Xanthone biosynthesis in *Hypericum perforatum* cells provides antioxidant and antimicrobial protection upon biotic stress

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**ABSTRACT**

Xanthone production in *Hypericum perforatum* (HP) suspension cultures in response to elicitation by *Agrobacterium tumefaciens* co-cultivation has been studied. RNA blot analyses of HP cells co-cultivated with *A. tumefaciens* have shown a rapid up-regulation of genes encoding important enzymes of the general phenylpropanoid pathway (PAL, phenylalanine ammonia lyase and 4CL, 4-coumarate:CoA ligase) and xanthone biosynthesis (BPS, benzophenone synthase). Analyses of HPLC chromatograms of methanolic extracts of control and elicited cells (HP cells that were co-cultivated for 24 h with *A. tumefaciens*) have revealed a 12-fold increase in total xanthone concentration and also the emergence of many xanthones after elicitation. Methanolic extract of elicited cells exhibited significantly higher antioxidant and antimicrobial competence than the equivalent extract of control HP cells indicating that these properties have been significantly increased in HP cells after elicitation. Four major de novo synthesized xanthones have been identified as 1,3,6,7-tetrahydroxy-8-prenyl xanthone, 1,3,6,7-tetrahydroxy-2-prenyl xanthone, 1,3,7-tri-hydroxy-6-methoxy-8-prenyl xanthone and paxanthone. Antioxidant and antimicrobial characterization of these de novo xanthones have revealed that xanthones play dual function in plant cells during biotic stress: (1) as antioxidants to protect the cells from oxidative damage and (2) as phytoalexins to impair the pathogen growth.

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1. Introduction

*Hypericum*, a genus of the family Clusiaceae, is widely used in traditional medicine throughout the world since ancient times. The genus is known to produce several xanthones (Dias et al., 2000, 2001; Dias, 2003; Ferrari et al., 2005; Tanaka and Takaishi, 2006).

Xanthones are a class of polyphenolics that exhibit well-documented pharmacological properties, mainly due to their oxygenated heterocyclic nature and diversity of functional groups. They have been described as strong scavengers of free radicals (Jiang et al., 2004). In addition, many of the xanthones have been reported to be active against bacteria including methicillin/multi-drug resistant *Staphylococcus aureus* (Rukachaisirikul et al., 2003, 2005; Sukpondma et al., 2005; Xiao et al., 2008), vancomycin resistant *Enterococci* (Sakagami et al., 2005), *Mycobacterium tuberculosis* (Suksamrarn et al., 2003), etc. Some xanthones even surpass the antimicrobial activity of traditional antibiotics (Iinuma et al., 1996; Xiao et al., 2008). Other pharmacological properties of xanthones include anti-inflammatory (Banerjee et al., 2000), cancer-chemopreventive (Ito et al., 2003), hepatoprotective (Tian et al., 2005), cardiovascular protective (Jiang et al., 2004), selective inhibition of cyclooxygenase-2 (Zou et al., 2005), inhibition of platelet-activating factor (PAF)-induced hypotension (Ishiguro et al., 2002; Oku et al., 2005) and cytotoxic activities (Yimjdo et al., 2004; Boonsri et al., 2006; Suksamrarn et al., 2006).

Plant cells respond to pathogens predominantly by mobilizing their secondary metabolites, which tend to protect the plant cells from pathogen attack. In this context, Beerhues and Berger (1995) observed an increase in xanthone accumulation in suspended cells of *Centaurium* species upon elicitation with yeast extract and methyl jasmonate (Mej). Conceição et al. (2006) observed a similar response (xanthone accumulation) in *Hypericum perforatum* suspended cells elicited with *Colletotrichum gloeosporioides* cell wall extracts. The induced xanthone accumulation was further increased (by at least 12-fold) when the HP cells were primed with either Mej or salicylic acid (well known defense signaling compounds in plants), an important observation, which explains possible role of xanthones in plant cells under biotic stress.

