Introduction

Plants are affected by a multitude of fungal diseases, which often cause high yield losses in all kinds of crops. Hofferek (1980) reported that spraying of barley plants with a root extract of barley can induce resistance toward the fungal pathogen *Puccinia striiformis* West. The plasma membranes of barley roots have an exceptionally high content of free phytosterols (57 mol% of total lipids in comparison to 7% in the membranes of spinach leaves). Analysis of the sterol composition showed that sitosterol accounts for nearly two thirds of the total amount (Rochester et al., 1987). Therefore it was not surprising that the resistance-inducing compounds of barley roots could be identified as 7α-hydroxysterol (1), 7β-hydroxysterol (2) and 7-oxosterol (3) (Schabdach et al., 1995). These oxysterols can only be found in the roots. Stem and leaves of untreated plants have a content below detection level of GC and GC-MS coupling. Their metabolism in barley roots has also been investigated (Vesper, 1998).

Fungal resistance was induced after spraying the leaves with 7-oxosterols. After leaf application of [3a,7α-2H]-7β-hydroxyosterol (6) and [3α-2H]-7-oxosterol (7) these sterols could never be detected in the roots. After root application the two 7-oxosterols were found in the leaves (König and Seifert, 1998). The 7-oxosterols applied via the roots should be transported through the xylem into leaves. The question is, whether oxysterols occur also in the phloem? In this publication we report on the analysis of phloem sap of barley collected by means of aphids. For example, this method has been used before for the detection of the indolizidine alkaloid swainsonine in honeydew excreted by phloem selective pea aphids (Dreyer et al., 1985). Instead of collecting the phloem exudate we have extracted and analysed the aphids. The aphid species used, *Sitobion avenae* Fabr., is a wide-spread herbivorous insect on grasses. It is a selective phloem feeder and commonly used for sampling pure phloem sap via the exuding stylet technique (Fisher and Frame, 1984). This practice of severing aphid stylets is often used to study phloem physiology (Barlow and McCully, 1972). In some cases other phloem-selective insects are also used, as for example leafhoppers and planthoppers on rice (Kawabe et al., 1980).

Results and Discussion

After root application of 7-oxosterol (3, 300 µg/ml) for 5 days the sterol could be detected by GC-MS in the roots and leaves of barley but not in the phloem sap feeding aphids. When the roots of barley plants were immersed for 2 days in a 1% aqu. ethanolic solution of [3α,7α-2H]-7β-hydroxyesterol (6) or [3α-2H]-7-oxosterol (7) (100 µg/ml per compound), 0.67 ± 0.08 mg fr. wt of 6 or 0.67 ± 0.08 µg/g fr. wt of 7 were found in the leaves (König and Seifert, 1998). In the experiment with the aphids, the concentration of 7-oxosterol...
Sterol in the phloem sap was too low to detect 3 in the aphids. If the roots of barley plants were cut off and the leaves were immersed for 9 days in a 0.5% aqu. ethanolic solution of 3 (300 μg/ml), 7-oxositosterol could be identified by GC-MS in the leaves and in the aphids. Since the aphids used are selective phloem feeders, 7-oxositosterol is a phloem-mobile compound.

After application of [7α-3H]-7β-hydroxysitosterol (4, 50 μg/ml) and [3α,6β-3H]-6α-hydroxylathosterol (5, 150 μg/ml) the radioactivity increased in the roots (4, 47.7 Bq/mg dry wt; 5, 5432 Bq/mg dry wt), leaves (4, 4.6 Bq/mg dry wt; 5, 267 Bq/mg dry wt) and aphids (4, 0.25 Bq/mg dry wt; 5, 11.3 Bq/mg dry wt) (Table I). The concentration of [3α,7α-2H]-7β-hydroxysitosterol (6) after exogenous application to the leaves was determined three days (0.39 ± 0.03 μg/g fr. wt) and for weeks (0.36 ± 0.04 μg/g fr. wt) after application (König and Seifert, 1998). The concentrations indicate the absence of any metabolization and deuterium-hydrogen exchange of 6 in the leaves. Therefore it is no reason to assume that the tritium of [7α-3H]-7β-hydroxysitosterol (4) or [3α,6β-3H]-6α-hydroxylathosterol (5) was completely transferred to other compounds and the detection of tritium in the aphids is based on artifacts.

