readily extractable lipid (5.7%), and bound lipid (5.6%). Only 7% of the wall dry weight was cellulose; the susceptibility of the walls to digestion of exo-β-1,3-glucanase or a mixture of endo-β-1,4-glucanase and endo-β-1,4-glucanase revealed that the remainder of the wall glucan consisted of β-1,3-linked polymers similar to those found in hyphal walls of Oomyctes.

Survival of Helminthosporium maydis, race T, in Georgia. R. H. Lurry & D. R. Sturges (Univ. Ga. Coastal Plains Exp. Sta., Tifton). Infectivity of southern leaf blight infected field corn residue was studied under natural and artificial conditions. Replicated field plots were disk-harrowed, rotary-chopped, or plowed. Leaf residues were also placed in wire cages in the field and stored in cloth bags in an open shed. Samples were collected 15 January–15 March 1971. Residue was separated into stalk and leaf-shuck tissue, washed in water and filtered and centrifuged, and total Helminthosporium conidia/ml were counted. The suspended conidion was sprayed on cultivars containing both Texas male sterile (T) and normal (N) cytoplasm. Lesions were counted, and selected lesions placed in moist chambers to determine conidial production. Numbers of lesions produced from 1 g of residue declined from 13 to 0.4 in tissue in field plots during the winter, but remained at 24–40 g in leaf tissue in cages and 55–97 g in leaf tissue in cloth bags. Filtrates from residue samples from nine counties collected in late February caused 0.2–13.8 lesions/g. Only occasional typical lesions were noted on N plants, and these lesions produced only one-half as many conidia as lesions on T plants. Infectivity was greatest from leaf and shuck residue, although more total Helminthosporium conidia were observed in filtrates from stalk tissue. No H. turcicum or H. carbonum conidia were observed.

Helminthosporium maydis and temperature, light, and humidity. R. J. Lukens, P. E. Waggoner, & J. G. Horsfall (Conn. Agr. Exp. Station, New Haven). Conidiospores of the T-strain form most abundantly at 18 to 20°C on washed hyphal fragments suspended in 0.02 M phosphate buffer (pH 6.3) and placed on filter paper. No conidiospores form at 10 and 35°C. Two to four times as many form in light as in darkness. Sporulation of conidiospores is inhibited by light at 23°C and warm. Conidial formation is max at 18 to 23°C in darkness, whereas no conidiation occurred at 30°C even in darkness. Conidia appear after 4 to 7 hr of darkness at 23°C. Four to 5 days after inoculation of maize leaves, conidia form within 24 hr after wetting the lesions at 23°C. Spores germinate imperfectly, and do not form appressoria on glass slides at 94% RH. In water, spores germinate about 50% in only 3 hr at 23 or 35°C. At 23°C, appressorial formation is 50% after 6 to 9 hr, but appressoria are rare after 9 hr at 35°C. Keeping leaves wet for 3 hr after inoculation permits some infection, but max infection follows 24 hr of wetness. Lesion enlargement is optimum at 30°C, and seeds about the same whether leaf surfaces are wet or dry.

Histopathology of infection of bean hypocotyls by Sclerotinia sclerotiorum. R. D. Lubsden & Roberta L. Dow (ARS, USDA, Beltsville, Md.). Dome-shaped infection cushions formed on bean hypocotyls within a few hours after inoculation with *S. sclerotiorum*-infected oats. Ingress into host tissue occurred by multiple infection pegs that formed by dichotomy of hyphae of the infection cushion next to host cuticle. The pegs forcibly penetrated the cuticle and formed large, swollen, granular infection hyphae beneath the cuticle. Some infection hyphae penetrated the epidermis and cortical tissue directly, whereas others radiated from the infection site and entered the cortex. The subcuticular hyphae advanced ahead of the cortical hyphae and moved more rapidly up the stem than across. After 24 to 48 hr of incubation, the advancing subcuticular hyphae appeared to move into the cortex where further advancement up the stem occurred. Movement of infection hyphae through cortical tissue was always intercellular; however, branches of small hyphae from the infected infection hyphae soon ramified throughout the invaded tissue, frequently in an intracellular manner. In older portions of the lesion, the small hyphae (branching and coiling hyphae) evolved, revealing that the ramifying infection hyphae probably are of secondary importance in pathogenesis, as compared to the infected infection hyphae.

*Datura stramonium*, a local lesion host for certain isolates of cauliflower mosaic virus. M. Lung & T. P. Pirtone (Univ. Ky., Lexington). Necrotic local lesions were produced on leaves of *Datura stramonium* when inoculated with the Campbell, CM1841, KK cabbage, and New York 8153 isolates of cauliflower mosaic virus (CMV). Lesions developed 10-15 days after inoculation, and lesion numbers were directly proportional to virus concentration. No lesions or other symptoms were produced on *D. stramonium* inoculated with the cabbage B isolate. Electron-microscopic
examination of cells within lesions induced by the Campbell isolate revealed the presence of inclusion bodies in the cytoplasm. These contained compactly aggregated virus particles, and resembled the inclusions found in cells of novel Herpesvirus (Baltimore II type-1 like) systemically infected with CIMV. The morphology of the particles was typical of CIMV. Typical CIMV symptoms were produced on mustard inoculated with extracts from local lesions produced on D. stramonium leaves inoculated with purified CIMV from Coriandrum sativum. Thus the second reported host of CIMV outside the Cruciferae, and the only reported local lesion host for this virus.

_Fusarium wilt of susceptible and resistant tomato isolines: host colonization._ M. E. Mack & J. A. Vecch (ARS, USDA, Beltsville, MD). The effects of the site of inoculation and inoculum concentration on disease development were investigated in susceptible and resistant host isolines. No killing, no stunting, but a nearly constant amount of vascular browning were observed in the resistant host inoculated with 5 x 10⁹, 5 x 10⁸, 5 x 10⁷, and 5 x 10⁶ spores/ml. In contrast, the susceptible host exhibited progressive sensitivity to increased inoculum concentration. In both host cultivars, the amount of mycelial colonization of the hosts decreased as the distance from the point of inoculation increased, even though there was nearly systemically distribution of fungus spores through the axillary system. The site of inoculation greatly affected the extent to which the host was colonized by the pathogen. Both the susceptible and resistant hosts showed increased resistance to colonization as the site of inoculation moved up the plant axis. It was concluded that two morphologically distinct zones, tissue above the cotyledonary node and tissue below the cotyledonary node, should be considered separately in comparative physiological or biochemical studies of susceptibility and resistance to _Fusarium_ wilt of tomato.

_Germination self-inhibitor of sunflower and snapdragon rust uredospores._ V. Macko, R. C. Staples, & J. A. A. Reznick (Boyce Thompson Inst., Yonkers, N.Y.). The endogenous germination self-inhibitor of _Puccinia helianthi_ and _P. antirrhini_ uredospores has been identified as methyl 3,4-dimethoxyxycinnamate. Purification involved extraction with water, partition into ether, molecular distillation, and thin-layer (TLC) and gas chromatography (GLC). Analyses of the highly purified natural compound by infrared and mass spectrometry showed that it was identical to synthetic 3,4-dimethoxyxycinnamate. In the absence of both natural and synthetic compounds had identical RF values in two different TLC systems, and identical retention times in two different GLC systems. The ED₅₀ of the natural and synthetic inhibitor for spores of both rust species was 5 x 10⁻⁹ µg/ml.

_Properties of DNA bacteriophages from Agrobacterium tumefaciens._ R. J. Manasse & R. C. Staples (Boyce Thompson Inst., Yonkers, N.Y.). DNA bacteriophages from related pathogenic and nonpathogenic strains of _Agrobacterium tumefaciens_ (JVB7 and ITBTV6, respectively) were isolated. The phage isolates examined, two from each bacterial strain, were shown to be composed of hexagonal heads with long flexible tails containing a set of six appendages. Phage isolate LIIBV7-1 had a sedimentation constant (S₂₀,w) of 596, a molecular wt of 72 x 10⁶ daltons, and a buoyant density of 1.5110. The adsorption rate constant is 7.1 x 10⁻¹⁰ cm² sec⁻¹, and the eclipse and latent periods are 60-75 and 135-150 min, respectively. Similar data on both the physical and biological properties of three other phage isolates have been obtained.

_Artifact pseudoclonies on solid agar media inoculated with aster yellows or healthy plant material._ K. Marom, H. Hirumi, & B. P. Plavec-Banjac (Boyce Thompson Inst. Plant Res., Yonkers, N.Y.). Attempts to culture the mycoplasmalike presumptive agents of aster yellows and corn stunt in liquid or solid media have not yet been successful. Commercial mycoplasma media containing serum agar were inoculated with phloem tissue and phloem sap from aster yellows-diseased _Nicotiana rustica_ plants. Controls consisted of media inoculated with phloem tissue and sap from healthy plants inoculated agar serum. Nonliving artifacts, resembling pseudoclonies described by Laidlaw (1925) and Brown et al. (1940), mimicking mycoplasma colonies in time of appearance and in size developed in all plates. Careful microscopic observation showed that these artifacts did not have the round granular "fried egg" or the vacuolated morphology of mycoplasma colonies. The pseudoclonies "grew" and could be "subcultured", but did not consist of mycoplasma-like bodies, as was ascertained by electron microscopy. Pseudoclonies are probably crystalline formations of calcium and magnesium soaps.

_The application of an electronic cell counter to the study of growth of insect cell monolayers._ G. Martinez-Lorena & L. M. Black (Univ. Ill., Urbana). The growth of Agallia consticta cell monolayers (AC20 cell line) was studied by using the Coulter Cell Counter at various times to count the number of AC20 cells present in cultures under the influence of several factors. The insect cells were grown in Chia & Black's modification of Schneider's medium. A satisfactory seeding density for a 4-day interval between subcultures was between 2.5 x 10⁶ and 3.0 x 10⁶ cells per flask with a 25 cm² area for cell attachment. The optimum temperature for cell growth was estimated to be between 28 and 29°C. No differences in growth were detected at any temperatures from 27 to 30°C, but at 24°C the rate of growth was less, and at 33°C, cells died. A 10% concentration of fetal bovine serum was as good or better than 20%, and permitted a considerable economy in preparing the medium. Studies on the effect of infection by wound tumor virus (WTV) on the growth of AC20 provided definite evidence for cytopathic viral action. This effect was clearly observed after cells were subcultured, since the attachment of WTV-infected cells was about 25 to 30% less than that of cells free of WTV.

_Effect of carboxin on mitochondrial activities._ D. E. Mathie (Montana State Univ., Bozeman). Carboxin was tested for its effect on the activities of mitochondria from several fungi, pinto beans, and rat liver. Succinate oxidation in the mitochondria from the sensitive fungi, _Fusarium graminearum_ and _Fusarium maydis_, was inhibited by low concentrations of carboxin, the Km (inhibition constant) being 0.32 µM. The inhibition was noncompetitive. Succinate oxidation was also inhibited in the mitochondria from other sources, but not to such a great extent as those from _Fusarium maydis_. Carboxin had little effect on the oxidation of reduced nitroinAGED aminodisulfonic acid, nitro-blue tetrazolium, and did not appear to uncouple oxidative phosphorylation in the presence of succinate in tightly coupled rat liver mitochondria, but did decrease the respiratory control ratio. It is believed that carboxin inhibits mitochondrial respiration at or close to the site of succinate oxidation, and does not greatly affect the remaining potassium transport system or the cytochrome oxidation reactions.

_Influence of growth media on ultrastructure of hyphal tips of _Sclerotinia sclerotiorum._ D. P. Maxwell & P. H. Williams (Univ. Wis., Madison). The possible functional role of vesicles and crystal-containing microbodies (CB) in the production of oxalate, endopolygalacturonase (PG), _Lipase_ (Cxc) by _Sclerotinia sclerotiorum_ was investigated. The fungus was grown on mineral salts-yeast extract medium with glucose, glucose-succinate, sodium polypectate, or carboxymethylcellulose (CMC) as the carbon source. Relative Cx production on these media was <0.5, 3.4, 179, respectively, while relative PG production was <0.5, <1, 211, and 23, respectively. Total Cx accumulation was 1.1, 1,600, 851, and 641 µg/mg dry wt of hyphae, respectively. No unique organelle was detected in tips grown on media inducing high PG or Cx produc-
tion. More CB's (about 3 times) were present in tips grown on polypsectate and CMC than in tips grown on the other media. More membrane-bound vesicles (0.69-0.18 μm diameter) were associated with tips grown on the glass bead medium than with tips grown on the other media. These results do not support a functional role for CB's in either PG, Chx, or oxalate production, but they do show that the growth medium can influence the ultrastructure of hyphal tips.

Relationship of moisture and light to ascospore discharge by Mycosphaerella lirigula. R. F. McCoy & A. W. Dimock (Cornell Univ., Ithaca, N.Y.). Ascospore discharge by Mycosphaerella lirigula on diseased Chrysanthemum morifolium tissue was measured under controlled environmental conditions by means of a spore trap placed within a dew chamber. Moisture and, in some cases, light were found to be controlling factors regulating violent discharge. Liquid water was required for discharge in all cases. Ascospore discharge occurred within 30 min after the approximate dry period of the tissue with water. When the tissue was wetted by dew, ca. 2 to 2.5 hr. condensation was required before discharge began. The rate of condensation of dew was 4.5 mg/cm² per hr., indicating a requirement for ca. 0.1 mm of moisture accumulation to initiate discharge. With 3 isolates (from California and New York) discharge was noted only in darkness after switching on lights (combination incandescent cool-white fluorescent, 1,750 μW/cm²) prevented or stopped discharge even after a dew period which provided the requisite moisture had occurred in darkness. Ascospore discharge by a fourth isolate was not inhibited by light.

The distribution and fate of southern bean mosaic virus in the bean seed. J. G. McDonald & R. I. Hamilton (McGill University, Montreal, Quebec, Can.). Southern bean mosaic virus (SBMV) was recovered from mature embryos and seedcoats harvested from SBMV-infected Phaseolus vulgaris 'Logan'. However, when the embryo and seedcoats were treated to remove surface contamination, infectivity was eliminated from the embryos, but not from the seedcoats. Crude extracts of immature infected seedcoats were analyzed by sucrose density-gradient centrifugation, revealing a component, C₁, sedimenting slightly faster than the SBMV monomer, C. With drying of these seedcoats, the relative amount of C₁ increased, while that of C₁ declined. A third component, C₂, sedimenting faster than C₁, was also observed in these preparations. These components were not observed in noninoculated controls, and they reacted positively with SBMV antibody. The OD 260:280 nm ratios of both C₁ and C₂ (1.2–1.3) were considerably lower than that for SBMV monomer (1.5). With maturation of infected seedcoats, only a peak corresponding to C₁ was observed, suggesting that the maturation process caused a drastic alteration in the morphology of SBMV.

Effect of soil porosity, moisture, and temperature on diffusion of 1,3-Dichloropropane nematocides. M. V. McKey & I. J. Thomas (Univ. Cal., Riverside). Using techniques for direct chemical analyses (gas chromatography), the diffusion pattern of cis- and trans-1,3-Dichloropropane and 1,2-Dichloropropane was monitored under field and laboratory conditions. The three components of Telone diffuse at independent rates in soil. Increasing the soil moisture slowed the rate of diffusion. The diffusion patterns in a dry clay loam or sandy loam were similar. As the temperature increased, the rate of diffusion increased, depending upon the total volume of continuous soil air spaces. In soils with very little air space, the gas did not diffuse until pore spaces were sufficiently drained. In a very dry soil, adsorption to soil particles resulted in a reduction in the rate of diffusion. Increases In temperature from 5 to 25°C resulted in a faster rate of diffusion, while increased the rate of degradation of the 1,3-Dichloropropanes. In a clay loam soil, the maximum concentrations of trans-1,3-Dichloropropane were achieved 6 inches from the point of injection in 1.5 days when dry, 2.5 days when moist, and 9 days when soil was wet.