Here, we show that the antioxidant and antimicrobial potentials have been significantly increased in HP cells after elicitation with *Agrobacterium tumefaciens* due to the rapid up-regulation of xanthone metabolism. Putative roles of xanthone metabolism in HP cells under conditions of biotic stress are discussed.
2. Results and discussion

2.1. Induction of xanthone biosynthesis is a defense response

2.1.1. Xanthone profile of HP cells was altered after co-cultivation with *A. tumefaciens*

Major classes of phenolic compounds produced by cultured HP cells under normal conditions are flavonoids and xanthones (Fig. 1a, F, flavonoid and X, xanthone). Though the flavonoid profile remained unaltered after elicitation, xanthone profile has significantly changed within 24 h (Fig. 1b). With new xanthones, the total xanthone content increased 12-fold (Fig. 2) indicating that xanthone biosynthesis has potential role in the biotic interaction. Similar patterns of xanthone accumulation has also been observed in HP cells that were elicited with fungal extracts or primed with plant defense signaling compounds such as MeJ and salicylic acid (Conceição et al., 2006). Likewise, xanthones were seen accumulated when cell cultures were treated with MeJ and yeast-extract in *Centaurium* species (Beerhues and Berger, 1995), *H. androsaemum* and *Centaurium erythraea* (Abd El-Mawla et al., 2001); the latter cell cultures exhibited new xanthones after MeJ treatment. All these observations support the participation of xanthones in plant defense response.

Although mangiferin (peak denoted as ‘M’) found along with few other unidentified xanthones in both control and treated cells, new xanthones appeared in the HP cells treated with *A. tumefaciens* (Figs. 1a and b). Amongst the up-regulated xanthones, four were identified (Fig. 1b, peaks 1–4, see Fig. 1c for structural formulae)

![Fig. 1. Phenolic profile of *H. perforatum* cells before and after co-cultivation with *A. tumefaciens*. HPLC chromatogram showing the major phenolic compounds (F, flavonoids; X, xanthones and M, mangiferin) in control (a) and *H. perforatum* cells co-cultivated with *A. tumefaciens* for 24 h (b). Note the appearance of many new compounds after elicitation. Peaks 1, 2, 3 and 4 were identified as 1,3,6,7-tetrahydroxy-8-prenyl xanthone, 1,3,6,7-tetrahydroxy-2-prenyl xanthone, 1,3,7-trihydroxy-6-methoxy-8-prenyl xanthone and paxanthone, respectively. Structural formulae of these xanthones are shown in (c).
as 1,3,6,7-tetrahydroxy-8-prenyl xanthone (peak 1), 1,3,6,7-tetrahydroxy-2-prenyl xanthone (peak 2), 1,3,7-trihydroxy-6-methoxy-8-prenyl xanthone (peak 3) and paxanthone (peak 4). Concentrations of those xanthones before and after elicitation are shown in Table 1. The major xanthone in the treated HP cells was 1,3,6,7-tetrahydroxy-8-prenylxanthone, which accounted for 23% of the total xanthones (Table 1). This was found to be the same compound that was elicited by Mej in H. androsaemum cell cultures (Abd El-Mawla et al., 2001), and, might be a common compound induced in both H. perforatum and H. androsaemum in response to the biotic stress.

2.1.2. Xanthone production in response to elicitation by A. tumefaciens is regulated by transcriptional control of enzymes participating in the xanthone biosynthesis pathway

BPS gene expression was found to be strongly up-regulated in HP cells upon treatment with A. tumefaciens as revealed by northern blot analysis (Fig. 3d). While BPS transcripts were undetectable in control HP cells, they accumulated within 4 h of A. tumefaciens challenge, and reached the maximum in 12 h and declined then on. A more or less similar pattern of expression was observed for PAL gene (Fig. 3b), which is participating in the biosynthesis of xanthone precursor and 4CL gene (Fig. 3c) that is related to the ubiquitous activation of general phenylpropanoid pathway following elicitation. The changes in the PAL mRNA levels agree quite well with the changes in the PAL activity level in elicited H. androsaemum cell cultures (Abd El-Mawla and Beerhues, 2002).