**Table I. Radioactivity of barley plants and aphids after root application of [7α-3H]-7β-hydroxysitosterol (4) and [3α,6β-3H]-6α-hydroxylathosterol (5).**

<table>
<thead>
<tr>
<th>Compound</th>
<th>Radioactivity [Bq/mg dry wt]</th>
<th>Concentration [μg/mg dry wt]</th>
<th>Number of Aphids</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>4</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Roots</td>
<td>47.7</td>
<td>0.71</td>
<td></td>
</tr>
<tr>
<td>Leaves</td>
<td>4.6</td>
<td>0.069</td>
<td></td>
</tr>
<tr>
<td>Aphids</td>
<td>0.25</td>
<td>0.0037</td>
<td>300</td>
</tr>
<tr>
<td>Roots control</td>
<td>0.005</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leaves control</td>
<td>0.002</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aphids control</td>
<td>0.002</td>
<td></td>
<td>200</td>
</tr>
<tr>
<td><strong>5</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Roots</td>
<td>5432</td>
<td>4.28</td>
<td></td>
</tr>
<tr>
<td>Leaves</td>
<td>267</td>
<td>0.21</td>
<td></td>
</tr>
<tr>
<td>Aphids</td>
<td>11.3</td>
<td>0.0089</td>
<td>40</td>
</tr>
<tr>
<td>Roots control</td>
<td>0.13</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Leaves control</td>
<td>0.01</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aphids control</td>
<td>0.22</td>
<td></td>
<td>40</td>
</tr>
</tbody>
</table>

1 mg dry wt of aphids = 8.38 aphids, 4.
1 mg dry wt of aphids = 8.69 aphids, 5.
Estimated concentration of 4 in the phloem: 68 ng/ml.
Estimated concentration of 5 in the phloem: 202 ng/ml.

The application of radiolabelled sterols can prove the downstream transport of sterols together with photosynthates. The aphids on the barley leaf surface incorporated radioactivity by feeding for 9 days (4) or 7 days (5) of the phloem sap containing the tritiated sterols 4 or 5. It is possible to calculate the amount of sterol which was absorbed by a single aphid. Aphids in the first group (50 μg/ml of 4) contained a mean of 0.44 ± 0.02 ng of sterol and aphids of the second group (150 μg/ml of 5) contained 1.02 ± 0.37 ng. The oxysterol concentration in the phloem sap could be estimated as follows: Unattended aphid species excrete between 20 and 40 nl honeydew per aphid and hour (Heimbach, 1986; Völkl et al., 1999; Fischer et al., 2000). A similar honeydew production can be assumed for Sitobion avenae resulting in 480 – 960 nl honeydew per aphid and day. Assuming a value of 720 nl per aphid and day, an aphid of the first group fed approximately 6480 nl (9 days) and of the second group 5040 nl (7 days). The concentrations of 4 and 5 in the phloem sap of barley seedlings should be about 68 ng/ml and 202 ng/ml.

The results are very consistent considering the different sterol concentrations. Our experiments demonstrate that small amounts of oxysterols are
downstream transported through the phloem in barley plants. As expected, the major part of oxy-
sterols is transported from the roots to the leaves through the xylem. This is the first time to our
knowledge that 7-oxysitosterol and 6-hydroxylathosterol transport in the phloem of intact barley
plants has been reported.

So far, mainly the transport of photosynthates has been followed with different methods (Orlich
and Komor, 1989). The aphid technique used is doubtless one of the most precise methods for col-
lecting phloem sap but also one of the most time consuming. Another common practice is to cut
open exposed phloem vessels and collect the exu-
date. For example, this procedure was used for the analysis of the translocation of quinolizidine alkalo-
oids in Lupinus albus L. (Wink and Witte, 1984). The presence of the alkaloid ricinine in the phloem sap of Ricinus communis L. seedlings has also been proven by using this technique (Holf-
felder et al., 1998). Cutting open vessels has always the risk of contamination with other liquids than phloem sap. The EDTA-method has been used for studies of phloem transport of tropane and pyri-
dine alkaloids (Kitamura et al., 1993). This tech-
nique does not deliver pure phloem sap, but it is
popular due to its simplicity. It is also quite un-
likely to work with a substance which is very well
transported in the phloem, but not at all in the xylem, as was the case with atropine in Duboisia
myoporoides (Kitamura et al., 1991).