An electrophoretic examination of selected isoenzymes in strobilus of diseased citrus. C. W. Meister, R. M. Allen, & R. L. Caldwell (Univ. Ariz., Tucson). Enzymes from citrus leaves are being examined to find a simple, inexpensive, and rapid method for diagnosing stubborn disease. Leaves of healthy or diseased 5-year-old citrus trees were homogenized in Tris buffer, pH 7.5, and clarified by centrifugation. Protein was precipitated with trichloroacetic acid and measured by the Lowry method. Equal quantities of protein were layered on 8% or 5-15% gradient polyacrylamide gel columns and subjected to electrophoresis with Tris glycine buffer, pH 8.3. Twenty-eight enzymes have been identified by using specific reaction stains. Isoenzymes extracted from leaves of healthy plants are being compared with those from similar leaves of plants suffering from stubborn or psorosis diseases, drought, or mineral deficiencies. Preliminary results indicate that leaves from stubborn diseased plants have higher activities of peroxidase, leucine amino peptidase, and shikimate dehydrogenase than those from healthy plants. Lower activities of catalase and malate dehydrogenase have been consistently noted in stubborn-diseased plants.

Epiphytic pseudomonads on soybean in the field. T. W. Mew & B. W. Kennedy (Univ. Minn., St. Paul). A modified Kado's medium, selective for Pseudomonas, was used to detect populations of bacteria on leaves of three field-grown cultivars of soybean (Glycine max). By use of UV irradiation and leaf prints, we were able to demonstrate that the pseudomonads were in fact epiphytes, and that more saprophytic pseudomonads were on leaves early in the season than late. In the greenhouse, Phaseolus vulgaris 'Bush Blue Lake' was readily infected with isolates of epiphytic pseudomonads pathogenic to soybeans, and showed more or less typical water-soaked lesions, but there were no symptoms of systemic infection. As indicated by five major characters of Pseudomonas, it was evident that the pathogenic epiphytes were among the Group-I pseudomonads. It is concluded that epiphytic pseudomonads pathogenic to soybeans compose a unique group, but vary with the season and with genotype of the host.

Pathogenicity of Heterodera virginiae to Va. 312 dark-fried tobacco. L. I. Miller (Va. Polytech. Inst. & State Univ., Blacksburg). The pathogenic effect of the horse-nettle cyst nematode (Heterodera virginiae) to Va. 312 dark-fried tobacco was evaluated in microplots (metal cylinders 35 cm in diameter imbedded in soil to a depth of 90 cm and filled with methyl bromide-fumigated Woodstown fine sandy-loam soil). Cysts were mixed with the soil of six plots so that the infested soil contained an average of 30 eggs and/or larvae per g of air-dried soil. Six other plots were not infested. Single tobacco seedlings were transplanted to each microplot the 2nd week of May. Plants in the noninfested plots were harvested the 3rd week of August when they were judged to be of maximum quality and yield. Plants in infested plots were harvested the 3rd week of September to obtain the maximum yield. The diameter of the stalks (from node to node) of the stalks from noninfested plots was 37 mm, and 21 mm from infested plots; dry wt of roots from noninfested plots was 45 g, and 20 g from infested plots; and dry wt of all aboveground parts was 205 g from noninfested plots and 76 g from infested plots. Seed oil from infested plots averaged 98% yield and/or larvae per g of air-dried soil.

Some physical properties of elm mosaic virus. H. F. Molinue, G. L. McDaniell, & D. E. Maytew (Iowa State Univ., Ames). A virus isolated from Ulmus americana (Moline elm) in Iowa is serologically related to the type strain of elm mosaic virus (EMV). Shadowed, negatively stained, and fixed tissue preparations were examined by
electron microscopy. The virus is a small polyhedron. It was associated with necrotic ringspots in inoculated leaves of Chenopodium quinoa. Infectivity is associated with EMV which consists of three densities determined by analytical ultracentrifugation: top, middle, and bottom components with S20, w values of 45, 65, and 92, respectively. The middle component made up more than 60% of the total volume of purified preparations. The buoyant density of this component is 1.53 g/cm³. The isolate has a dilution end point of 10⁻⁶. 10⁻⁵ g/ml of EMV systemically infected C. quinoa leaves. The longevity in vitro of EMV is 4 days in neutral 0.01 M phosphate buffer; it is inactivated by heating for 10 min at 62°C but not at 60°C. The Iowa EMV isolate does not seem to differ markedly from others previously described.

**Preparation and storage of phytopathogenic bacteria in liquid nitrogen.** L. W. Moore & R. V. Carlson (Oregon State Univ., Corvallis). Liquid nitrogen was evaluated as a method of storage for species of Agrobacterium, Erwinia, Pseudomonas, and Xanthomonas. Bacteria in exponential growth were harvested (10⁸ to 10⁹ colony-forming units/ml) from a liquid shake-culture consisting of 4% yeast extract, 20% dextrose, 0.4% peptone, and 0.5% (NH₄)₂SO₄ at pH 7. This medium plus 1.5% agar was used as a viability counts (spread-plate method) before and after freezing. The harvested bacteria were frozen directly in their growth medium, or centrifuged and the pellet resuspended in the following media: fresh medium, fresh medium containing 1% dimethyl sulfoxide, 12% skim milk, or distilled water. Glass ampules containing 0.5-ml aliquots of a bacterial suspension were heat-sealed and plunged into liquid nitrogen. Recovery of viable bacteria was periodically determined over a 2-month period by thawing the ampule contents for 12 sec at 40 ± 1°C, diluting, and plating. Optimum recovery from a given freezing medium varied among the nine species of the four genera; recovery percentages were within a range of 65 to 118% of the initial viable counts.

**Protoplast enzyme production by Verticillium albo-atrum.** H. W. Muesell & Blanchie Strouse (Boyce Thompson Inst., Ithaca, N.Y.). Twelve isolates of Verticillium albo-atrum from cotton produced extracellular protoplastic enzymes when grown using soluble or insoluble proteins as a sole carbon source. Protolytic enzymes were not detected when glucose was used as a sole carbon source, but low levels of glucose with proteins enhanced enzyme production 3- to 6-fold over that induced by proteins alone. Certain amino acids nor the polypeptide glutathione induced protoplastic enzymes when used as the only carbon source or with glucose. Protoplastic activity had a pH optimum of 8.5, and was stable in solution for several months under aseptic conditions. Polyalactonaractone activity was higher on media that induced protoplastic enzymes than on comparable media containing only glucose. Cellulase activity, on the other hand, was uniformly low on glucose and protein-based media.

**Etiology of Monochaetia mail in apple.** R. Naik & D. Powell (Univ. Ill., Urbana). Monochaetia mail has been known since 1900, but its etiology in apples has not been studied. In the current studies, the organism infected twigs through ovipositor wounds caused by cicada, and produced a canker lesion occurred only through wounded tissue. Cankers were produced by inoculation of mature apple tree limbs. Inoculation of fruit produced lesions which were convex, light brown, and enlarged with age. Aecervi, which formed within 8-10 days after inoculation, produced characteristic conditions in which were several-celled and dark, with hyaline-pointed end cells and a single apical appendage. Similar fruiting structures were found at the margins of limb cankers. The fungus grew well on lima bean agar. Optimum temperature for growth in vitro was between 25 and 30°C, and optimum pH was 6 with extremes of 3 to 9. Susceptibility of inoculated fruit was directly correlated with increase in tissue pH and warm temperature of ca. 30°C. The incubation period was shorter in the field on mature fruit (4 days) than on 20-day younger fruit (5-7 days).

**Pseudomonas glycinea inhibits germination of soybean seed.** J. F. Nicholson & J. B. Sinclair (Univ. Ill., Urbana). A new role of Pseudomonas glycinea in soybean pathology was studied. Two isolates of an internally seedborne bacterium in soybean (Glycine max) inhibited the germination of Lee 68 and Asmoy soybean seed. In culture, one isolate had a smooth surface and margin, whereas the other had a rough surface and margin. Both isolates were identified as P. glycinea through a series of standard biochemical tests compared to stock cultures of P. tabaci and P. glycinea. Our isolates and the stock isolate of P. glycinea did not grow on Kado's selective medium D4 for Pseudomonads. When suspensions of three P. glycinea isolates were infiltrated by vacuum into sterilized Asmoy seed, germination was significantly inhibited by the rough-margined and stock culture isolates to 38 and 65%, respectively, whereas the smooth-margined isolate had normal germination (84%) compared with the control (88%). The bacterium was isolated from 17 seed lots of Lee 68 soybean collected from five states. The occurrence in individual seed lots was as high as 64% and as low as 3%.

**The effect of β-sitosterol on the activity of certain enzymes of Phytophthora cactorum.** C. Norman & R. A. Calderone (Washington State Univ., Morgantown). β-Sitosterol (10 mg/liter) increased activity of aldolase, r-glutamic acid transaminase, and glucose-6-phosphate dehydrogenase of Phytophthora cactorum when added to cultures growing in a liquid glucose-asparagine medium at 25°C. The increase in activity generally occurred 48-72 hr after the addition of the sterol to 5-day-old cultures. Specific activities (units of activity/mg protein of cell-free extract) of both aldolase and transaminase was 1.6-1.7 times greater in cultures containing β-sitosterol than in the controls (no sterol). Likewise, the specific activity of glucose-6-phosphate dehydrogenase was 1.3-1.4 times greater than the control. The sterol-induced increases in enzyme activity paralleled increases in growth (mg dry wt) of the fungus over control cultures.

**The influence of exogenous cytokinins on the development of the bacterial hypersensitive reaction.** A. Novacky (Univ. Mo., Columbia). When tobacco leaves were treated with 5-10 μg/ml of cytokinins (kinetin, 6-β-(γ-dimethylamino propyl)purine, and benzylaminopurine), 48 hr prior to infiltration with 5 x 10⁶ or 10⁷ cells/ml of Pseudomonas pisi, development of the hypersensitive reaction (HR) was overcome. The initial appearance of HR was identical in leaves infiltrated with water or with cytokinins. In the former, the HR progressed rapidly from the initial shininess and faint wrinkling of lower leaf surface to necrosis. In the latter, the initial reaction did not progress beyond the initial symptoms. Bacterial growth was not affected by cytokinins. Age of the treated leaves was critical in demonstrating the cytokinin effect on HR. Only fully expanded leaves showed this phenomenon. Extending the pretreatment period with cytokinin to 72 hr with an additional one or two applications of cytokinins prevented HR induction by 10⁶ cells/ml. Other consequences of treatment of the tobacco with cytokinins were an increased resistance to water stress and a reduction in electrolyte leakage when compared with water-treated controls. Cytokinin-treated tissues also released lower peptidase activity. From these experiments we have postulated that cytokinins in some way preserve the integrity of vital cellular membranes which are generally altered as a consequence of HR.

**Remission of symptoms of pear decline in pear and peach X-disease in peach after treatment with a tetra-cycline.** G. Nyland (Univ. Cal., Davis). Recent reports indicate that mycoplasma-like organisms infect pear decline
and peach X-disease. One tree from each pair of 15 pairs of Comice and 15 pairs of Bartlett pear trees was treated with 3 liters of tetracycline hydrochloride solution (100 µg/ml) in May 1969. The solution was fed into the xylem of each tree by gravity via plastic tube connectors and plastic tubing attached to reservoirs suspended 3-4 feet above 4 equally spaced holes drilled into the trunk. All trees had shown symptoms of the leaf curl type of pear decline during each of the previous 2 seasons. Symptoms of leaf curl developed in less than any of the 30 trees treated with tetracycline hydrochloride. Typical symptoms of leaf curl developed in 29 of 30 untreated trees. Four peach trees showing X-disease symptoms were treated with 2 liters of tetracycline hydrochloride solution (34 µg/ml) fed into diseased branches in September 1969 plus 2 additional liters (1 µg/ml) fed in April 1970. Only untreated trees showed symptoms of X-disease in 1970. The treated ones remained symptomless.

Biological activity of conversion products of benomyl. J. M. Ogasawa, Elaine Bose, B. T. Manji, E. R. White, & W. Killoore (Univ. Calif., Davis). A mixture of 8 oz of 50% benomyl and Bordeaux mixture 10-10-10 (foliage dosages) formed a compound identified as 5-triazinooxy 1,10-dihydro-2-benzimidazole cyanamide. Technical benomyl placed in contact with water adjustment to pH 12 with NaOH formed the same compound. The spectrum of activity of STB on test fungi was the same as that of benomyl. For total suppression of Monilinia laxa on potato-dextrose agar, 10 to 50 ppm of STB were required, whereas for Botrytis cinerea and Magnaporthe (2-benzimidazole cyanamide), only 0.1 ppm was required. Using Botrytis cinerea as a test pathogen, 30 detached Drake almond blossoms/treatment were sprayed with each chemical, inoculated with a spore suspension, and incubated at 20°C and over 95% RH. At concentrations of 1 and 10 ppm, STB reduced infection by 10 and 90%; and benomyl, 23% and 80%, respectively. Seven single tree replications of Drake almond trees sprayed with field dosages for the control of M. laxa at the pink bud stage of bloom showed an average of 60, 23, 90, and 593 blighted shoots for STB, benomyl, benomyl plus Bordeaux mixture, and control, respectively. STB was used at the same concentration as benomyl.

Host specificity of Polyergus betulinus. R. L. Payne & W. W. Johnson (Pa. State Univ. Park). Polyergus betulinus is restricted to Betula spp. in nature. Our previous studies of the mechanism of this host specificity showed that spores of the fungus germinated on host and nonhost species. Sawdust and heartwood of 40 trees each of Betula lenta, Picea abies, and Pinus sibirica were placed with polythene bags or glass containers of P. betulinus. The fungus could not be recovered from Betula or Pinus 6 months later; it was recovered from P. abies after 6 months from four wounds on three trees inoculated with spores or grain inoculum and after 1 year from one heartwood wound inoculated with spores. In the latter, the fungus was recovered from the margins of a pocket of yellowish-brown decay 8 x 3 cm in diameter. Identification of all isolates was confirmed by hyphal anastomoses with known isolates of P. betulinus. Host specificity of P. betulinus can be controlled by factors other than those acting to prevent spore germination and subsequent colonization. The multitude of competing microorganisms in wounds from which P. betulinus could not be recovered suggests that host specificity may be due partially to factors which control growth of these competitors.

Enhancement of Pseudomonas phaseolicola infection of bean by 3'-5' cyclic AMP. N. J. Panopoulos & M. N. Schrot (Univ. Calif., Berkeley). Infection of bean by Pseudomonas phaseolicola is noted as a weakening, suggesting catalase repressor. Bacterial inocula (10^3 cells/ml) were suspended in 2 to 5 mM solutions of glucose, glycerol, or other sugars, and infiltrated into the intercellular spaces of primary leaves of red kidney beans. The number of lesions was reduced by 50 to 60% in the presence of glucose or glycerol, compared to water and osmotic controls. Since catalase repression of inducible enzyme formation in bacteria has been experimentally related to lowering of the intracellular level of 3'-5' cyclic AMP, it was examined whether infection was affected by this or other cyclic nucleotides. 3',5'-Cyclic AMP, 2',3'-cyclic AMP, and 3',5'-cyclic UMP at concentrations ranging from 2 to 5 mM (pH adjusted to 6.8-7.0 with NaOH) enhanced infection 1- to 5-fold over water controls. In contrast, ATP was the biochemical precursor of 3',5'-AMP, inhibited infection by 50 to 70% under these conditions.