The rapidly induced expression of PAL, 4CL and BPS genes in HP cells co-cultivated with A. tumefaciens indicates that the xanthone biosynthetic pathway is activated immediately after sensing the pathogen. PAL, the first enzyme of the general phenylpropanoid pathway, catalyses the deamination of phe to cinnamic acid. Cinnamic acid, which is used in the synthesis of benzoic acids, the precursors of xanthone biosynthesis (Abd El-Mawla et al., 2001) thus, plays a key role in diverting photosynthate from primary metabolism to phenylpropanoid metabolism. BPS is an important enzyme, which catalyses the stepwise condensation of activated benzoic acids with three molecules of malonyl-CoA (Beerhues, 1996; Schmidt and Beerhues, 1997) and the resulting benzenophenones undergo intramolecular cyclization to give xanthones (Peters et al., 1998). The enzyme 4CL plays a key role in channelling carbon flow into diverse branch pathways of phenylpropanoid metabolism, which serves important functions in plant growth and adaptation to environmental perturbations (Ehlting et al., 1999).

Activation of phenylpropanoid metabolism has been described as an almost ubiquitous feature of plant response to stress conditions (Dixon and Paiva, 1995). Marked increase in PAL synthesis and corresponding mRNA accumulation occur in response to microbial or endogenous elicitors has been reported in many plant-pathogen systems (Sharan et al., 1998). Thus, xanthone biosynthesis in response to A. tumefaciens treatment is believed to be regulated by transcriptional control of enzymes participating in the pathway.

2.2. Xanthones protect HP cells from oxidative stress

2.2.1. ROS scavenging potential of HP cells increases after elicitation with A. tumefaciens

HP cells taken at different time intervals (0, 15, 30, 60, 120, 180 and 240 min) following A. tumefaciens challenge were incubated with H$_2$DCFDA, a probe that measures the cellular ROS levels internally. HP cells produced a biphasic oxidative burst after the challenge (Fig. 4a) an observation already reported (Franklin et al., 2008). Because of the highly cytotoxic and reactive nature of ROS, their accumulation must be under tight control (Shao et al., 2008). If not scavenged, the ROS would eventually lead to critical levels of H$_2$O$_2$ (Scandalios, 1993). Such levels are detrimental to

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**Table 1**

<table>
<thead>
<tr>
<th>Xanthone content (µg/g dry wt) of HP cells before and after elicitation.</th>
<th>Control HP cells</th>
<th>Elicited HP cells</th>
</tr>
</thead>
<tbody>
<tr>
<td>1,3,6,7-Tetrahydroxy-8-prenyl xanthone</td>
<td>0</td>
<td>950</td>
</tr>
<tr>
<td>1,3,6,7-Tetrahydroxy-2-prenyl xanthone</td>
<td>0</td>
<td>270</td>
</tr>
<tr>
<td>1,3,7-Trihydroxy-6-methoxy-8-prenyl xanthone</td>
<td>0</td>
<td>270</td>
</tr>
<tr>
<td>Paxanthenone</td>
<td>0</td>
<td>110</td>
</tr>
<tr>
<td>Total xanthones</td>
<td>320</td>
<td>4080</td>
</tr>
</tbody>
</table>
Intracellular reactive oxygen species (ROS) production in *H. perforatum* cells during co-cultivation with *A. tumefaciens* has been reported before (Franklin et al., 2008). The rate of membrane lipid peroxidation in untreated (control) HP cells doubled (from approximately 23 to 47 nmol MDA/g fr. wt) as the cultures aged (during the experimental period of 5 days, Fig. 4b); whereas in the HP cells that have been co-cultivated with *A. tumefaciens*, the level of membrane lipid peroxidation remained similar to the control on day one and thereafter significantly declined to about 70% of its initial value (16 vs. 23 nmol MDA/g fr. wt) over the period of 5 days, in spite of the intense oxidative burst observed. Moreover, HP cells retained 100% viability throughout the experimental with and without the pathogen. This observation has been reported before (Franklin et al., 2008).

Analysis of the culture medium of the elicited HP cultures (by xylenol orange method) did not show the presence of H$_2$O$_2$. This indicated that the intense oxidative burst produced internally in the HP cells was not transported outside of the cells. Absence of H$_2$O$_2$ in the extracellular medium further suggests that the ROS produced by HP cells might have been readily scavenged either before or immediately after their conversion into H$_2$O$_2$.