After root application of [3α,7α,2H]-7β-hydroxy-
sitosterol (6) and [3α-2H]-7-oxositosterol (7)
(100 μg/ml per compound) the ratio of the endoge-
nous sterols stigmasterol and sitosterol in the barley
leaves was changed to 1.71 in comparison with 0.52
in the control plants (König, 1997). Nutrient defi-
ciency caused a similar change in the stigmasterol/
sitosterol ratio in wheat plants (König et al.,
1996). Salt-stressed barley roots have a higher
membrane sterol content than the controls (Brown
and DuPont, 1989). Apparently, there are many
factors which may alter the amount and composi-
tion of the sterols in the roots and leaves of barley
and may in some cases trigger fungal resistance.

However, further activities of these sterols may
also include interactions with life-history traits of
herbivorous insects as well. Behmer and Grebenok (1998) were able to demonstrate that
differences in the amount of definite sterols in arti-
ficial diets contributed to the survival in the dia-
mondback moth Plutella xylostella. Moreover, the
biochemical deficiency for sterol production in in-
ssects could be used in feeding experiments using
the approximation of sterol uptake of aphids as
calculated above to study the impact of different
sterol concentrations on the preference and per-
formance of these herbivorous species.

A study reported on different reproduction rates for aphids on different warm-season grasses,
which are important transitional hosts (Kieck-
hefer, 1984). Interestingly, with young seedlings
the differences were not so pronounced, but on
mature plants there was a significant discrimina-
tion between different grasses. The same has been
reported in a study comparing wheat, barley,
maize and sorghum as hosts for four different
aphid species (Kieckhefer and Gellner, 1988). All
these studies are quite important because most
aphids are vectors, which can transmit different
plant viruses. Therefore, feeding deterrence can
prevent a plant from getting infected with a virus.
However, a virus-disease can cause stress to the
plant. The consequences of that could be increased
or changed sterol content, population control of
insects or fungal resistance.
Experimental

Seeds of barley (*Hordeum vulgare* L. cv “Mammut”) were germinated on moist filter paper and after germination transferred onto perforated polystyrene plates. The seedlings were grown hydroponically in Münchner/2 solution (3 mM KNO$_3$, 2 mM Ca(NO$_3$)$_2$, 4 mM MgCl$_2$, 0.5 mM KH$_2$PO$_4$, 0.06 mM EDTA-Fe(III)-Na salt, 0.023 mM H$_3$BO$_3$, 0.005 mM MnCl$_2$, 0.4 μM ZnSO$_4$, 0.2 μM CuSO$_4$, 0.05 μM Na$_2$MoO$_4$) for two weeks. The barley plants were separated in root-collected, frozen in liquid N$_2$ in an insect cage for 5 days. At the end, aphids were consumed (approximately 1 ml/day per plant). This was continued until the end of the experiment (4, 9 days; 5, 7 days). Aphids and plants were treated as described above. After 9 days the experiment was finished and the same concentration was added (approximately 1 ml/day per plant). This was continued until the end of the experiment (4, 9 days; 5, 7 days). Aphids and plants were harvested and dried as described above.

Treatment with 7-oxositosterol (3)

The nutrient solution of barley plants consisted of Münchner/2 solution containing 0.25% of detergent (Palmolive®), 0.5% of EtOH, 300 μg/ml of 3 and was prepared as follows: 7-Oxositosterol (3) was dissolved in EtOH and diluted with the detergent containing Münchner/2 solution. The roots of the seedlings were placed in the Münchner/2 solution. After the nutrient solution was consumed (approximately 1 ml/day per plant), fresh solution was added. 40 adult aphids were put on each group of 15 plants at the same day as the sterol was applied. Plants were kept in a closed insect cage for 5 days. At the end, aphids were collected, frozen in liquid N$_2$ and dried at 60° for 2 days. The barley plants were separated in root- and shoot-parts and also dried at 60° for 2 days. Controls were grown in the same nutrient solution as described above but without 3.