Plant water stress and the development of Fusarium rot in sweet wheat. R. J. Parton, G. C. Mattos (ARS, USDA, NATO Fellow, Pullman, Wash.). Foot rot of winter wheat (Tritium aestivum) caused by Fusarium culmorum is most severe in low rainfall areas in fields with high N fertility and early fall-seeded to semidwarf cultivars with high tillering capacity. Such conditions hasten extraction of the limited soil water and onset of plant water stress. Soil water depletion rates during May in field plots of Nuggines and Moro were generally greater with higher N (up to 200 lb/acre) and narrower row spacings (12 vs. 24 inches). Leaf water potentials measured by thermocouple psychrometers for biological and chemical treatments (MC 20, 2-benzimidazole cyanamide, 2-benzamide) were between -10 to -27 bars in early May to -33 to -35 bars or less in mid-June. Potentials were generally lower, and onset of stress was earliest in plots with high N and narrow row spacings. Infection of crowns by F. culmorum was uniform through all treatments on 1 April, and remained latent through May, after which severe basal culm decay began occurring under greatest water stress. Five cultivars showed a positive correlation between onset of stress, onset of severe foot rot, and yield reduction, suggesting that disease control may be possible by breeding for cultivars having high water-use efficiency. A sixth variety that showed early stress had negligible foot rot and yielded well, suggesting that stress alone could not account for the yield reduction.

Exogenous carbon and nitrogen requirements for germination of conidia by Aspergillus flavus. T. Pass & G. J. Griffin (Va. Polytech. Inst., Va. State Univ., Blacksburg). In a phosphate-buffered (pH 5.7), inorganic salts solution (B), almost full dependence on exogenous carbon and partial dependence on exogenous nitrogen was found for complete germination of washed conidia by A. flavus over a range of conidial concentrations. At equivalent carbon, an amino-acid mixture supported higher germination than a sugar-organic acid-alcohol mixture plus NH4Cl, proline or serine alone supported higher germination than several other single amino acids or single sugars plus NH4Cl. Glucose plus NH4Cl was the most stimulatory of the latter. Germination was more dependent on exogenous nitrogen as the conidial density increased from 10^6 to 10^9 conidia/ml. At equivalent nitrogen, proline alone or an amino acid mixture supported higher germination than NH4+ plus NO3- or several other single amino acids tested in B solution plus glucose. Percentage germination decreased as the conidial density decreased from 10^6 to 10^2 conidia/ml when glucose or glucose plus NH4Cl were supplied at constant factors. However, little or no decrease in percentage germination was observed in glucose plus an amino acid mixture, and only a slight decrease was observed in glucose plus peptone. Germination was best at 30-35°C, while a broad pH optimum of 3.0-8.0 was found.

Role of benzyloxyloxyacetylene in the resistance of papaya fruit to postharvest fungal diseases. S. S. Patil, C. S. Tang, & J. E. Hunter (Univ. Hawaii, Honolulu, & Hilo). The level of benzyloxyloxyacetylene (BITC), a compound ubiquitously in papaya fruit, decreases steadily from harvest to harvested fruit, and this decrease is inversely correlated with the incidence of postharvest diseases. In vitro studies on sensitivity of papaya pathogens to BITC showed that Rhizopus stolonifer and Phytophthora palmivora were completely inhibited at 40 ppm of BITC. At the same con-
centron, *Stemphyllum* sp. and *Phomopsis* sp. were inhibited ca. 20%, and colonies of *Colletotrichum gloeosporioides* and *Fusarium oxysporum* were inhibited 30-40%. Control experiments were conducted on mature-green fruits which were wounded inoculated with *K. stolonifer*, incubated for 24-48 hr at 25°C then sealed for 6 days in plastic boxes with tape. Distilled water was applied to the wounded area, and the boxes were placed in a constant humidity incubator at 100% relative humidity. The results showed that wounded inoculation with *K. stolonifer* did not significantly increase the incidence of infections, indicating that invasion by the pathogen is not required for its production. The inhibitory effects of the compound are destroyed by heating and by acid hydrolysis.

A medium for the selective growth of *Monilinia* species. D. J. PHILLIPS (USDA, Fosno, Cal.). A medium containing 1,000 ppm PCNB (pentachloronitrobenzene), 20 g strained canned peaches, 100 ppm neomycin, and 500 ppm streptomycin was developed for the selective growth of *Monilinia fructicola* and *M. laxa*. While allowing rapid growth of *Monilinia* sp., the medium represses the growth of *Mucorales* and restricts other contaminating fungi and bacteria to discrete colonies. Counts of viable spores made with a hemocytometer correspond to plate counts on the medium over a dilution range from 1:10 to 1:10,000 (1 = 4600 spores/ml). Counts of colonies of the medium served to measure the inoculum density when spores were made up a very small portion of the total population. For example, a 6-month-old peach nummy had 25,000 viable spores in March 1971 when the total *Monilinia* spore population was estimated with the hemocytometer to be 8,500×10^4. Washing freshly harvested peach, nectarine, and plum fruits in 1970 showed that the inoculum density ranged from 300 to 27,000 viable spores/fruit, which caused 1 to 50% decay after a simulated storage and marketing period of 24 hr at 2°C and 4 days at 25°C.

A disease of soybean caused by *Neocosmospora vasinfecta*. D. V. PHILLIPS (Univ. Ga., Ga. Exp. Sta., Experiment). *Neocosmospora vasinfecta* was isolated from several soybean (*Glycine max*) plants with discoloration in the pith and xylem of the stem. Soybean plants inoculated with *N. vasinfecta* developed internal stem discoloration, and the fungus was reisolated. Single ascospore isolates were used in subsequent pathogenicity tests. Fifteen soybean cultivars were susceptible. Symptom expression was greater at 27°C than at 21 or 15°C. Stem puncture inoculations resulted in a much higher percentage of diseased plants than did inoculation of injured roots. The length of internal stem browning in greenhouse-grown plants 4 to 7 weeks after inoculation did not exceed 1 cm. In field-grown plants inoculated in June and examined in October, internal browning extended a maximum of 47 cm above the inoculation point. *Neocosmospora vasinfecta* had little effect on plant height, and did not kill the soybean plant. Internal stem discoloration caused by *N. vasinfecta* in soybean plants is similar to that of brown stem rot caused by *Cephalosporium gregarium*, but *N. vasinfecta* hyphae are confined to the pith, whereas *C. gregarium* hyphae are found primarily in the xylem of young plants. *Neocosmospora vasinfecta* caused similar symptoms in cultivars of *Phaseolus vulgaris* and *Vigna sinensis*, but not in 11 other crop plants.

The differentiation between bacterial hypersensitive reactions and pathogenesis by the use of cycloheximide. Y. PRZYBYLAK & A. NOWACKY (Univ. Mo., Columbia). Injection of cycloheximide (70 mg/ml) into tobacco leaf tissue at or up to 3 hr after inoculation with *Pseudomonas pisi* (10^8 cells/ml) delayed the hypersensitive reaction (HR) for 18 hr. In vitro and in vivo experiments showed no inhibition of cell division or growth by bacterial multiplication. On the contrary, multiplication was higher on CHI-treated tissues. These data suggest that the delay caused by CHI is a result of its effect on the plant tissue; hence, HR is a plant response to the incompatible bacteria. Analogous delay of the HR symptoms was observed with *Erwinia amylovora*, another incompatible pathogen in to-
bacco. Since CHL did not delay symptoms in tobacco inoculated with the compatible pathogen Pseudomonas tabaci, it would appear that HR mediated by this pathogen is a distinctly different phenomenon. It was found that CHL strongly inhibited 14C leucine incorporation into tobacco leaf proteins, which suggests that HR is associated with plant protein synthesis. The labeled soluble proteins from bacterially infected tissue were separated by isoelectric focusing in polyacrylamide gel. The protein patterns were similar in controls and in tissues undergoing HR. However, the level of protein biosynthesis was increased in tissue undergoing HR. This increase is in large part due to plant biosynthesis.

Effect of EDTA on transmission of purified cucumber mosaic virus by aphids. T. P. PIRONE (Univ. Ky., Lexington). Cucumber mosaic virus (CMV) (1 mg/ml in 5 mm borate buffer, pH 9), acquired by aphids (Myzus persic Crus through a Parafilm membrane, was transmitted by the aphids to tobacco seedlings with an efficiency of 20-50%. Aphids did not transmit CMV when the virus suspensions contained 1 umole/ml EDTA at pH 9. The addition of CaCl2 (1 mmole/ml) to virus preparations which contained 1 umole/ml EDTA restored aphid transmissibility of the virus. In some experiments, the presence of 0.1 umole/ml CaCl2 in virus suspensions not pretreated with EDTA resulted in levels of aphid transmission which were higher than those obtained with preparations which did not contain CaCl2. Infectivity of the virus, as measured by local lesions on tobacco, was neither decreased by the presence of 1 umole/ml EDTA, nor increased by the presence of 0.1 umole/ml CaCl2.

Characterization of barley stripe mosaic virus replicative form RNA. D. R. PEARSON (Univ. Neb., Lincoln). An apparent replicative form (RF) of barley stripe mosaic virus RNA was produced by in vivo 32P-labeling of the inoculated leaf one of Black Hulls barley. The RF was resistant to deoxyribonuclease and to change in sedimentation coefficient upon formaldehyde treatment. The RF was resistant to ribonuclease in high salt conditions, susceptible to the enzyme in low salt conditions, and was alkali-labile. Estimated molecular wt of single-stranded viral RNA is about 8.5 x 10^6, based on sedimentation after formaldehyde treatment. The sedimentation coefficient of the RF was 12.75, which gives a molecular wt of about 1.7 x 10^6 by Studier's formula. Buoyant density of the RF in CsSO4 was very close to 1.600. Maximum 32P incorporation into RF occurred during a 72-96 hr postinoculation labeling period, or shortly before the maximum level of single-stranded RNA was apparent in the infected tissue.

Reduction of aflatoxin production in peanuts in the presence of dimethyl sulfoxide (DMSO) on the production of aflatoxin by Aspergillus flavus (ATCC 2221) in peanuts was determined. Fifty g of dry peanuts, cultivar Early Runner, were soaked for 30 min in distilled water containing 0.0, 0.6, 1.2, 2.5, 5.0, 10.0, or 20.0% DMSO. After soaking, they were autoclaved and inoculated with a spore suspension of A. flavus. The cultures were incubated for 7 days at 24°C, then extracted with 70% aqueous acetone and spotted on thin-layer chromatographic plates and developed in chloroform-methanol (97:3) solvent. Developed plates were examined under UV irradiation, and relative amounts of aflatoxin were determined by amounts of fluorescence. The total amount of aflatoxin was determined spectrophotometrically at 365 nm and compared to aflatoxin standards. Aflatoxin production in culture containing 0.6 and 1.2% DMSO was comparable to the controls. However, at concentrations above 1.5%, aflatoxin production decreased rapidly. At 20.0% DMSO, little or no fungal growth or aflatoxin was detected. These results were verified by a bioassay using chick embryos.

Tobacco etch virus tolerance in tomato. W. B. RAYMER (Campbell Inst. Agr. Res., Cinnaminson, N J). A New Jersey isolate of tobacco etch virus (TEV) was used to sap-inoculate 20 plants each of 288 tobacco cultivars and P. L. lines. These accessions were evaluated for tolerance to TEV by means of a 0-5 rating system based on severity of mottling, rugosity, and leaf distortion. Only 5 of 125 Lycopersicon esculentum lines were considered highly tolerant, with ratings of 2 or less, whereas 9 of 17 L. hirsutum, 25 of 56 L. peruvianum, and 6 of 73 L. pimpinellifolium lines were in this category. All 7 lines of L. glabratum and 1 each of L. cheesemani f. minor and Solanum pennellii were intolerant, with ratings of 3 or higher. No resistance to infection by sap-inoculation was found in these materials, as TEV was recovered from all symptomless plants. P. L. 125955 remained symptomless for 4 weeks following inoculation, and was the most tolerant L. esculentum tested. P. L. 127827, 7 other L. hirsutum lines, and P. L. 247057 (L. peruvianum) also remained symptomless. Of the P. L. lines reported to be resistant to TEV in Florida, only P. L. 27527 was highly tolerant in these tests, with a 0 rating. P. L. lines 166989 and 183692 (L. esculentum) were given an intolerant rating of 4, and P. L. 134417 (L. hirsutum f. glabratum) was given an intermediate 3 rating.

An improved method for purification of wound tumor virus. D. V. R. REDDY & J. A. LESKAW (Univ. Ill., Urbana). The yield of wound tumor virus (WTV) purified by earlier methods was 2 to 3% of the virus contained in starting material. Moreover, the specific infectivity of the virus was only 1% of the initial virus. A simpler method of purifying WTV was developed using polyethylene glycol (PEG) 6000. Root tumors from sweet clover were ground in buffer first used for WTV by I. Kimura (0.1 m histidine and 0.01 m MgCl2, pH 7.0). After low-speed centrifugation, the virus was precipitated using 0.3 m NaCl and 4% PEG. After standing 2 hr at 4°C, the precipitate was concentrated by low-speed centrifugation, further purification was achieved by a single equilibrium zonal density-gradient centrifugation in sucrose. Infectivity was measured by inoculating monolayers of vector cells (AC20) and counting the foot of infection after fluorescent antibody staining. The virus concentration in purified preparations was obtained from the optical density of their RNA, measured after its extraction with 1 n HCl. Purity of the virus was ascertained from determinations of protein/RNA ratios and of base ratios of RNA. According to these criteria, no impurities were detected in the preparations. About 50% of the virus present in the starting tumor material could be recovered, and the purified virus retained about 90% of the original specific infectivity.

Induction of the soybean phytoalexin hydroxyphaseolin with fungicides. J. J. REELLY & W. L. KLAMANN (Univ. Maryland, College Park). Technical-grade fungicides were tested for their ability to induce hydroxyphaseolin (HP) in the soybean cultivar Harosoy 63. Cotyledons and hypocotyls were removed from 6-day-old soybean plants and incubated at 22°C in petri dishes each containing 7 ml of various concentrations of fungicides. Maneb, sodium dimethyldithiocarbamate dihydrate, nabam, Terrazole (5-ethyl-3-trichlormethyl-1,2,4-thiadiazole), Mersolite-19 (phenyl mercuric salicylate), and benomyl induce HP as determined by bioassay, UV spectroscopy, and gas liquid chromatography. Zineb, ferbam, and chlorothalonil (benomyl-chlorothalonilbenzenzene) did not induce detectable amounts of HP. The amine degradation products of benomyl and maneb were potent inducers. In a subsequent study of six amines, isobutylamine was the most effective inducer of HP. The MBC (minimum bactericidal concentration) of benomyl was determined. Induction of the benomyl molecule is reported to be as fungistatic as benomyl in vitro; however, MBC did not induce HP.

Oospore production, viability, and germination in relation to establishment of F. cultures of Phytophthora infestans. O. K. RIBEIRO, M. E. GALLEGGY, & R. J. YOUNG
Pathogenicity of Pseudomonas syringae on Phaseolus vulgaris as influenced by bacterial isolate source. S. M. SAAD and D. J. HACEDORN (Univ. Wisc., Madison). Pathogenicity of 30 Pseudomonas syringae isolates from 10 hosts was compared on Tenderwheat bean in the greenhouse, using inoculum levels of 107 and 108 cells/ml. Bacterial suspensions were sprayed on both surfaces of the first trifoliate leaf. All bean isolates at both inoculum levels induced typical olive-green, water-soaked lesions within 3-4 days. These lesions soon became necrotic (dark brown), with marginal chlorosis. Isolates from pear, apple, sour and sweet cherry, lilac, peach, plum, walnut, and soybean produced only tiny, dark necrotic flecks within 24 to 48 hr with inoculum containing 108 cells/ml. No macroscopic symptoms were obtained with the lower level of inoculum. Different isolates from the same host showed minor pathogenic differences. Growth characteristics of several isolates were studied in bean pods. In the compatible combination, the bacteria multiplied logarithmically for 4 days, inducing water-soaked lesions without brown necrosis. Logarithmic growth of incompatible isolates from pear, lilac, and sour cherry terminated after 2 days and induced brown, sunken necrotic lesions which fluoresced when exposed to UV light. These studies indicate a positive relationship of high pathogenicity to bean with P. syringae isolates from bean alone.