### 2.2.2. Increased ROS scavenging potential of elicited HP cells is due to induced xanthone biosynthesis

In order to determine whether the xanthones synthesized anew by the cells after elicitation (Fig. 1b) have ROS scavenging activity, we compared the antioxidant potential of the total phenolics of both the control and elicited HP cells that was extracted with MeOH from (co-cultivated with *A. tumefaciens* for 24 h) in two different antioxidant model assays (DPPH reduction and TBARS). Methanolic extracts of elicited cells showed a higher degree of radical scavenging activity (DPPH reduction) with a greater capacity for preventing synaptosomal lipid peroxidation, than equivalent extracts obtained from non-elicited (control) cells (Figs. 5a and b).

In DPPH reduction assay, the EC$_{50}$ values were 0.7 mg dry wt/ml and 0.44 mg dry wt/ml respectively for the control and elicited cells (Fig. 5a). Similarly in synaptosomal lipid peroxidation inhibition assay, the EC$_{50}$ values were 0.17 mg dry wt/ml and 0.04 mg dry wt/ml respectively for the control and elicited cells (Fig. 5b). Altogether, the free radical scavenging capacity was increased about 0.6-fold and the protection against membrane lipid peroxidation was increased about 3.25-folds in HP cells after elicitation. The increased antioxidant activity can possibly be attributed to the large amounts of xanthones produced in HP cells after elicitation.

In the present study, the first ROS burst (Fig. 4a) occurred relatively earlier (in 15 min) than the up-regulation of xanthone biosynthesis (BPS mRNA starts to accumulate at 4 h and attain maximum at 12 h) after treating with the bacterium. In fact, this burst was seen to be short-lived and might have been buffered by the existing phenolics (xanthones plus flavonoids). Our hypoth-
esis that the oxidative burst that occurred in the initial phase of the incompatible plant-pathogen interactions is responsible for the induction of defense reactions agrees with that of Arias et al. (2005). Moreover, ROS has complex downstream effects on secondary metabolism as signalling molecules (Chen et al., 1993; Vanderauwera et al., 2005; Foyer and Noctor, 2005). Hence, the first ROS burst could have acted as signal to prime the xanthone biosynthesis, and the massive xanthone production could be related to the second phase of ROS production, which is more intense and long-lasting, as we observed. During such state and thereafter, the cells would need additional antioxidant support conferred by xanthones produced anew, to achieve cellular ROS homeostasis.

2.2.3. De novo synthesized xanthones have antioxidant properties

In order to check the contributions of the de novo xanthones in the hiked antioxidant potential of elicited cells, we studied the antioxidant efficacies of the four xanthones identified in elicited HP cells methanolic extract (Fig. 1c). These xanthones showed anti-radical activity (DPPH reduction) to varying degrees (Fig. 6a) and were very effective in protecting against synaptosomal lipid peroxidation (Fig. 6b). The synaptosomal lipid peroxidation inhibitory activity of 1,3,6,7-tetrahydroxy-8-prenyl xanthone (EC\textsubscript{50} = 0.08 \mu g/ml, Fig. 6b) was the highest among tested xanthones. Complete inhibition of induced synaptosome lipid peroxidation by this xanthone even at considerably low concentration (1.0 \mu g/ml) indicates that even a small increase could have significant antioxidant protection for the host cells.

Previous studies (Pinto et al., 2005; Foti et al., 2005) have shown that xanthones could effectively suppress ROS production and prevent lipid peroxidation. Hence, we conclude that the up-regulation of xanthone biosynthesis aids HP cells to defend against biotic stress produced during the pathogenic challenge. This hypothesis could also explain the long-term viability/survival of HP cells during extended periods of pathogenic interactions (Franklin et al., 2007, 2008).

2.3. Xanthones provide antimicrobial protection to HP cells

2.3.1. Antimicrobial potential of HP cells increased after elicitation

It was remarkable to find the viability of A. tumefaciens gradually declining reaching (<1%) after 12 h of co-cultivation (Franklin et al., 2008) and the decline was about 50% at 5–6 h, at a time when xanthones start accumulating (Abd El-Mawla and Beerhues, 2002). To check whether the induced xanthone biosynthesis of HP cells is responsible for the reduced bacterial viability, we compared the antimicrobial potentials of methanolic extracts of control and cells co-cultivated with A. tumefaciens for 24 h period. Concentrations of methanolic extracts equivalent to 20 mg dry wt elicited HP cells/ml and 200 mg dry wt control HP cells/ml could kill 250 \times 10^7 A. tumefaciens cells within 24 h of incubation.