Barley plants, grown in potting soil for two weeks, were cut off under water just above the roots. The water contained 0.01% detergent to avoid air-bubbles which could plug the xylem vessels. The shoot parts were placed in the above described Münchner/2 nutrient solution which contained 300 μg/ml of 3. The level of solution was maintained constant during the whole experiment. 30 adult aphids were put on each group of 15 plants at the same day as the sterol was applied. After 9 days the experiment was finished and the aphids and barley plants were treated as described above.

Aphids and plants were ground to a fine powder with mortar and pestle and were extracted four times with CHCl$_3$. The CHCl$_3$ extracts were evaporated and the residues purified by TLC (20 × 20 cm silica plates, cyclohexane/EtOAc 1:1 v/v). The bands with the $R_f$ 0.45–0.50 were scraped off and eluted with CHCl$_3$. The combined eluates were evaporated to dryness under reduced pressure. Samples were trimethylsilylated with N-methyl-N-trimethylsilyl trifluoroacetamide for 12 h at room temp. and analysed by GC and GC-MS. 3-O-trimethylsilyl-7-oxositosterol was identified by comparison of $R_T$ and MS with the reference compound.

Capillary GC measurements were carried out under the following conditions: Fused silica-glass column DB1 (J&W Scientific, 29 m × 0.32 mm, 0.1 μm film thickness); injector temp. 280°; column temp. 3 min 80°, heat rate 3° min$^{-1}$ to 280°; FID temp. 290°; carrier gas H$_2$, flow rate 2 ml min$^{-1}$, split ratio 1:20. GC-MS measurements were performed on an instrument coupled to a GC fitted with a fused silica-glass column DB-5 (30 m × 0.32 mm, 0.1 μm film thickness); inj. temp. 280°; carrier gas H$_2$, flow rate 2 ml min$^{-1}$, splitless injection.

3-O-Trimethylsilyl-7-oxositosterol

$R_T$ (GC, DB1) 28.93; MS m/z (rel. int.): 500 (M$^+$, 65), 485 (12), 444 (6), 410 (18), 395 (26), 281 (34), 207 (100), 129 (15).

Treatment with [7α-3H]-7β-hydroxysitosterol (4) and [3α,6β-3H$_2$]-6α-hydroxylathosterol (5)

The $^3$H-labelled sterols (Schröder, 1995) were purified with prep. TLC (cyclohexane-EtOAc 1:1 v/v). A 50 μg/ml solution of 4 (specific radioactivity 67 kBq/mg) consisted of 0.5% EtOH and 0.1% detergent (Palmolive®) in Münchner/2 solution. A 150 μg/ml solution of 5 (specific radioactivity 1.27 MBq/mg) consisted of 0.5% EtOH and 0.25% detergent in Münchner/2 solution. Control plants were nourished with Münchner/2 solution containing 0.5% EtOH and 0.1% detergent. The roots of 2-week-old barley plants were placed in the appropriate solution, ensuring that the leaves had no contact with the solution. After the radioactive labelled sterol solution was completely consumed, fresh solution with the same unlabelled sterol in the same concentration was added (approximately 1 ml/day per plant). This was continued until the end of the experiment (4, 9 days; 5, 7 days). Aphids and plants were harvested and dried as described above.
Dried aphids and plant parts were weighed and wrapped in ash-free filter-paper. The samples were oxidized in a sample oxidizer which collected the total combustion water. The tritium activity was counted in a liquid scintillation counter. A Packard Sample Oxidizer Model 307 was used according to the manual with a flow setting of 18 ml of monophase S and 2 ml of permafluor E. N₂ and O₂ pressure were set at 5 bar and the carburation time was 45 sec. The yield of tritiated water was determined by combustion of known standards (on filter-paper as well). The tritium content was counted with a Packard Liquid Scintillation Analyzer 2500 TR. Counting time was 10 min, with samples kept in 30 ml HD-PE scintillation tubes.