Remote sensing of southern corn leaf blight. G. R. SATR, G. SUTS, & A. H. ELLINGHOR (Mich. State Univ., East Lansing, Univ. Mich., Ann Arbor). Corn plants of the hybrid W64A X OH 43 with Normal "N" or Texas male sterile "T" cytoplasts were inoculated with Helminthosporium maydis race "T". The reflectance of leaves at wavelengths between 0.35 and 2.6 μm of inoculated and non-inoculated plants was determined with a Beckman recording spectrophotometer. Greater reflectance, primarily at the chlorophyll and water-absorbing wavelengths, was obtained from leaf areas with lesions than from healthy leaves. Portion of leaves not having lesions was less affected. A greater difference in reflectance in the water-absorbing wavelengths was observed if the inoculated corn plants had "T" cytoplasm than if the plants had "N" cytoplasm. The earliest detection of a difference in reflection between diseased and healthy tissues was ca. 36 hr after inoculation.

Identification, characterization, and transmissibility of a virus infecting St. Augustine grass in Florida. J. L. SALADINI & F. W. ZETTLER (Univ. Fla., Gainesville). A virus of St. Augustine grass isolated in 1963 by E. S. Field was identified as st. Augustinella mosaic virus RuBEC (SMV-E). It induced discrete localized lesions in Atlas sorghum, and a local and systemic necrosis followed by recovery in CP 31-294 and CP 31-588 sugarcane; it did not infect Johnson grass. Of 138 particles measured in leaf extracts, 89 were 675-782 nm long with a main maximum at 711 nm. Pinwheel, circle, bundle, and laminated aggregate inclusions occurred in diseased corn leaves. SMV-E was transmitted from corn to corn by Schizaphis graminum aphids in a stilt-borne manner. St. Augustine grass was more resistant than was corn to infection by mechanical inoculations to adaxial leaf surfaces. In trials using identical inocula, only 10 of 200 St. Augustine grass plants became infected as compared to 195 of 200 for corn. Once infected, however, symptoms persisted in St. Augustine grass plants, and SMV-E was always transmitted leading to a reduction in yield of corn. Scanning electron micrographs revealed extensive wax deposits on adaxial but not on abaxial leaf surfaces of St. Augustine grass.

Protein content of tobacco roots following heat treatment and inoculation with Phytophthora parasitica var. nicotianae. G. E. SANDEN & L. D. MOORE (Va. Polytech. Inst., Va. State Univ., Blacksburg). Roots of intact plants of susceptible and resistant cultivars of flue-cured tobacco, Nicotiana tabacum, were subjected to the following treatments: (i) roots were heated for 30 min at 95°C; (ii) roots were not heated; (iii) roots were inoculated with P. parasitica var. nicotianae; (iv) roots were not inoculated. The results indicated that the protein content of tobacco roots can be increased by both heat and inoculation with P. parasitica var. nicotianae.

Temperature and moisture requirements for infection of detached pineapple inoculums by pink disease bacteria. K. E. ROSENBACH & Joan PREPENTER (Pineapple Res. Inst., Wahilla, Hawaii). Studies on the disease of pineapple, caused by various strains of acetic acid bacteria (Acetomonas sp.), have been limited because the disease occurs sporadically and little is known about environmental conditions necessary for flower infections. To determine conditions required for flower infections, pineapple inoculums were detached and bacterial suspension was applied to the open flowers. The inoculums were then subjected to various temperature and moisture regimes for time periods of up to 72 hr. Isolation of the bacterium in nectary gland and placental tissue was interpreted as positive infection, although a pathogenic relationship was not verified. At high humidity (polyethylene bag), 92% of the nectary glands were infected at 18°C for 48 hr, whereas only 47% occurred at 29°C. Infections decreased 4-fold at both temperatures at low humidity (no polyethylene bag). Only slight differences of infection occurred with an alternating 29°C day and 18°C night temperature when compared to a constant 18°C temperature (high humidity, 72%). Pathogen recovery from nectary glands was consistently higher than from placental tissue. When the length of time at high humidity was varied following inoculation, maximum infections occurred at 6 hr.
trols were untreated. The root tissue was assayed immediately and 6 days after treatment for total protein content by the Lowry method. Extracts from the susceptible cultivar, Virginia Gold, which were inoculated only, heated only, or heated and inoculated, showed a decrease in total proteins as compared to extracts from roots which received no treatment. Extracts from the resistant cultivar, Coker 187, showed a decrease in total protein levels from those roots which were heated only or inoculated only as compared with unheated checks, while roots which were both heated and inoculated showed an increase. The protein content of heated-only and inoculated-only roots was slightly lower than that of the resistant cultivar than for the susceptible cultivar. The low concentration of protein in roots of Virginia Gold, which were both heated and inoculated, may have been due to the degradation of the roots.

**Impaired amino acid biosynthesis in phytopathogenic pseudomonads. D. C. Sands & M. L. Zipper (Conn. Agr. Exp. Sta., New Haven).** Phytopathogenic fluorescent pseudomonads related to *Pseudomonas syringae* differ distinctly in amino acid metabolism from saprophytic members of the group. The pathogens cannot use a number of amino acids as carbon or energy sources. The saprophytes can. A number of amino acids were toxic to 11 strains of pathogens but not to saprophytes. t-Homoserine was toxic to all pathogens. Threonine, valine, alanine, glycine, and a combination of phenylalanine plus tyrosine were strongly inhibitory to the growth of more of the pseudomonads. In some cases, the inhibitory effect was reversed by the presence of certain amino acids in the same biosynthetic pathway. Studies with mutants of saprophytes have shown that toxicities and reversals are indicative of lesions in the regulation of the branched pathways of biosynthesis of amino acids. Perhaps the lesions result from selection, due to the high levels of amino acids in plants. These lesions may be linked to toxin production and host specificity factors.

**Survival of Erwinia rubrfaciens in exudate on diseased walnut trees. N. W. Schaad (Univ. Cal., Davis).** *Erwinia rubrfaciens* occurs in large numbers in a slimy substance which exudes through cracks and accumulates on the bark of infected walnut trees. To determine the longevity of the bacteria in dried gummy exudate, samples of exudate were collected at various times after injury. After 62 days, numerous bacteria were still viable in exudate protected from further surface replenishment but left in its natural environment. For example, the original concentration of 1.4 x 10^9 viable cells/mg decreased only to 1.0 x 10^7 after 41 days; after 41 days, when expressed as a survival rate, the data show that the half-life of the bacteria was 5.7 days for the first 41 days. When exudate was sampled after having been placed on a healthy tree or stored in the laboratory (room temperature), the results were somewhat different. For example, it took 5.6 days for half the bacteria to die after 48 days in the exudate placed at room temperature, but only 3.8 days for bacteria in exudate transferred to a healthy tree. It is concluded that *E. rubrfaciens* can survive for at least 123 days in exudate on the bark of diseased trees.

**The influence of dew and atmospheric moisture periods on infection of slash pine seedlings by Cronartium fasiiforme. R. A. Schiefer (Univ. Fla., Gainesville).** Two-month-old seedlings of slash pine (Pinus elliottii var. elliottii) were exposed for 1-2 hr in a dew chamber beneath trays of oak leaves bearing generating tefla of *Cronartium fasiiforme*. Inoculated seedlings (50/treatment) were placed in the environmental conditions described below and planted in the nursery, where the percentage of seedlings with galls was determined after 9 months. Dew periods (x) of 0, 1, 2, 4, 5, 6, 8, 10, 12, 16, and 24 hr resulted in disease percentages of 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, and 0, respectively. Per cent disease (y) was related to positive exponential equations, x(t) in a direct linear manner (y < 4, y = 0; x1 > 5, y = 15.71 + 5.20x1; and x1 > 2, y = 0; x1 > 4, y = 5.16 + 2.96x1). The significant correlation coefficients (r) were 0.67 and 0.751, respectively. Postinoculation drying periods (t) of 0, 1, 2, 3, 4, 5, 12, 16, 24 hr prior to inoculation at favorable moisture conditions resulted in disease percentages of 62, 63, 70, 74, 53, and 51, respectively (y = 58.0 - 0.408x, and r = 0.264).

**Halo blight of mung bean induced by a new strain of Pseudomonas phaseolica L. E. Schonbeck, H. A. H. Hoitink, & M. E. Klaert (Ohio Agr. Res. Dev. Cent. Wooster).** Introduction of mung bean (*Phaseolus aureus*) has not been successful in Ohio because of severe losses from halo blight. The pathogen isolated was similar to both *Pseudomonas phaseolica* and *P. glycinea* in cultural characteristics. Mung bean isolates caused water-soaking on mung bean, Red Kidney bean, Fordhook 243 lima bean, and Acme soybean, but not on other soybeans used to differentiate *P. glycinea* races. None of six *P. phaseolica* isolates or six *P. glycinea* races caused water-soaking on mung bean. Since only the mung bean isolate was virulent to mung bean, it is proposed that they be named the mung bean strain of *P. phaseolica*. The pathogen was introduced on seed. In experimental plots at 11 locations in Ohio initial infection occurred in plots of Berken, Jumbo, and Oriental cultivars from seed produced in Oklahoma. In experimental plots in Brazil, Peru, and Thailand seed from greenhouse-produced Berken seed had no initial infection. Halo blight eventually developed in most cultivars at all locations. However, at four locations, plots from greenhouse Berken seed isolated from a source of primary inoculum remained free of halo blight. At these four locations, Berken with halo blight (Okahoma seed) yielded ca. 60% less than healthy Berken (greenhouse seed).

**Multidense satellite of tobacco ring spot virus: a series of components with a uniform distribution in density between each component. I. R. Schneider, R. Hulse, & R. Markham (ARS, USDA, Beltsville, Md., John Innes Inst., Colney Lane, Norwich, England).** The satellite (STRSV) tobacco ring spot virus (TRSV) ratio found in the inoculated plant was higher when the STRSV/TRSV ratio in the inoculum was increased. STRSV is composed of 11-14 homogeneous components ranging in buoyant density (BD) in CsCl from 1.408 through 1.529. Each component differs in density from its nearest neighbor by ca. 0.006. The BD's are not identical to 6.6. The artifacts caused by CsCl, but correlate with the position of the particles sedimenting in a sucrose gradient. Two components of STRSV cannot be distinguished in BD from bottom component of TRSV; another component of STRSV cannot be distinguished in BD from middle component of TRSV. Since some TRSV is always present in STRSV infections, a satellite population has 11 components at the least, but may have as many as 14 distinct components. The data indicate that each STRSV particle of lowest BD has ca. 11 nucleic acid strands (molecular wt) of 86,000 daltons/strand. Each STRSV particle of next higher buoyant density has one more nucleic acid strand than its nearest neighbor of lower density. Each particle of highest density contains 24 nucleic acid strands.

**Etiology of Cephalosporium gregatum in soybean. R. W. Schneider & J. B. Sinclair (Univ. Ill., Urbana).** In greenhouse and field experiments, internal browning in soybean stems, caused by *Cephalosporium gregatum*, developed to a greater extent in plants inoculated at a younger stage (6 weeks) than in those inoculated at later stages (8-12 weeks). These results suggest that if maximum symptom development is to occur in the field, the fungus must enter the roots at an early age. Evidence for conidial movement within the transpiration stream after root penetration suggests that this may be a factor. *Cephalosporium gregatum* spread to the tops (ninth node) of plants in the early pod-
filling stage within 1 and 2 days, respectively, when plants were grown in a fungal suspension or when the hypocotyls were dipped into a conidial suspension (25,000/ml).

Though the fungus enters the roots, internal browning may be delayed by warm temperatures. There was a direct relation between spruce time of infected plants to cool temperatures (18-24 C) and extent of internal browning.

**Invasion of rough rice in storage by strains of Aspergillus flavus, H. W. Schroeder & R. A. Boile (ARS, USDA, Citrus Ctr., Sarasota, Fla.).** Belle Patna rough rice was inoculated with spores of each of the three strains of A. flavus, separately and in all combinations. P-70-53I (I), green-spored, produces large quantities of aflatoxin; P-70-25F (II), green-spored, produces an orange fluorescent compound but no aflatoxin; and AF-2 (III), a white-spored mutant, produces little or no aflatoxin. In all treatments, invasion of kernels increased significantly only during the 1st week in storage in a relative humidity of 90% at 25 C. The white-spored mutant strain (III) failed to invade as readily as the two green-spored wild-type strains (I, II) when each strain was applied separately. Maximum toxin (33 ppb) was detected in rice inoculated with two non-producers of toxins (II, III). When mixed inoculum was used, all strains appeared equally capable of invading rice under the conditions of this experiment. The separate strains appeared to retain their identity and to mix physically rather than through sexual recombination of genetic materials.

**Evidence for an infectious double-stranded RNA virus in tomatoes, L. S. Semancik & L. W. Weathers (Univ. Neb., Lincoln, Univ. Cal., Riverside).** The replication of the free-RNA citrus exocortis virus (CEV) was determined by bioassay of protein extracts from Gymnopus avrangianus. Infectivity was max 45-60 days postinfection (p.i.), with 50% still remaining at 140 days p.i. When the nucleic acid was fractionated into 2 x LDI (DNA-containing) and supernatant fraction, the supernatant fraction contained twice the total infectivity of the supernatant. Infectivity was reduced slightly when preparations were heated to 20-100 C for 10 min. Infectivity was linear from 100 to about 140 C. Concentrations of diethylpyrocarbonate which reduced the infectivity of single-stranded viral RNA by 95-100% inhibited CEV by only 40-70%. CEV infectivity was eluted from methylated albumin at 0.72 M NaCl as compared to 0.69 M and double-stranded RNA at 0.77 M. The infectivity distribution of CsCl equilibrium sedimentation conformed to a broad density range (1.55-1.64) and indicated the presence of single- and double-stranded RNA species. These data suggest that the major infective unit of CEV is a free-RNA double-stranded structure.

**Diseased leaves as a source of ascospores of Leptosphaeria (Pseudopeziza) medicaginis for alfalfa inoculations. G. Skemer (S. Dak. State Univ., Brookings).** Diseased leaves hand-picked from Medicago sativa plants in late June and early July in western South Dakota were air-dried, then wetted and dried flat between blotting paper and secured in numbers of ca. 30 between 4 x 5-inch plastic window screens (screens 37 mesh/inch) sewed together with nylon thread. The leaf-carrying screen(s) placed (leaf adaxial side up) between heavy galvanized wire mesh-wide screens and set outdoors in early August on open ground in the shade. Mature apothecia appeared by late August 1969 and late October 1970. Their number increased to ca. 5/leaflet (range 0-25) over an additional 2 weeks' exposure, and at that time the screened leaves were stored in a cool, dry place for use in greenhouse plant inoculations. Screened leaves were arranged on light-wt wire mesh screens over plants, and the whole was enclosed for 3 days with a transparent plastic sheet. Seven hundred potted alfalfa plants in two 4 x 12-ft horticultural cages were inoculated the first time from 145 screens, first one bench, then the other; and a second time from 470 screens, both benches at once. The results suggest uniform ascospore discharge and yellow leaf blight development over the bench areas.