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**Fig. 6.** DPPH reduction (a) and synaptosomal lipid peroxidation inhibition (b) potentials of four identified xanthones; 1,3,6,7-tetrahydroxy-8-prenyl xanthone (1), 1,3,6,7-tetrahydroxy-2-prenyl xanthone (2), 1,3,7-trihydroxy-6-methoxy-8-prenyl xanthone (3) and paxanthone (4).

**Fig. 7.** Antimicrobial potential of H. perforatum cells after 24 h co-cultivation with A. tumefaciens. Methanolic extract of elicited H. perforatum cells showing higher antimicrobial activity than the equivalent extract of control H. perforatum cells, against A. tumefaciens colony formation.
Thus, the antimicrobial potential of HP cells has been increased 10 times after elicitation, which can be attributed to the xanthones accumulation.

Since the methanolic extract of control cells also showed inhibition of *A. tumefaciens* at higher concentrations, it can be concluded that antimicrobial compounds are present in control cells too; but at relatively low concentrations, acting as phytoanticipins (Dixon, 2001). With biotic elicitation, the amount of total xanthones increased and new xanthones are produced, eventually to achieve a higher biological response, i.e. killing of bacterial cells.

2.3.2. *De novo* synthesized paxanthone display antimicrobial activity

From earlier studies, it is found that many xanthones possess antimicrobial activities against bacteria (Suksamrarn et al., 2003, 2006; Pinheiro et al., 2003; Laphookhieo et al., 2006; Xiao et al., 2008). Hence, we studied the antimicrobial activities of the four xanthones identified in the elicited HP cells (peaks 1–4 of Figs. 2008). Hence, we studied the antimicrobial activities of the four xanthones against bacteria (Suksamrarn et al., 2003, 2006; Pinheiro et al., 2003; Laphookhieo et al., 2006; Xiao et al., 2008). Hence, we studied the antimicrobial activities of the four xanthones identified in the elicited HP cells (peaks 1–4 of Figs. 1b and c) by disc diffusion method. Of the four tested, only paxanthone (4) showed a clear inhibition zone against *A. tumefaciens* growth (Fig. 8a). So, not all of the xanthones produced by HP cells after elicitation have antimicrobial activity. The dose-dependent antimicrobial activity of paxanthone was studied by incorporating varying concentrations of paxanthone in *A. tumefaciens* suspension.

As seen in Fig. 8b, with the maximum concentration of paxanthone tested (200 µg/ml), we did not observe 100% reduction of *A. tumefaciens* viability. In contrast, a concentration of methanolic extract that is equivalent to 20 mg dry wt elicited cells/ml was sufficient to achieve 100% reduction in *A. tumefaciens* viability in 24 h (Fig. 7). Hence we believe antimicrobial compound(s) other than paxanthone or a synergism between this xanthone and other compounds of elicited cells might exist.

Pinheiro et al. (2003) had reported that assignuxanthone-B, a xanthone isolated from *Kielmeyera variabilis* showed only a moderate antibacterial activity (minimal inhibitory concentration = 25.0 µg/ml) against *Bacillus subtilis*, in contrast to the methanolic extract of the plant (minimal inhibitory concentration = 1.95 µg/ml). They found that the antimicrobial potential of assignuxanthone-B could be increased 4-fold, when combined with 1,3,5,6-tetrahydroxy-2-prenyl xanthone. Mixture of the two was able to inhibit *B. subtilis* at 6.25 µg/ml concentration. This observation demonstrates that the combination was active at much lower concentrations than each of the single compounds. Other evidences (Briskin, 2000) also indicate that many complex phytomedicines (plant extracts) exert biological or pharmaceutical effects through synergistic actions of several compounds.

