The severity of the damages to the cultivation and the difficulty of the chemical and agronomic control of the pathogen, led to purify the phytotoxin produced in the culture filtrates, to use the active compound in a programme of genetic improvements and to study its role in the pathogenesis.

The fungus was grown on a synthetic medium containing sucrose, L-asparagine, yeast extract, mineral salts and vitamins. The shaken cultures were incubated for 12 days at 26° C.

The culture filtrates, which were issted for toxicity on tomate cuttings, did not show the occurrence of any lipsophile phytoxists, here of an hydrophilic and polymeric compound, stable at 70° C for 30°. The culture filtrates were submitted to precipitation with cold accronce, the resulting phytoxoxis supermature was dialyzed against hidsittled water, using a nobe with a cut off of 3500 dathon. The resulting non-permature descripts was posified by ged filtration, affending a very active fras-

tion.

The chemical and electrophoretic analysis strongly suggested for such fraction a nature of an acidic glycoprotein with an apparent molecular weight of 13000-14000 dahon.

The toxin showed strong phytotoxicity still at 0.05 mg/ml. Work is in progress to further define the chemical and biological characterization of the active compound.

EFFECT OF FUSARIC ACID ON THE VACUOLAR MEMBRANE

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The effect of fusaric acid, a phytotoxin of Fusarium spp., on the tonoplast was investigated.

In the classical test of betacyanin release from red beet root disks, the toxin fed at millimolar concentrations induced a marked release of betacyanins, suggesting that the totoplast functionality in the intact cell was somehow affected (Radice and Peci, this same issue).

A possible direct effect on the tonoplast was thus tested both at morphologic and bishemical levels. Fusairs acid taps 10 3 mild bit and effect the interness of Acer pundaplatatus included wasoles, observed at the optical microscoral preparations from cultures cells of both Acer proaduplatatus indicated wasoles, observed at the optical microscoral preparations from cultures cells of both Acer proaduplatatus and Ace bidgets the disease, fusairs acid at concentrations from 0.25 mild to 5 mil inhibited (-10 to -35%) the internessentially innocellat TDEs, whereas it was infectively concelled. TDEs, whereas it was infectively

on another tonoplast enzyme activity, the K*-stimulated pyrophosphatase.

The structural analogues picolinic and nicotinic acids at the same concentrations showed a similar behaviour (inhibition of nitrate-sensitive ATPase and no effect on K*-stimulated PPase).

These results indicate that fusaric acid can directly affect some tonoplast function without inducing a general alteration of the state of the membrane.