**Uncoating of TMV-RNA in tobacco leaves treated with Carbosulfan or chloroform. J. G. Shaw (Univ. Ky., Lexington).** Tobacco leaves were treated with three different ways prior to being rubbed with tobacco mosaic virus which was labeled with 14C in its protein moiety. Leaves were (i) rinsed with water and dusted with Carbosulfan; (ii) rinsed with chloroform followed by water; or (iii) rinsed with water and finely ground. Three min after inoculation, the leaves were rinsed with water and finely ground. The TMV-protein released from the virus was determined by centrifugal analysis of extracts of the leaves. Uncoating activity in leaves rinsed with chloroform was twice that in leaves dusted with Carbosulfan and up to 10 times that in leaves to which neither chloroform nor abrasive had been applied.

**Successions of microorganisms and patterns of discoloration and decay after wounding in red oak and white oak. W. S. Weyant (USD, Northeast Forest Exp. Sta., Durham, N.H.).** Trunks and roots of 23 mature red oak, Quercus rubra, and white oak, Q. alba, were dissected to determine the patterns of discolored and decayed (D & D) wood associated with 22-year-old basal fire wounds and fire-scorched wounds inflicted during subsequent salvage operations. The columns of D & D wood had advanced at least 2 ft farther along the sapwood-heartwood boundary present at the time of wounding. The D & D tissues associated with the wounds were confined to the tissues present when the wounds occurred. The heartwood cylinder constricted abruptly below the root collar. Heartwood formation was retarded about the wounds. Complex patterns of D & D wood occurred at the root-trunk transition zone. Isolations for microorganisms were made in a systematic way from columns of D & D wood in 19 trees. Bacteria and non-hymenomycetous fungi were isolated consistently from columns that contained only discolored wood, and from the discolored wood at the distal margins of columns that contained decay. Hymenomycetous fungi were isolated commonly from the tissues at the border of D & D wood. A large variety of microorganisms was isolated from the advanced decay. The results indicate that basic patterns of discoloration and decay and successions of microorganisms follow wounding in oaks.

**The role of dew and temperature in the epidemiology of Botrytis leaf blight of onion. P. B. Shoemaker & J. W. Lorbeer (Cornell Univ., Ithaca, N.Y.).** Conidia of Botrytis squamosa (isolate 64a) were atomized onto leaves of onion plants (Allium cepa) which were then held for various periods in a dew chamber at 20 ± 1 C. Only plants whose leaves remained wet continuously for 6 or more hr developed an appreciable number of lesions when intervals of 0, 3, 6, 9, 12, and 18 hr were tested; lesion numbers were determined 40 hr after inoculation of the plants. There was a significant increase in lesion numbers for each interval beyond 6 hr. Necrosis (blighting), measured from the leaf tip, also increased significantly as the dew period was increased. When plants whose leaves remained continuously wet were kept 40 hr at constant temperatures of 7-30 C, lesions developed only in the range from 9 to 23 C, inclusive. Lesion numbers and amount of blighting were significantly greater on older outer leaves than on younger inner leaves. Leaf age and position may be important factors determining susceptibility. When conidia were brushed onto dry leaf surfaces, lesions developed only if the plants were continuously kept wet. Germination of conidia was maximum at 15 C, and occurred over the range 12-27 C. Growth in culture was maximum at 24 C and occurred from 9-31 C.

**Distribution and metabolic fate of benomyl in dwarf pea plants. M. R. Siegel & A. J. Zabala, Jr. (Univ. Ky., Lexington).** 14C benomyl was added to Knowl's solution in
which 21-day-old peas (Pisum sativum ‘Laxton Progress’) were growing. Plants were treated with benomyl for 4 days and harvested at 12-day intervals until senescence of the plants occurred (52 days after treatment). Neither 14C benomyl nor 14CO2 was recovered from the plants at any time. A fungitoxic derivative, 14C methyl 2-benimidazole-carboxylate (MBC), was present in large quantities in organic solvent extracts of the plants at all time periods. At senescence, 76% of the label was present in MBC and 7% in unknown water-soluble metabolites; 15% of the label was bound to the plant residue. When the plant residue was treated with NaOH, a portion of the bound label was precipitated as benomyl. Benomyl and dicofol residue in plants was dependent on the age when harvested. The labeled products were translocated to the stems and leaves of the plant. Fifty-two days after treatment, 94% of the label was in nonroot portions (stems, leaves, and pods). However, only 12% of the total label was present in tissue from the plants that were treated. These data suggest that long term chemotherapeutic effects could be dependent on the distribution, persistence, and concentration of MBC in plant tissue.

**Effect of light on growth and sporulation of Colletotrichum destructivum in culture.** R. C. Sievert (ARS, USDA, Univ. Tenn., Greeneville). Cultures of Colletotrichum destructivum were grown on numerous solid media (generally 1.5% agar, w/v), at initial pH values from 4.0 to 7.2, at constant temperature of 24 C. Variations in the carbohydrate and/or plant extract included in the growth medium resulted in differences in growth, but addition of 2.5% diethylamine gel columns and sediments on a 5-20% sucrose-density-gradient showed that the mol wt of purified RNA corresponded to 4-5 Svedberg units. Further purification by exonuclease digestion showed that the RNA was not completely resistant to exonuclease digestion. Infection by C. destructivum of the different fractions produced by purification were conducted on the local lesion host, Scoparia bicornis, and showed a 35-fold concentration from the crude sap to the DEAE-cellulose-treated fraction.

**Infectious low molecular weight ribonucleic acid from tomato infected with potato spindle tuber virus.** R. P. Single & M. C. Clark (Can. Dep. Agr., Fredericton, New Brunswick). Infectious low mol wt ribonucleic acid (RNA) from Sheyenne tomato tissues infected with potato spindle tuber virus was extracted and purified by precipitation with 5% aqueous cetyltrimethylammonium bromide followed by chromatography on diethylaminoethyl cellulose. Electronmicroscopy of 2.4% polyacrylamide gel columns and sedimentation on a 5-20% sucrose-density-gradient showed that the mol wt of purified RNA corresponded to 4-5 Svedberg units. Further purification by exonuclease digestion showed that the RNA was not completely resistant to exonuclease digestion. Infection by C. destructivum of the different fractions produced by purification were conducted on the local lesion host, Scoparia bicornis, and showed a 35-fold concentration from the crude sap to the DEAE-cellulose-treated fraction.

**Effects of Alternaria alternata infection and other leaf injuries on subsequent growth of tobacco.** L. J. Slama & J. R. Stavely (ARS, USDA, Beltsville, Md.). We reported that heavy infection by Alternaria alternata severely inhibited the expansion of the youngest inoculated leaves. To determine further effects on plant growth, we treated greenhouse-grown Coker 187 Hicks tobacco plants (having about 11 leaves) by inoculating with A. alternata or Colletotrichum destructivum, wounding, or removing the youngest leaves. Dimensions and numbers of leaves on these plants and controls were recorded after 0, 2, 4, and 6 weeks. At flowering, we recorded the date, plant height, and number of leaves per plant. All treatments inhibited expansion of the youngest treated leaves, but caused production of more posttreatment leaves than the controls. The increase varied with the number of leaves on the controls. Alternatively, the increase in the number of leaves above the number on the controls was 0.5 and 6 on lightly and heavily infected plants, respectively. Heavy A. alternata infection induced twice as many additional leaves as did leaf removal, and three times as many as severe wounding or C. destructivum infection. Stem elongation was inhibited in the youngest treated portion, but was not affected above this area. Severe A. alternata infection caused a reduction in the size of leaves produced above infected tissues, but other treatments had no effect.

**Influence of nitrogen fertilizers on rhizosphere pH and take-all of wheat caused by Ophiobolus graminis.** R. W. Rouse & R. J. Cook (W. State Univ., USDA, ARS, Pullman, Wash.). Take-all of cereals and Ophiobolus-patch of turf can be controlled in the field by NH4+-N, but not commonly by NO3--N. Take-all control with (NH4)2SO4 in greenhouse pots of Puyallup fine sandy loam (native pH 5.7) was negated by limiting the soil pH to 5.0. The treatment with 0.84% (NH4)2SO4 was obtained with no H and with Ca(NO3)2 in Ritzville soil to pH 5.7; but without acidification, only (NH4)2SO4 was effective at the native pH 7.5. Disease was less at pHs <6.0 (bulk soil pH), but severity differed for each soil, and was only moderately controlled with (NH4)2SO4. Disease ratings were highly correlated with pH (Rhizosphere soil pH); and regardless of soil type, pH4, N fer-
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S. A. Soileau, D. D. Van Etten, & D. F. Bateman (Cornell Univ., Ithaca, N.Y.)

Phaseolus vulgaris hypocotyls infected with R. solani were triturated in 95% ethanol (1:4, v/v). The extract was filtered and centrifuged at 15,000 g for 10 min. Water (v/v) was added to the supernatant, and the ethanol removed in vacuo at 40°C. Phaeollin was extracted from the aqueous phase by partitioning twice with petroleum ether (1:4, v/v). Bioassays of the residual aqueous fraction revealed a second antifungal component. This fraction was not removed with CHCl₃ (1:2, v/v), but the CHCl₃ partitioned twice with 2 M NaCO₃ (1:1, v/v). The NaCO₃ fraction was adjusted to pH 6.0 with HCl, extracted twice with CHCl₃ (1:1, v/v), and the CHCl₃ fraction dried in vacuo at 40°C. The residue was dissolved in 95% ethanol and subjected to thin-layer chromatography gel using a 1-bromo-3-dioxan-5-ol/8:1 solution. The Ref for R. solani corresponds to an absorbance of 6.0 OD units/ml at 294 nm.

Aerobiology of peanut leaf spot fungi. D. H. Smith & F. L. Crossy (Univ. Ga., Nat. Weather Serv., Experiment.) Air was continuously sampled with a Hirst spore trap in a peanut (Arachis hypogaea 'Argentine') field during the growing seasons of 1969 and 1970. Cocercospora arachidicola conidia and Lepiophaeraulina crassicae ascorpores were present in the air during June, July, August, and September of both years. Cocercosporidium personatum conidia were not trapped in either year. The concentration (spores per m³ of air per 24-hr period) of C. arachidicola conidia and L. crassicae ascospores varied widely from day to day. The ascospore concentration of C. arachidicola conidia >300 m³ occurred from 3-4 September 1970. Lepiophaeraulina crassicae ascospores were trapped on 71 days in 1969 and 79 days in 1970. Low numbers of L. crassicae ascospores were present in the air until the level of peanut leaflet abscision reached 37%. In late July of 1969 and early August of 1970. The peak concentration (1,300 m³) of L. crassicae ascospores occurred from 23-24 August 1970. Very few phragmosporous L. crassicae ascospores were trapped.

Trehalase activity in natural and artificially fungus-colonized soil. R. E. Smith & R. Rodriguez-Karaba (Auburn Univ., Auburn, Ala.). Trehalase activity (TA) was measured in natural soil and in soil incubated with Scelerotium rolfsii (SR), Fusarium oxysporum f. sp. phaseoli (FOV), and Trichoderma viride (TV). In natural soil, incubated over a temperature range of 20-30°C, TA increased to maximum at 50°C, then declined. Acetate, propionate, citrate, succinate, maleate, and phosphate buffer at pH 5 were used to determine maximal TA over a pH range of 3.5-7.5. Maximal TA occurred in propionate buffer at pH 4.95, and was at a high level in acetate buffer. When pH was held constant at 5.5 (acetate), andionic strength increased from 0 to 5 M, TA decreased. When trehalase concentration was varied in 7.5 M Tris buffer, maximal rates of increase in TA occurred between 0 and 20% (w/v). At constant volume, increases in amount of soil resulted in increased TA. There was a linear increase in TA with incubation time from 0 to 22 hr. Determination of TA in colonized soils was performed using 125-ml flasks containing 5 g soil, 5 ml 2% (w/v) trehalose, and 1 ml toluene. Soil pH values were 4.1, 5.3, and 6.4 for SR, FOV, and TV, respectively. During a 3-hr incubation period at 35°C, TA in soil colonized by the three fungi was in the following order: SR > FOV > TV.

Lead, mercury, and sodium contamination of urban trees. W. H. Smith (Yale Univ., New Haven, Conn.) Acer platanoides planted in New Haven were analyzed for Pb and Na contamination. During the fall of 1970, branch samples were collected ca. 2 m aboveground from 50 trees located within 7 m of streets. Analysis was by atomic absorption spectrophotometry. Lead content of the growing season was analyzed as follows (µg/g dry wt basis, mean ± se, unwashed and washed, respectively): intact branches 163 ± 13, 167 ± 12; leaves 113 ± 33, 96 ± 20; twigs 25 ± 7; V. agnes leaves as follows: intact branches 374 ± 30, 362 ± 35; leaves 469 ± 183, 275 ± 73; twigs 319 ± 71, 347 ± 57. The Pb concentrations are in excess of most reports of tree Pb levels. Since the concentrations of unwashed and washed samples were not significantly different, either the cations were precipitated inside the leaves or the washing procedure was inadequate to remove surface deposits. Hg analyses revealed the following concentrations (µg/g) on single, unwashed samples: Acer saccharum leaves, 0.81; Alnus glutinosa leaves, 0.62; Platanus racemosa leaves, 0.73; twigs, 0.13; Picea abies intact branches, 0.22; Pinus nigra leaves, 0.17; twigs, 0.08; Quercus palustris leaves, 0.77; Taxus sp. intact branches, 0.84; and Tilia cordata leaves, 1.10.

Competition of peaches and nectarines in packing houses. W. L. Smith, Jr., J. M. Wells, & R. W. Penney (ARS, USDA, Beltsville, Md.). Surveys indicate that peaches and nectarines are often essentially free of decay-producing organisms when they enter packing houses, but can be contaminated when exposed to unsanitary conditions in packing houses. Such contamination can cause a high percentage of the decay occurring during marketing. In these tests, most of the decay due to contamination was brown rot, caused by Monilia fructicola. Chief sources of contamination were hydrocoolers and tanks for dumping bulk bins. After 3-9 days at shipping temperatures, fruit sampled from these areas had from 2 to 10 times as much brown rot as fruit sampled directly from the orchards. Additional contamination sometimes occurred during dehusking. Suggestions for reducing contamination include (i) more frequent cleaning of hydrocoolers and dump tanks; (ii) chlorination of cooling or dump tank water; (iii) hot water treatments (52°C for 1-2 min); and (iv) cooling of fruit in air rather than water.

Inheritance of leaf rust resistance in Red River 68, a semidwarf red spring wheat. G. D. Stoller (N.D. State Univ., Fargo). Inheritance of seedling and adult plant resistance to wheat leaf rust incited by Puccinia recondita f. sp. tritici was investigated in Red River 68, a semidwarf hard red spring wheat resistant to prevalent races of wheat leaf rust. Red River 68 was crossed and backcrossed to the susceptible variety Thatcher for the genetic analysis. The F₁, F₂, and backcross progenies of the crosses were inoculated in the seedling and adult stages with a culture of race 15. Plants that had been purified by three successive, single-pustule isolations. Seedling and adult resistance to physiology race 15 of P. recondita was conditioned by a single dominant gene in Red River 68 as shown by three criteria. The F₁ plants were all resistant to race 15; the 104 backcross plants segregated into a 1:1 ratio (P between .50 and .75) of resistant to susceptible; and the 308 F₂ plants seg-
regated to fit a 3:1 ratio (P between .50 and .75). The same gene was apparently operative in both seedling and adult stages, providing Red River 68 with resistance to wheat leaf rust.