3. Conclusion

In summary, we provide evidence for the activation of secondary metabolism in HP cells in general and xanthone biosynthesis in particular under elicitation by *A. tumefaciens* co-cultivation. We also present data that show the activation of xanthone biosynthesis having dual defense function in HP as described in the model (Fig. 9). On the one hand, xanthones constitute a powerful antioxidant system of protecting the host cells from self-generated ROS and subsequent oxidative damage. On the other hand, xanthones have also the potential to act as phytoalexin to help the host to impair and eventually kill the pathogen. In fact, both functions may help to prevent disease development by their concerted action.

4. Experimental

4.1. Growth and elicitation of HP cells

*H. perforatum* cell suspension culture was established from the variety ‘Helos’ (Richters seeds, Ontario, Canada) and maintained as per Dias et al. (2001). Five days after subculture, cells suspensions from several flasks were aseptically collected into a sterile beaker.
and the cell density was estimated in an improved Neubauer haemocytometer (Neubauer, Wertheim, Germany). After adjusting the cell concentration to about 5000 HP cells/ml with sterile MS medium, 80 ml aliquots were redistributed to each flask. A. tumefaciens stock was prepared as described before (Franklin et al., 2008). A set of 10 flasks, each containing 80 ml of this HP cell suspension, were inoculated A. tumefaciens such that to reach 250 × 10^6 bacterial cells/ml. Another set of 10 flasks each containing 80 ml of just HP cell suspension served as control. Treated and control cultures were incubated in growth chamber at 25 °C under a photoperiod of 16 h light. After 24 h (or at the time indicated) cells were harvested by vacuum filtration and freeze-dried.

4.2. Measurements of reactive oxygen species (ROS) and membrane lipid peroxidation

4.2.1. Detection of ROS in the intracellular space

Intracellular ROS production in control and co-cultivated cells was measured using the fluorogenic dichlorodihydrofluorescein diacetate (H_2DCFDA) probe (Invitrogen, Germany). In this method, the de-acetylation of H_2DCFDA by cellular esterases results in the production of an oxidant-sensitive compound dichlorodihydrofluorescein (H_2DCF), which can be detected spectrofluorimetrically.

To measure the intracellular ROS, 1.0 ml HP cell suspension culture from the treatment and control were transferred to Eppendorf tubes containing 10 μl of 200 μM H_2DCFDA at different time intervals (0, 15, 30, 60, 120, 180 and 240 min) of A. tumefaciens inoculation. The mixture was vortexed 20 s and incubated in dark for 15 min. Then, the cells were spun down in a table-top centrifuge at 6000g for 30 s and 0.5 ml supernatant was taken in a plastic cuvette containing 2.5 ml sterile distilled H_2O. The fluorescence was read in a spectrofluorimeter (Perkin Elmer LS50 Luminescence spectrometer, Buckinghamshire, UK) with 488 nm excitation and 525 nm emission for 1 s.

4.2.2. Detection of ROS in the extracellular space

Hydrogen peroxide (H_2O_2) concentration in the medium was measured by Ferric-xylene orange (FOX1) method (Wolff, 1994). This method is based on the peroxide-mediated oxidation of Fe^{3+}, followed by the reaction of Fe^{3+} with xylene orange. This method is extremely sensitive and used to measure low levels of H_2O-Soluble hydperoxide present in the aqueous phase.

From the control and treated cultures, 1.0 ml suspension was taken in an Eppendorf tube and centrifuged in order to pellet the cells down. Then, 500 μl of the supernatant was transferred to Eppendorf tubes containing 500 μl of xylanol orange reagent (500 μM NH_4Fe(SO_4)_2, 50 mM H_2SO_4, 200 μM xylanol orange, and 200 mM sorbitol). The mixture was vortexed briefly and incubated in dark for 45 min. Absorbance of the Fe^{3+}-xylene orange complex was measured in a spectrophotometer at 560 nm and the concentrations of H_2O_2 in the medium was calculated by plotting those values on H_2O_2 standard graph.

4.2.3. Detection of ROS-induced membrane lipid peroxidation in HP cells

HP cells were harvested by vacuum filtration after 24, 48, 72 and 120 h of A. tumefaciens inoculation. Membrane lipid peroxidation was determined by measuring the amount of MDA as described by Unyayar et al. (2006). Briefly, 200 mg fr. wt of cells were homogenized in 1 ml of 5% trichloroacetic acid (TCA) solution. The homogenates were centrifuged at 12000 rpm for 15 min at room temperature. A 0.5 ml aliquot was mixed with 0.5 ml freshly prepared TBA reagent (0.5% thiobarbituric acid (TBA) in 20% TCA) in a screw cap tube and heated at 96 °C in a boiling water bath. After 25 min, the tubes were snap-cooled in ice to stop the reaction and centrifuged in a table-top centrifuge at 10000 rpm for 5 min. Absorbance of the supernatant was recorded at 532 nm and corrected for non-specific turbidity by subtracting the absorbance at 600 nm. Reaction mixture containing 0.5 ml 5% TCA and 0.5 ml of TBA reagent served as blank. MDA content was determined using the absorption coefficient of 1.56 × 10^5 cm^-2 mol^-1.

4.3. Extraction and analysis of phenolics from HP cells

Equal quantity of HP biomass of control and elicited cells were extracted under dark in 90% MeOH. Briefly, 4 g dry wt was taken in a Falcon tube that contained 40 ml 90% MeOH and sonicated 1 h and kept under dark for 24 h with frequent mixing. Cell debris was spun down by centrifugation at 3000 g for 15 min in a table-top centrifuge. The supernatant was evaporated under vacuum, and dissolved in 2 ml of DMSO to obtain 2 mg dry wt/μl final concentration. Phenolic compounds were analyzed by HPLC–DAD as described previously (Dias et al., 1999). Chromatographic separation was carried out on an RP C18 column (Merck, Germany), using H_2O/HCO_2H (99:1) and MeOH as the mobile phases and absorbance was recorded at 254, 350 and 590 nm.

The quantification of the phenolic constituents was done by the external standard method, using a solution containing chlorogenic acid, p-coumaric acid, quercetin, quercetrin, hypericin and mangiferin as reference compounds in MeOH. Flavonoids were quantified as quercetin equivalents at 350 nm and xanthones as mangiferin equivalents at 254 nm.

4.4. RNA isolation and blot hybridization

About 500 mg fr. wt HP cells were collected after 0, 4, 12 and 24 h A. tumefaciens addition to the cultures and frozen in liquid nitrogen. Total RNA was isolated according to Çakir et al. (2003), denatured, separated on agarose-formaldehyde gel and transferred to nylon membrane (Hybond–N°, Amersham Biosciences, Buckingham, UK) by capillary transfer. DNA fragments specific to phenylalanine ammonia lyase (PAL), 4-coumarate:CoA ligase (4CL) and benzophenone synthase (BPS) cDNAs (from H. perforatum) were labeled with [α-^32P] dCTP (Amersham Biosciences) using Prime-a-Gene labeling kit (Promega, Madison, WI, USA) and used as hybridization probe in 2X SSC at 65 °C. Blots were washed twice with 2X SSC + 0.1% SDS each for 15 min followed by 0.1X SSC + 0.1% SDS for 5 min at 65 °C. The blots were exposed to the imaging screen for 12 h and scanned in a Personal Molecular Imager (Bio–Rad, Hercules, CA, USA).

4.5. Free radical-scavenger activity (DPPH assay)

The antiradical activity of methanolic extracts from HP cells and isolated xanthones (1,3,6,7-tetrahydroxy-8-prenyl xanthone, 1,3,6,7-tetrahydroxy-2-prenyl xanthone, 1,3,7-trihydroxy-6-methoxy-8-prenyl xanthone and paxanthone) were determined by the 1,1-diphenyl-2-picrylhydrazyl (DPPH) assay, as described previously (Silva et al., 2005). The antiradical activity of each sample was evaluated using a series of dilutions, in order to obtain a large spectrum of sample concentrations. The reaction solution consisted of 0.1 ml diluted sample and 1.4 ml of 80 μM DPPH (dissolved in 100% EtOH). The absorbance was monitored continuously at 515 nm with a Perkin–Elmer UV/VIS Spectrometer Lambda2, assuring that the reaction was complete (plateau state). The percentage of reduced DPPH at steady state (DPPH–R) was calculated and these values were plotted against the concentrations methanolic extracts or of pure compounds. A decrease by 50% of the initial DPPH concentration was defined as the half maximal effective concentration (EC_{50}).
4.6. Inhibition of synaptosomal lipid peroxidation (TBARS assay)