**Effect of Cercospora nicotianae infection on chemical constituents of cured tobacco leaf.** J. R. STAVELY & J. F. CHAPLIN (ARS, USDA, Beltvile, Md., Oxford, N. C.). In 1970 we grew three burley, three Maryland, and four cured tobacco cultivars in the field in two blocks of six plants/each of five replicates per cultivar. One block was inoculated with *C. nicotianae*, the incitant of frogeye and green leaf spots. All inoculated cultivars were heavily infected, but the most frogeye occurred on the burley, and the least green spot occurred on Maryland types. The second block remained disease-free. Leaves were harvested twice from equivalent positions on all plants. Half the harvested leaf from each replicate was air-dried; half was flue-cured. Analyses of all cured leaf showed that the disease caused respective decreases in total alkaloids of 44, 10, and 15% in the first and 36, 30, and 18% in the second harvest from burley, flue-cured, and Maryland types. Soluble phenols were reduced by the disease 85% in air-cured and 57% in flue-cured leaf. Reducing sugar content was decreased an average of 24% in all leaf that was flue-cured. The greatest reduction occurred in flue-cured leaf of flue-cured cultivars, which had 2-4 times more sugar than the other types. Reducing sugar content and its reduction by disease was greater in the second than in the first harvest. Leaf spot caused a significant 3 and 9% increase in total nitrogen in the first and second harvest, respectively.

**Identification of tenuazonic acid as an important toxic substance produced by Alternaria spp.** J. A. STEELE & C. J. MUNNIA (Univ. Minn., St. Paul). Certain isolates of *Alternaria* spp. when grown on moist autoclaved corn or rice mixtures, produce substances toxic to rats, swine, and turkeys. One of these substances was previously isolated but not fully characterized. Fifty mg of the unknown toxin were obtained from moist autoclaved corn which had been inoculated with an *Alternaria* isolate known to produce large quantities of the toxin. This material was submitted for elemental analysis, flame emission spectroscopy, UV spectroscopy, and infrared spectroscopy. Chemical degradation studies were made by acid hydrolysis and treatment with sodium chloride. The spectral data and degradation products indicate that the toxin was the mixed potassium, sodium, and calcium salt of tenuazonic acid (3-pyrylino-2-one, 3-acetyl-L-5 sec-butyl-4-hydroxy).

**Dutch elm disease: control with soil-injected fungicides and surfactants.** R. J. STIPES & DONNA R. OBERWALD (Va. Polytech. Inst., Va. State Univ., Blacksburg). The control of Dutch elm disease (DED) by soil-injected fungicides was investigated. Methyl 1-(butylcarbamoyl)-2-benzimidazolecarbamate (B), N-[(chloromethylthio)-4-cyclohexene-1, 2-dicarbamoxide (C), and thiaboradazole [2-(4-Thiazolyl), benzimidazole] (T) suspensions alone or in 1.0% (v/v) Tween 80 (polyoxyethylene sorbitan monooleate) surfactant, were injected into the root zone of 3-year-old up-rooted *Ulmus americana* trees 14 days prior to artificial borer entry. Seven days later the tree was injected with 6,000 ppm active ingredient suspension were administered to each of nine completely randomized elms/treatment. Mean visually estimated foliar symptoms in B+, T- and C-treated and check trees were 1.0, .52, 78, and 68%, respectively. Four days after inoculation, infected inoculated leaves were found by 11 and 85% over C and T alone, but C, umit was isolated in significantly higher frequencies in all Tween 80-treated trees. Bioassays of solvent-extracted leaves revealed the presence of B, C, and T 1 day after application. These findings suggest promise of controlling DED in plantations with B by this method.

**Dothiorella canker of dawn redwood (Metasequoia glyptostroboides).** R. J. STIPES, F. S. SANTAMOUR, JR., & R. C. LAMBRE (Va. Polytech. Inst., Va. State Univ., Blacksburg, U.S. Nat. Arboretum, Washington, D.C.). We observed a new, destructive, perennar canker on 20-year-old dawn redwood (*Metasequoia glyptostroboides*) in the United States National Arboretum, Washington, D.C. Several stems (20-20 cm diam) with wilting and necrotic leaves were ultimately girdled. Isolations from cankers consistently yielded a *Dothiorella* in pure culture. The colony on potato-glucose agar was white, then turned olivaceous green with a darkly pigmented margin. Koch's postulates were confirmed by inoculation of seedlings in the greenhouse. The fungus produced typical white rot lesions on apple (Malus sylvestris) fruits, and also cankers on stems of apple and white pine (Pinus strobus). The apple white rot fungus *Botryosphaeria dothidea* (= *B. ribis*) on which we believe the pathogen to be, also produced similar cankers on *Metasequoia* stems. The syndrome of *Metasequoia* also included marked resin exudation and extensive discoloration of xylem and phloem tissues in advance of the visible canker lesion. Rapid blighting resulted commonly from xylem-inoculated stems. Neither pycnidia nor perithecia were observed on natural infections, whereas pycnidia were abundant on apple fruit and stem lesions.

**Effects of fluoride on pollen germination and pollen tube growth.** C. W. SULZBACH & MERRILL R. PACK (Wash. State Univ., Pullman). Tomato plants continuously exposed in atmospheres having 5ppm or more of F/3, as HF, produced smaller fruits or fruits with fewer seeds, or both, than similar control plants. Such treatment reduced pollen germination and subsequent pollen tube growth. Pests of self- and manually pollinated flowers (the latter massed when flowers were fully open) were removed 2 to 10 days after pollination, stained with aniline blue, and observed under UV light. The amount of pollen on the stigma, per cent of grains which germinated, number of grs reaching the style, and number of tubes reaching ovules were determined. Pollen from HF-exposed plants had lower germination percentages in vitro than pollen of the controls. A similar result was obtained when NaF was added to the germinating media. Less pronounced results were obtained in cross pollinations (both pollen of controls to HF exposed and the reciprocal), suggesting that the F affected both the male and female parts of the flower. Inhibition of germination and tube growth appears to be proportionately related to the concentration of the F; however, the degree of the F effect can be partially regulated by the amount of Ca within the nutrient solution or media.

**Ecology of root rot of turnips grown for processing.** D. R. SUMMER (Univ. Ga. Coastal Plain Exp. Sta., Tifton). Turnips roots grown for processing (with greens) frequently are severely damaged by root rot in the Georgia Coastal Plain. Brown external discoloration is usually first observed near the top of the roots, and the disease progresses down the vascular tissue in the interior of the root. Both *Colletotrichum higginsianum* and *Erwinia* sp. were commonly isolated from infected roots. Roots were inoculated by wounding or nonwounding with both pathogens alone and together. Typical symptoms were produced only with *C. higginsianum* and *C. higginsianum* + *Erwinia* sp. The latter caused the disease in the roots. Seven days later the root with a distinctive putrid odor. In a field test, 71% of the roots were unacceptable for processing when the fungus + the bacteria were injected into the roots, as compared to 17% when the fungus alone was injected and 1.9% when the fungus was injected without inoculating the fungus. The disease was most pronounced on upper leaves and tops of the roots. When the fungus was sprayed on only upper leaves or when plants were not inoculated, only 15% of the roots were unacceptable. In greenhouse tests, soil moisture had no significant effect on root rot. Boron deficiency plus the pathogens significantly increased the percentage of unacceptable roots as compared to boron deficiency or the pathogens alone.

**Etiology of crazy top of corn.** M. H. SUN & A. J. ULLSTROM (Purdue Univ., Lafayette, Ind.). *Sclerotophora*...
macrosperma has long been implicated as the incitant of crazy top of corn. However, proof of pathogenesis based on fulfillment of Koch's postulates has not been established. Artificial inoculation was achieved by the following technique: young "green island" leaf tissue of diseased seedlings growing from infected seeds was incubated in water at 24-26°C for 24 hr for production of sporangia. The sporangial suspension was then incubated at 16°C for 5-12 hr to stimulate sporangial germination and zoospore production. The motile zoospores were injected into coleoptiles of 2- to 3-day-old corn seedlings (Pioneer Hybrid 3306) with a syringe. The inoculated plants were incubated at 16°C for an additional 48 hr before transfer to the greenhouse (24-26°C). The first green island symptoms appeared on the fourth or fifth leaf about 4 weeks after inoculation. Thereafter, systemic symptom development was similar to that of the diseased plants growing from infected seeds. Sporangia were isolated from the artificially inoculated plants and successfully used for re-inoculation. Infection has been as high as 70% with the method, and the technique is being used for testing susceptibility of corn genotypes.

Purification and characterization of a severe strain of peanut mosaic virus. M. K. SUN and T. T. HUEY (N.C. State Univ., Raleigh). A severe disease of peanut (Arachis hypogaea) having chlorotic, sometimes necrotic, mosaic leaf symptoms is caused by a severe strain of peanut mosaic virus (PMV-s). PMV-s infected only nine species of leguminosae of 30 mechanically inoculated species in eight plant families. Yield reduction was 90% and 70% in Floridant and NCs peanut in the field, respectively. Seed transmission was 0.001% in NCs. In unbudded sap from Pisum sativum 'Alaska Early', properties were: thermal inactivation between 60 and 65°C, dilution end point between 10-3 and 10-4, and longevity at room temperature 12 hr with a half life of 83 min. Virus was purified from Alaska Early peas by chloroform-butanol clarification, polyethylene glycol precipitation, and resuspension of the virus in 0.025 M NaHPO4, pH 7.4, buffer containing 0.025 M sodium sulfate and either 0.1-0.6 M urea or guanidine-3HCl. PMV-s particles are flexible rods with a normal length of 740 nm. Virus inclusions were similar to members of the potato virus Y group. PMV-s was not serologically related to soybean mosaic, bean common mosaic, potato Y, or tobacco etch viruses.

Extractable phenols in clear, discolored, and decayed tissues of sugar maple and red maple. T. A. TATTAR (Univ. New Hampshire, Durham). Ethyl acetate extracts were prepared from hot water extracts of 10-g samples of clear, discolored, and decaying tissue of sugar maple, Acer saccharum, and red maple, A. rubrum. Extracts were run on 2-way cellulose thin-layer chromatography plates with butanol:acetic acid:water (6:1:2) and 7% acetic acid: 0.05% sodium acetate. Total phenols were determined by the Folin-Ciocalteau method on methanol extracts of clear red maple tissue. Gallic acid, its esters, and catechin were identified as the major phenols in clear tissue of both red maple and sugar maple. These phenols were absent from discolored and decayed tissue. An unidentified flavonoid appeared only in discolored tissues. Total phenols in clear red maple tissue were highest (10 mg/g) near the cambium, and decreased progressively toward the pith to 0. Total phenols in clear sugar maple tissue were highest (10 mg/g) near the cambium, and decreased progressively toward the pith to 0.

The processes of discoloration and decay result in decreases in extractable phenols confined to clear unaffected tissue. Total phenols in clear tissue of red maple are highest in the most physiologically active tissues near the cambium. The low level of phenols in discolored wood may permit the growth of decay fungi unable to grow at the phenol concentration in clear tissue.

Synnemata induction in Ceratocystis ulmi. P. A. TAYLOR, E. B. SMALLEY, & F. M. STRONG (Univ. Wis., Madison). Why type isolates of Ceratocystis inornata and Torr. do not readily form synnemata (cornia) on common culture media unless such media contain elm wood fragments or extracts of elm wood. Aqueous and methanolic extracts of elm wood, chromatographed on a polyamide column, yielded a "phenolic fraction" and a "water fraction" containing nutrients. The fungus grew well on 1% agarose gel containing the "water fraction", but did not produce synnemata unless the "phenolic fraction" was also present. Fructose in the "water fraction" was essential for synnemata production, but not for fungal growth. A polyamide extract was isolated by successive chromatography on polyamide and silica gel columns. When added to the "water fraction" agar gel or to potato-dextrose agar (PDA), this phenol induced C. ulmi to form synnemata. Synnemata production in PDA was initially inhibited when the added glucose in PDA was replaced by fructose. The active phenol was identified as catechin (7,4,5,7-pentahydroxyflavay) by UV and proton magnetic resonance spectroscopy, mass spectrometry, and cochromatography. Authentic 4-(4-)catechin was identical with the isolated compound in both biological and chemical aspects.

A chloronene-resistant mutant of Ustilago maydis. R. W. TILMAN & H. D. SISLER (Univ. Md., College Park). A mutant of Ustilago maydis resistant to chlorone was isolated from medium containing 8 µg/ml of toxicant. Growth of the wild type was prevented by 8 µg/ml. There was no appreciable difference in the uptake of ring-labeled or methyl-labeled chlorone-14C by the mutant or wild type. All radiolabeled methylme were added to mutants, cultures could be recovered after 24 hr as unaltered chlorone. The chlorone-resistant mutant was also resistant to 2,6-dichloro-4-nitroaniline, diphenyl, hexachloroazene, naphthale, p-dichlorobenzene, pentachlorothiophene, and sodium-o-phenylphenate. Wild type and mutant were equally sensitive to 2,4,6-trichloro-o-phenol; 2,4,5,6-tetrachlorothiophenyl; o-benzyl-chloro-phenyl; methyl-2-benzimidazole carbamate; and 1,2,4-trichloro-3,5-dinitrobenzene. Tolerance to chlorone in progeny from a cross of the resistant mutant to a sensitive wild type indicated single gene resistance.

The capacity of leaf extracts of tobacco to react with ozone. H. TOMLINSON & S. RICH (Conn. Agr. Exp. Sta., New Haven). The capacity of leaf extracts of tobacco to react with ozone was tested by measuring the ability of these extracts to inhibit the peroxidation of methyl linolate exposed to ozone. Methyl linolate (60 µg) suspended in 3 ml neutral 0.025 M phosphate buffer plus 40 µg of sodium lauryl sulfate and 10% ethanol gave a linear increase in peroxidation when ozonated for 1-4 min by an ozone generator containing 8 ppm ozone during 30 min at the rate of 1 liter/min. Peroxidation was measured as the increase in malonyl didehye (MDA). Cysteine (20, 40, and 60 µg), added before ozonation, inhibited MDA formation 45, 55, and 65%, respectively, when ozonated for 2 min, and 30, and 48% when ozonated for 4 min. Juice expressed from a single frozen disc (1 cm-diameter) of toacco leaf (Nicotiana tabacum) also inhibited formation of MDA when added before ozonation. After 2 min of ozonation, juice from an ozone-resistant and an ozone-sensitive cultivar inhibited the formation of MDA 52 and 55%, respectively, after 4 min of ozonation, MDA formation was inhibited 4% and 28%, respectively. Therefore the capacity of sap from the susceptible cultivar to react with ozone remained equivalent to 40 µg of cysteine, whereas the capacity of the resistant cultivar was depleted.

Processes of abnormal cell wall thickening in potato virus M-infected lesions in Red Kidney bean. J. C. Tu & C. HIRUKI (Univ. Alberta, Edmonton, Can.). Abnormal secondary wall thickening, inhibited in secondary walls of normal potato bodies was found in cells in the peripheral zone of potato virus M-infected lesions in Red Kidney bean. The initiation of the deposition in an affected cell was closely associated with increased cellular metabolic activities evidenced by increased numbers of ribosomes and mitochondria. Slight thickening of the plasmalemma was observed at this stage.
Later, the roughening extended into invagination of various degrees and, therefore, numerous invaginated periplasmic spaces were formed in the cell. Increase in numbers of dictyosomes and in deposition of dictyosome-related vesicles and microtubules occurred near or in the invaginated portions of the plasmalemma. Multiplication of paramural bodies on the normal secondary wall became thickened, extended, and then coalesced with the adjacent deposits and finally coated the inner cell wall. The deposition initially resembled a loosely-packed spongelike matrix, later filled with fine fibrinous materials, and eventually became very dense.