Synaptosomal fraction of male Wistar rat brains was isolated and thiobarbituric acid reactive substance (TBARS) assay was performed as described previously (Silva et al., 2005). Briefly, lipid peroxidation was induced in synaptosomes using an oxidant pair (800 μM ascorbic acid and 2.5 μM FeSO₄) in Krebs buffer (132 mM NaCl, 4 mM KCl, 1.4 mM MgCl₂, 1 mM CaCl₂, 6 mM glucose, 10 mM Hepes–Na, pH 7.4) at 37 °C for 15 min. Synaptosomes, without the oxidant pair served as negative control to calculate the basal lipid peroxidation level. The effect of methanolic extracts and isolated xanthones (1,3,6,7-tetrahydroxy-8-prenyl xanthone, 1,3,6,7-tetrahydroxy-2-prenyl xanthone, 1,3,7-trihydroxy-6-methoxy-8-prenyl xanthone and paxathone) on lipid peroxidation was recorded by adding them to the reaction mixture at different concentration. The malondialdehyde (MDA)–thiobarbituric acid (TBA) complex was recorded as a change in absorbance at 530 nm, after reaction of samples with trichloroacetic acid (40%, w/v) and TBA (0.67%, w/v) (96°C, 10 min) and centrifugation (3000g, 10 min). The amount of MDA formed was calculated using the absorption coefficient of 1.56 × 10³ cm mol⁻¹. Values were corrected for the levels of basal peroxidation, obtained during the assay. A decrease by 50% of the value of lipid peroxidation, induced by ascorbate–Fe²⁺, was defined as the EC₅₀.

4.7. Determination of antibacterial activity (colony forming units and disc diffusion)

The stock A. tumefaciens suspension obtained as described before (Franklin et al., 2008) was diluted in MS medium to approximately 250 × 10⁶ CFU/ml. Aliquots (100 μl) from this bacterial suspension were then added to 900 μl of sterile MS medium containing methanolic extracts of control and elicited HP cells, to achieve a range of final concentrations (0, 5, 10, 25, 50 and 100 μl/ml). Similarly, paxathone (5 mg/ml dissolved in DMSO) was incorporated into bacterial suspension at different concentrations (0, 10, 25, 50 and 100 μg/ml). Bacterial suspensions that received the corresponding concentration of DMSO (0, 2.5, 20 and 40 μl/ml) served as control for paxathone treatment. All the samples were maintained on a shaker at 200 rpm at 28°C for 2 days. Aliquots (100 μl) from each sample was serially diluted to 10⁻⁶ and spread on Petri dishes containing 20 ml Luria–Bertani (LB) medium augmented with 25 mg/l rifampicin and 50 mg/l kanamycin. The colony forming units on each plate were counted after incubation for 2 days at 28°C and the percentage of viability was calculated.

Qualitative antimicrobial activity of isolated xanthones was determined by a classical disc diffusion assay. All the isolated xanthones (1,3,6,7-tetrahydroxy-8-prenyl xanthone, 1,3,6,7-tetrahydroxy-2-prenyl xanthone, 1,3,7-trihydroxy-6-methoxy-8-prenyl xanthone and paxathone) were dissolved separately in DMSO (0.1%) and 20 μl of each xanthone solutions were spotted on Petri dishes containing 20 ml Luria–Bertani (LB) medium augmented with 25 mg/l rifampicin and 50 mg/l kanamycin. The colony forming units on each plate were counted after incubation for 2 days at 28°C.

4.8. Statistical analyses

For all the treatments, at least three independent experiments each with three replica were done. Flavonoids and xanthone content of control and elicited HP cells were compared statistically by two way ANOVA followed by Bonferroni post-hoc test. In the same manner, the levels of membrane lipid peroxidation in HP cells after different time periods of A. tumefaciens inoculation were compared. Statistical analyses were performed using GraphPad Prism version, 5.00 for Windows (GraphPad Software, San Diego, CA, USA).

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