*Effect of streptomycin on development of fire blight in artificially injured pear fruit.* T. VAN DER ZWET & H. L. KEL (FSRD, ARS, USDA, Beltsville, Md.). Immature Bartlett pear fruit (4.5 cm) in the orchard were injured by bruising with the head of a thumbtack or by puncturing with a needle or nail at intervals of 0, 1, or 2 days before or after inoculation. The fruit were inoculated by dipping for about 3 sec in an aqueous cell suspension (5 x 10⁷ cells/ml) of Erwinia amylovora. At the time of inoculation, the fruit were dipped for 5 sec in a suspension of 100 ppm streptomycin. The antibiotic provided unsatisfactory control when applied at time of inoculation (0 days), regardless of the type of injury. Streptomycin applied 1 day before inoculation protected most of the infected fruit and those inoculated without fruit, but not those with or without streptomycin, showed considerably more blight than did punctured fruit, regardless of the interval between injury and inoculation. In laboratory tests, when fruit were injured, inoculated, and dipped in streptomycin at the same time, 3 times more fruit blighted when skin rupture accompanied bruising than without rupture. Without streptomycin, all fruit blighted regardless of degree of bruising. Under the severe conditions of these tests, streptomycin was most effective in preventing development of fruit blight only when it was present at the time of injury.

*Studies on the mode of action of phaeosolin.* H. D. VAN ETTEN & D. F. BATEMAN (Cornell Univ., Ithaca, N.Y.). Phaeosolin at 9, 15, or 47 μg/ml inhibited the ability of Rikhtoria solani to remove α-glucose-1-14C from a liquid medium (15 μg/ml is the min concentration needed to prevent an increase in dry wt of *R. solani* in shake culture). Microscopic examination of growing hyphae exposed to phaeosolin revealed immediate (<1.0-min) cessation of protoplasmic streaming, often accompanied by shrinkage of the protoplasts from the hyphal tips. Phaeosolin (15 μg/ml) incubated with autoclaved or nonautoclaved *R. solani* mycelium was taken up rapidly by the mycelium during the first 10-15 min, after which the rate of uptake leveled off; after 1 hr ca., 30% of the phaeosolin remained in solution. Biological degradation of phaeosolin (15 μg/ml) was not detected 1 hr after exposure to live mycelium. When *R. solani* was exposed to phaeosolin-14C (15 μg/ml) for 1 hr and then fractionated, most of the 14C (73%) was associated with the hyphal fraction removed by centrifugation at 5000 g for 15 min. Phaeosolin (23 μg/ml or above) rapidly lysed sheep red blood cells. The data support the hypothesis that the inhibitory action of phaeosolin is associated with its ability to disrupt normal membrane function.

*Comparative histopathology of the resistant and susceptible response of Pelargonium spp. to Xanthomonas pelargonii,* S. H. WAINWRIGHT & P. E. NELSON (Pa. State Univ., University Park). The histology of Pelargonium spp. susceptible, semi-resistant, and noninoculated plants. The most striking difference was the presence of tanninlike materials in the tissues of moderately resistant and resistant species, and their absence in susceptible species. The mode of spread of the pathogen was similar in all species, initially involving movement of the pathogen throughout the plant in xylem vessel elements and subsequent movement laterally into adjoining parenchyma cells. The relative numbers of the pathogen and the numbers of fascicles initially invaded were low in resistant species and high in susceptible species. In susceptible species, bacterial pockets formed around affected primary xylem vessel elements, enlarging to encompass all xylem cells in the fascicle, while portions of the phloem, phloem parenchyma, and *X. pelargonii* were invaded. *PeIargonum* spp. responded to infection by proliferation of a ring of cells around affected portions of fascicles, with cells immediately inside this ring having a suberinelike material formed on their walls. This response was greatest in the most susceptible plants, and decreased as resistance increased.

*Composition of the surface of the cotton boll in relation to spore germination and resistance to Diplodia gossypina infection.* S.-Y. C. WANG & J. A. PINECKARD (La. Agr. Exp. Sta., Baton Rouge). Diplodia gossypina attacks the bolls of many cultivars of cotton, being most destructive of very young bolls, less than 15 days of age, and bolls over 25 days old. The nature of the resistance of bolls of intermediate age, 15 to 25 days, was the purpose of this investigation. Surface washings of intermediate aged bolls and the nectarial fluids of the cultivar Deltapine 16 showed the presence of fructose, glucose, galactose, sucrose, raffinose, and 2 unknowns which stimulated spore germination. Upon removal of boll waxes with hexane and analysis of the extracts, similar sugars were found, with the exception of raffinose. Maximal amounts of waxes and cutin acids were found on bolls of 17 and 20 days of age. These materials inhibited spore germination and mycelial growth, and may account for the observed resistance. Exopolygalacturonase, one of the important cell-wall degrading enzymes produced by *D. gossypina*, was detected during spore germination, but only in the presence of boll leachates, suggesting that surface waxes and cutin are potential barriers to infection. A further study of the waxes and cutin of bolls of several cultivars may indicate materials useful for improved resistance.

*Enhanced growth of Armillaria mellea on extracts from roots of defoliated sugar maple trees.* P. M. WARGO (USDA Forest Serv., Northeastern Forest Exp. Sta., Hamden, Conn.). Defoliation of trees by insects appears to predispose them to attack by *Armillaria mellea*. Certain sugars, amino acids, and fatty acids that occur in roots stimulate growth of *A. mellea* in vitro. Defoliation can alter some of these chemical constituents, and these changes may be related to increased attack by *A. mellea* on defoliated trees. Root tissues from defoliated and nondefoliated sugar maple saplings were analyzed for changes in starch sugars, amino acids, and fatty acids. Defoliation caused a significant decrease in the starch content of whole roots, and a corresponding increase in glucose and fructose in the outer wood but not in the bark. Both total number and concentration of some amino acids increased in roots of defoliated trees, especially in the outer wood. Defoliation had no apparent effect on fatty acids, and concentrations were less than 0.001% dry wt. Growth of *A. mellea* was significantly greater on outer wood extracts from defoliated trees, but not on bark extracts. It appears that defoliation causes chemical changes in the outer wood of roots that are favorable for the growth of *A. mellea*.

*Influence of soil temperature on root rot of peach caused by Cylindrocladium floridanum.* D. J. WEATHER (ARS, USDA, Byron, Ga.). Six-week-old Elberta peach seedlings were transplanted into steam-sterilized soil, either noninfested or infested with mesococcoid of *C. floridanum*, and maintained in watered pots at 15, 20, 25, and 30°C in the greenhouse. Six weeks later, roots were washed free of soil, rated for severity of rotting, dried, and weighed. Samples of infected roots from each inoculated treatment were subjected to routine isolation procedures. *Cylindrocladium floridanum* was readily isolated from roots of inoculated seedlings. Root rot was slight at 15°C, moderate at 20°C, and severe at 25 and 30°C. Generally, soil temperatures that favored root rot were less favorable for growth.
of roots. The fungus was grown on potato-dextrose agar for 7 days at temperatures ranging from 9 to 36 C. Greatest colony diameters occurred at 27 C.

Studies on the relationship of the hoja blanca virus to the plant hopper vector, Sogatodes oryzae. A. J. Weiber, Jr., V. D. Daiser, & C. L. Graham (Plant Sci. Lab., Ft. Detrick, Frederick, Md.). The effects of the hoja blanca virus (HBV) of rice on Sogatodes oryzae were studied through several generations with replicated single-pair matings and reciprocal crosses of transmitting and nontransmitting individuals in all combinations. Reduced fecundity, nymph viability, and adult longevity in the transmitted line indicated that the virus adversely affected the vector. Both mean daily and total nymph production by viruliferous females was less than that by nonviruliferous females regardless of the virus-transmitting ability of the fertilizing male. Virus acquisition studies with nontransmitting insects from populations containing 8 and 64% transmitters (through transovarial passage of HBV) and from a virus-free population resulted in acquisition rates of 7.5, 6.0, and 7.0%, respectively. Deleterious effects of the virus on the vector and the low rate of virus acquisition may be compensating factors stabilizing the virus level in a randomly breeding population.

Influence of conditioning environment of Bermudagrass on sex differentiation of Meloidogyne graminis. A. J. Weiber, Jr., & J. A. Fox (Va. Polytech. Inst., Va. State Univ., Blacksburg). Sex differentiation of Meloidogyne graminis was found to differ on Tifgreen Bermudagrass conditioned under different temperature and nutrient regimes. Bermudagrass spires from plants conditioned at 26 and 32 C or at a low and high nutrient level were rooted in distilled water. Rooted spires were inoculated with 150 larvae at 26 C and incubated at 26 or 32 C. Different sex ratios obtained at an incubation temperature of 26 C were not significant, but at an incubation temperature of 32 C the nematodes from spires conditioned at 26 C averaged 44% males, whereas those from spires conditioned at 32 C averaged 21% males. Nematodes from spires conditioned at a low nutrient level averaged 45% males, whereas those from spires conditioned at a high nutrient level averaged 16% males. The indirect influence of environment on sex differentiation may be the result of differences in physiological or biochemical factors of host origin.

Virulence repression in Rhizoctonia solani by 3-methyl glucose. A. R. Whistling & T. Bowan (Univ. Cal., Berkeley). Long-term studies of the disease-repressing action of 3-methyl glucose on virulence in Rhizoctonia solani, 5-day-old cotton seedlings (Gossypium hirsutum ‘Acala 4-42’) were supported by quartz sand on glass plates and inoculated with mycelium grown in liquid culture. The inoculum was placed in contact with sand saturated with water or solutions of test materials. Previous studies have shown that an exogenous supply of glucose (sand saturated with a 10 g/liter solution) reduced disease severity and repressed the production of pectolytic enzymes by R. solani in vitro. When the sand was saturated with a solution of 3-methyl glucose (5 g/liter), disease development was completely inhibited. Growth of the pathogen was somewhat reduced in the presence of 3-methyl glucose (3-0-MG), but this was overcome by addition of a small amount of glucose (2.0 g/liter). The action of 3-MG differs from that of glucose. The pathogen readily grows on the surface of a cotton stem in both cases. With 3-0-MG, however, infection cushions did not form. The production of pectolytic enzymes in vitro is not repressed by 3-0-MG. These results suggest that 3-0-MG interferes with infection cushion formation. The results indicate that pectolytic enzyme production is not the sole factor in virulence of R. solani.

Heated wax-emulsions with benomyl and 2,6-dichloro-4-nitroaniline for control of postharvest decay of peaches. J. M. Wells (ARS, USDA, Beltsville, Md.). Mean lesion diameters of peaches, inoculated with Monilinia fructicola or Rhizopus stolonifer and treated with heated (52-C) or unheated (24-C) dips or sprays for 10, 20, or 30 sec with 225, 450, or 900 ppm 2,6-dichloro-4-nitroaniline (DCNA) suspended in a water emulsion were significantly smaller than those on fruit similarly treated with DCNA suspended in water. Similar wax-benomyl treatments at 10, 20, or 100 ppm were also more effective than benomyl treatments alone. Heated DCNA treatments, whether as dips or sprays, with or without wax, were more effective than unheated treatments. Heat improved the effectiveness of all benomyl-dip or of 10-sec wax spray treatments. Means of per cent decay of noninoculated peaches and nectarines due to Monilinia and Rhizopus infections were significantly less when treated with a 10-sec heated wax spray containing 450 ppm DCNA and 100 ppm benomyl than with a 3-sec unheated spray.

Isolation of Alternaria alternata from Nicotiana species. R. E. Wiley & H. W. Sprouse, Jr. (USDA, N.C. State Univ., Raleigh). We determined the incidence of the hop blight organism, Alternaria alternata, on leaves of susceptible (C-298) and resistant (PD 121) cultivars of field-grown, flue-cured tobacco, and on leaves of Nicotiana rustica, N. glutinosa, and N. sylvestris. Discs (9-mm diam) were cut from apparently healthy leaves surface-disinfected with 1% NaOCl for 1 min, rinsed in tap water, and placed on either rose-bengal streptomycin or Czapek + 6% NaCl agar or both. No measurable host differences were noted as the fungus was recovered from 97 to 100% of the discs cultured regardless of host. In a study of the effect of leaf age on the frequency of isolation of the fungus, leaf discs were randomly cut from leaves of C-298 and PD 121 in the plant bed before and after the ferbam applications for blue mold control were discontinued, and at 10 weekly intervals after the treatments. The percentage of leaf discs from which A. alternata grew was 0 when ferbam was applied, 29% after it was discontinued, 64-94% for 3-week-old leaves, and 93-100% for 7-week-old leaves (mature). Thus, the frequency of isolation of A. alternata is high in tobacco leaves of all ages, except in those protected by selected chemicals.

Ultrastructural investigation of susceptible and resistant isolines of soybean inoculated with Xanthomonas. W. P. Wergin & M. E. Mace (ARS, USDA, Beltsville, Md.). A fine structural study was made of the host-parasite interaction between soybean inoculated with Xanthomonas phaseoli var. sojevis. Fully expanded leaves of susceptible (Clark) and resistant (Clark 63) isolines of soybean were spray-inoculated with log-phase cultures of the bacterial pustule pathogen. One day after inoculation, examination of these isolines revealed no hyperplasia of the cells in the mesophyll or superficial cells. Sections of inoculated leaves showed the development of the new cells adjacent to the wall cells of the spongy parenchyma cells in the lower substomatal chambers. Regions of the wall of the abraded by the bacteria appeared to undergo gradual hydrolysis. Within 48 hr, progressive loosening and dispersal of wall microfibrils resulted in the rupture of several mesophyll cells. Three days after inoculation, structural differences between the isolines could be observed in the remaining mesophyll cells. In susceptible plants, the cells enlarged, pushed the lower epidermis outward, and formed a pustule. In the resistant plants, cellular expansion did not occur; however, the bacteria within the cells were significantly smaller and less abundant than those present in the experimental plants. Since these organisms are associated with protein synthesis and extracellular secretion, they may be related to a mechanism of disease resistance.

Lima beans resistant to Phytophthora phaseoli, strain C. R. E. Wester, V. J. Fisher, & V. L. Blount (ARS, USDA, Beltsville, Md., Univ. of Nebraska). Of 34 Lima bean plant introductions from Central America that were tested in October 1970 at Beltsville for resistance to downy mildew, strain C, four showed complete resistance. These introductions were also resistant to downy mildew strains A and B. The others tended to be completely susceptible.
A few seedlings survived inoculation, but apparently these were escapes, as all progeny from them were completely susceptible to strain C. Two of the resistant introductions were white-seeded (P.I. 195342 and P.I. 197025); two were black-seeded (P.I. 195345 and P.I. 197023). The white-seeded introductions closely resembled small-seeded commercial varieties, and bloomed earlier than the black-seeded introductions. The white-seeded introductions were chosen as parents for crossing to Dover, U.S.D.A. line 1068, Del. line 255-84, and U.S.D.A. lines 169, 269, and 369. F1 seeds from these crosses were resistant to all known strains of downy mildew (A, B, C). The F2 generation segregated in a ratio of three resistant to one susceptible, indicating that resistance is controlled by a single dominant gene as reported earlier for strains A and B.

Host responses and ultrastructural changes in maize leaves infected with Colletotrichum graminicola. H. Wheeler, D. J. Politis, & A. S. Williams (Univ. Ky., Lexington). Young maize plants (10 days old) sprayed with black-spore suspension (2 x 10^8 spores/ml) from an isolate of Colletotrichum graminicola obtained from naturally infected maize developed visible lesions within 35 hr, and many were killed within 5 days. Older plants (15-20 days) usually survived, but lower leaves were killed and mailes inoculated formed on upper leaves. All maize cultivars tested (14 commercial hybrid lines and one inbred line) were susceptible, but some were damaged more than others. Appressoria formed within 5 hr, epidermal cells were invaded within 8 hr, and acervuli with setae developed within 24 hr after inoculation. Spores, germ tubes, and appressoria were no unusual ultrastructural features. The earliest host response was an increase in electron density and a swelling of the cell wall beneath appressorium. As penetration progressed, many lomasomalike wall structure developed. The ability of this isolate to attack intact, nonwounded tissues and to kill young seedlings indicates that it is more pathogenic to maize than others of C. graminicola previously found in the USA.

Synergistic effect between Heterodera schachtii and Pythium ultimum on damping-off of sugarcane vs. additive effect of H. schachtii and P. aphanidermatum. E. D. Whitney (ARS, USDA, Salinas, Cal.). Damping-off tests of hybrid sugarcane inoculated with H. schachtii-P. ultimum and H. schachtii-P. aphanidermatum combinations were run simultaneously at various inoculum levels. The effect of the first combination on damping-off was synergistic but the second was additive. The synergistic effect was apparent at most inoculum levels and for both pre- and postemergence damping-off. In four tests, the presence of the nematode (10 larvae/g of soil) had reduced damping-off by an average of 260% of 10 days after planting. No reduction in seedling numbers resulted from the nematode alone. The mean number of infection centers per flat of beet was approximately the same whether the nematode was present or not. This suggested that the synergistic effect was due to increased rate of spread of the fungus in the soil around the infection centers when the nematode was present.

Inheritance of resistance to Xanthomonas campestris in cabbage. P. H. Williams & T. Stauber (Univ. Wis., Madison). Resistance in cabbage to black rot, caused by Xanthomonas campestris, was obtained from the Japanese cultivar, Early Fuji. When bacteria were inoculated through the hydathodes, resistance was expressed as either minute dark brown lesions or no visible injury. Lesions were obvious on the back surface of the leaves and were clearly visible from the underside. Resistance was not transferred to the backcross progenies, resistance was found to be controlled by one major recessive gene, r, the expression of which in the heterozygous condition was influenced by one recessive and one dominant modifier gene. Of over 350 cultivars and breeding lines screened for black rot resistance, none contained the major r gene for resistance; however, some contained the modifier genes.

Comparison of peroxidase and o-diphenol oxidase in postharvest apple decay by Penicillium and Phytophthora. D. M. Wilson & W. B. Darby (Univ. Md., Burlington). McIntosh apples inoculated with Penicillium sp. or Phytophthora sp. were used. Penicillium causes a light soft rot. As P. obscura decay progresses, the skin is brown and the pulp light; later the skin turns black and the flesh darkens. Peroxidase, o-diphenol oxidase, and catalase activities were measured using skin and pulp extracts of healthy apples, decayed areas, uninfected areas, and samples from the first 5 mm of firm tissue starting at the edge of visible change. Using gaiacol and H2O2 as substrates, peroxidase activity was low in decayed areas. Peroxidase activity in edge or uninfected tissue and skin of bothHOST increased 2.5 times over controls. o-Diphenol oxidase activity using 3,4-dihydroxyphenylalanine as the substrate was undetectable in rotted areas. When P. obscura-rotted apples were darkening, o-diphenol oxidase activity was again present. Inhibition of o-diphenol oxidase by low molecular weight substances was seen in edge or near necrotic Penicillium-infected tissue. Catalase was absent in rotted tissues. Soluble protein decreased with P. obscura only. These results indicate that enzymes in uninfected tissue respond differently during pathogenesis. Inhibition of o-diphenol oxidase by Penicillium may contribute to the light color; the dark P. obscura mycelia are responsible for most of the black color in black rot.

Relationship of basidiospore germination of Fomes igniarius to host specificity. J. E. Winch & P. Mantion (State Univ. N. Y., Coll. Forest., Syracuse). Basidiospores collected from fruiting bodies of Fomes igniarius on aspen, birch, beech, and butternut were used to inoculate birch, beech, aspen, and maple in all possible combinations. Spores were introduced into holes drilled into the sapwood of living trees in the field. Spores also were placed on cellulose membranes in contact with freshly cut cross sections of stem pieces and were applied directly to cross sections of wood. In both laboratory tests, a short piece of 6-mm tygon tubing, closed at the upper end, was placed over the inoculation site forming a miniature chamber to maintain favorable relative humidity. Germination was determined after 1 week. In the field, germination occurred when spores from fruiting bodies on aspen were inoculated into aspen and failed with all other combinations. Spore suspensions placed on stem sections with membranes germinated in all combinations used, but an initial laboratory test using the spore source from aspen indicated that germination does not occur when spores are placed directly on the cross sections of the 4 hosts.

Electron microscopic studies on the effects of tetracycline HCl on the mycoplasmalelike bodies in corn stunt and aster yellows-infected plants. B. S. Wolanski, M. Klein, & K. Marako-Moroschi (Boyle Thompson Inst. Plant Res., Yonkers, N.Y.). The effect of tetracycline HCl on the presumptive agents of aster yellows and corn stunt disease was studied by electron microscopy of thin sections of treated aster and corn plants. The effects of the antibiotics on the morphology of the mycoplasmalelike bodies could be observed 24 hr after treatment, and were most pronounced after 48 hr. After 72 hr, most of the phloem tissue examined was almost free of mycoplasmalelike bodies; any still present were in obvious disruptions or in disintegration of the morphology of mycoplasmalelike bodies resulting from the tetracycline treatment included condensation of the granular and fibrous contents of the structures into sporelike masses. This was followed by complete emptying of the bodies and transection of their membranes. Tetracycline treatment suppressed symptoms of the disease in asters for a period of 2 weeks, at the end of which time typical mycoplasmalelike bodies could again be found in the phloem tissue.
Regulation of the perfect stage in Fusarium roseum 'Graminearum' by F-2 (secalalene). J. C. Wolfe & C. J. Mukouchi (Univ. Minn., St. Paul). Fusarium roseum 'Graminearum' was seeded on 1.5-cm discs of Coons' medium agar. F-2 in quantities ranging from 10^-3 to 10^-4 µg was delivered onto the medium in diethylthioldimethylsulfoxide (9:3, v/v). In general, amt from 10^-4 to 10^-2 µg enhanced the production of perithecia, whereas amt from 10^-1 to 1 µg inhibited perithecial production. The period of effective inhibition of perithecial production was longer when amt of 10 µg and greater were used. The time of F-2 application to the developing colony was important. Greater amt of F-2 were needed to inhibit production of perithecia when introduced into the culture day after seeding than 3 days after seeding.

Identification of ozone-type air pollution injury to vegetation in Philadelphia. F. A. Wood (Pa. State Univ., Univ. Park). Exposes of 35 species of broad-leaf woody plants to ozone (O3) during the past 5 years indicated that chlorotic-to-purple stippling of the upper leaf surface of middle-aged to older leaves was the usual response. Recently established vegetation in Independence National Historical Park, Philadelphia, Pa., was examined 4 times during the summer of 1970. Purple stippling of the leaf surface was observed on American basswood, tree of heaven, and a Prunus sp. in July. In addition, ozone-type stippling was observed on Carolina silverbell, English oak, European mountain ash, flowering dogwood, grape, hawthorn, mulberry, and rhododendron in August. Forty white ash seedlings were planted as O3 indicators at seven locations in the park. Stippling typical of O3 injury developed on these plants within 4 weeks after planting. Fluoride-type symptoms also were observed, and fluoride levels in some of the plants indicated a fluoride problem, but ozone-type stippling was the most common symptom. These results substantiate the philosophy that one should examine the relative sensitivity of an array of species in the laboratory and then survey for symptoms in the field rather than attempt to elucidate the specific cause of each symptom complex as it is encountered in the field.

Genetic complementation between the nucleoprotein components of the cowpea mosaic virus, H. A. Wood & J. B. Bancroft (Boyce Thompson Inst., Yonkers, N. Y., Purdue Univ., Lafayette, Indiana). The bottom (B) and middle (M) nucleoprotein components of cowpea mosaic virus strain Vu (wild isolate) interact, resulting in increased levels of infectivity. Two local lesion mutants, designated 3d and 7b, were isolated following nitrous acid treatment. The 3d nuclei had an altered M-nucleoprotein to B-nucleoprotein ratio. In systemic assays, homologous and heterologous mixtures of the wild and mutant B and M components were infective at a 100-fold lower concentration than the individual components in the mixtures. Mixed component (MC) isolates from the infectivity dilution-end points produced only one lesion type. The lesion types and component ratios of the MC isolates were generally the same as the isolate from which the B component in the mixture was taken. However, the MC isolates arising from 3d-B and 7b-M component mixtures had wild lesion types and components. The nucleoprotein components from this MC isolate were mixed and assayed with the wild and mutant B and M components. The backcrossed-MC isolates had lesion types and component ratios which indicated the occurrence of genetic complementation, as the M component had the same properties as the original 3d-B and 7b-M components, respectively.

Use of dew point and ambient temperature differences to explain epidemic development of mushroom bacterial blotch. P. J. Wreger (Pa. State Univ., University Park). Bacterial blotch of the commercial mushroom is caused by Pseudomonas toliasi. All commercially grown strains of Agaricus bisporus are susceptible to the disease. The pathogen is soil-borne and a normal component of casing (soil) microflora, so disease occurs when environmental conditions permit. Disease occurrence and development were studied in two experiments in which PSU 310 was the suspect. Temperature-dew point in the casing room did not exceed 2° F and relative humidity did not vary more than 4%. Mushrooms were harvested daily, scored for disease incidence, and weighed. Data from harvests were summarized weekly. Disease development was measured by calculating four values. Dew points were computed hourly from temperature and RH values. Disease development was compared with various dewpoint/ambient temperature (DP/AT) differentials. When compared with days of DP/AT differentials of 3° F or more for 15 hr or more, disease increased when only 2 and 3 days, respectively, occurred during two periods of 21 days each. Disease was constant when 16 such days occurred during a different 21-day period. Disease decreased when such days occurred every day during a 14-day period.

The lack of effect of phosphate inhibitor on retention of tobacco mosaic virus by inoculated bean leaves. J. C. Wyatt & J. G. Shaw (Univ. Ky., Lexington). Pinto bean leaves rubbed with 14C-labeled viral inhibitor from Phytolacca americana retained from 5 to 15% of the label after thorough rinsing with water. The percentages varied with experiments, but did not vary with changes in concentration of inhibitor. The amount of inhibitor retained was not altered by the presence of up to 10 mg/ml of TMV in the inoculum. The amount of 32P-labeled TMV retained by an inoculated leaf after rinsing was not significantly affected by the addition of inhibitor to inocula. Nor did the presence of inhibitor affect the amount of nucleic-acid-susceptible 32P-RNA found in extracts of leaves 10 min after inoculation with 32P-labeled TMV. Inhibition of infection, therefore, does not appear to depend on alteration of the amount of TMV retained by inoculated leaves or alteration of the amount of uncoated TMV-RNA in extracts of the leaves.

Selective stimulation of ion uptake by Helminthosporium carbonum toxin. O. C. Youker & R. P. Scheeper (Mich. State Univ., E. Lansing). Susceptible corn roots from 4-day-old plants were treated for 4 hr with the host-specific toxin of Helminthosporium carbonum (2 µg/ml), then placed in NO3- solutions (0.04-10 mM) for 30 min. Control roots absorbed 200-300 nmoles NO3- g fresh wt/hr, whereas toxin-treated roots absorbed up to 3 times that amount. The same toxin concentration did not stimulate ion uptake by roots inoculated with the same amount. Ion uptake appears to be caused by increased influx rather than decreased efflux or decreased NO3- reduction, because a 50-min desorption period gave slight but equal loss of NO3- from treated and control roots. NO3- uptake by toxin-treated and control roots was greatly reduced at 5 C. Na+, K+, Ca2+, and Cl- were not required for toxin-stimulated NO3- uptake. Toxin increased Na+ absorption from 0.2-20 mM solutions by 50-60%. Toxin did not affect, or slightly inhibited, absorption of K+, SO42-, and PO43-, and did not affect P incorporation into perichorial acid-soluble and insoluble organic materials. Toxin did affect ion efflux under the conditions used. Enhanced absorption of NO3- and of Na+ was not required for toxicity because toxin affected susceptible tissues in the absence of either ion. Data indicate that toxin has selective effects on membrane properties, and does not generally derange the structure.

Production of tryptophol and indoleacetic acid in culture by Pythium debaryanum, K. Yoshit & D. J. Hagedorn (Univ. Wis., Madison). Pythium debaryanum produced 0.34 µg/ml indole-3-acetic acid (IAA) after 3 days of incubation in potato-dextrose broth (PDA). Addition of tryptophan to the medium invariably resulted in a significant increase, to 0.57 µg/ml in the production of IAA, suggesting that this fungus synthesizes IAA from tryptophan. Indoleacetonamide and indoleacetonitrile added to the medium at 0.01 mM increased IAA production to 2.5 and 20.0
μg/ml, respectively, but tryptophol and tryptamine had little effect. Therefore, indoleaceta mine and indoleacetonitrile may possibly be precursors for IAA synthesis by this organism. *Pythium debaryanum* also produced a neutral indole compound in PDA. The compound was extracted with ethyl acetate from the culture filtrate at pH 7.0, eluted with chloroform from silicic acid columns, and further purified by silicic acid partition chromatography with formate and a mixture of n-hexane and ethyl acetate as the stationary and mobile phases, respectively. It was identified as tryptophol by its melting point, chromatographic behavior, and ultraviolet and mass spectra.

Tobacco mosaic virus replication in suspensions of separated tobacco leaf cells. M. Zaitlen, A. O. Jackson, & R. I. B. Francki (Univ. Ariz., Tucson). Cells were separated from tobacco mosaic virus (TMV)-infected tobacco leaves with the aid of a pectinase procedure 3-4 days after inoculation. Suspensions of cells were presented with 3H-uridine. At various times after incubation in the light at 25°C, unlabeled virus was added as a carrier to facilitate the isolation of the virus from the cells and to determine the 3H-uridine incorporated into virus. By this means, virus was found to multiply at a constant rate for up to 45 hr of incubation (the longest time tested). Infected cells were also allowed to incorporate radioactive amino acids and uridine in order to detect virus-specific proteins and nucleic acids. Analysis by polyacrylamide gel electrophoresis revealed at least three high molecular wt virus-related proteins in addition to the coat protein, and two viral-induced RNA species plus the viral RNA. One of these RNA species is of very high molecular wt (ca. 4 × 10^6) and is considered to be the TMV replicative form. The other is a lower molecular wt RNA (2.5-3.0 × 10^5) of unknown origin and function.

Growth and movement of *Xanthomonas campestris* in natural fluids in relation to infection of cabbage. B. G. Zoller & T. Kosuge (Univ. Cal., Davis). Movement of *Xanthomonas campestris* in natural cabbage fluids was studied. Cells were grown in minimal medium (0.05% sucrose as carbon source) and resuspended (4 × 10^8 colony-forming units/ml) in conditioned water collected in the field from cabbage leaves. Ends of capillary tubes (2.38 × 10^-4 cm^2 in cross section) containing various field-collected samples of guttation fluid were immersed in the bacterial suspension. After 2-10 min at 24°C, up to 100 times as many bacteria had moved into capillary tubes containing field-collected cabbage guttation fluid than moved into capillary tubes containing the resuspending fluid. Conductivity bridge measurements revealed that the conductance of the guttation fluid was higher than that of the conditioned water from the leaves. Thus, the number of bacteria moving into the guttation fluid appeared to depend, at least in part, on the conductance of the fluids. Similar results were obtained in identical experiments using salt solutions having the same conductances as the natural fluids. The bacteria had a generation time of 2.7 hr at 26°C in the conditioned water leaf fluid, and 3.1 hr at 26°C in guttation fluid.