In vitro metacestodicidal activity of genistein and other isoflavones against

_Echinococcus multilocularis_ and _Echinococcus granulosus_

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Echinococcus multilocularis and E. granulosus metacestode infections in humans cause alveolar echinococcosis and cystic echinococcosis, respectively, where metacestode development in visceral organs often results in particular organ failure. Further, cystic hydatidosis in farm animals causes severe economic losses. Although benzimidazole derivatives such as mebendazole and albendazole are being used as therapeutic agents, there is often no complete recovery after treatment. Hence, by searching for novel treatment options, we examined the in vitro efficacy of a number of isoflavones against Echinococcus metacestodes and protoscoleces. The most prominent isoflavone, genistein, exhibits significant anti-metacestodicidal activity in vitro. However, genistein binds to the estrogen receptor and can thus induce estrogenic effects, which is a major concern during long-term chemotherapy. We have therefore investigated the activities of a number of synthetic genistein-derivatives carrying a modified estrogen receptor binding site. One of these, Rm 6423, induced dramatic breakdown of the structural integrity of the metacestode germinal layer of both species within 5-7 days of in vitro treatment. Further, examination of culture medium revealed increased leakage of parasite proteins into the medium during treatment, but zymography demonstrated a loss in the activity of metalloproteases. Moreover, two of the genistein-derivatives, Rm6423 and Rm6426 induced considerable damage in E. granulosus protoscoleces, rendering them non-viable. These findings demonstrate that synthetic isoflavones exhibit distinct in vitro effects on Echinococcus metacestodes and protoscoleces, which could potentially be exploited further for the development of novel chemotherapeutical tools against larval stage Echinococcus infection.
INTRODUCTION

Echinococcosis is a cosmopolitan zoonotic disease in ungulates and man that is acquired by infection with the larval stages of the genus *Echinococcus*. Cystic echinococcosis (CE), the causative agent of which is *Echinococcus granulosus*, is distributed worldwide. In contrast, alveolar echinococcosis (AE) and *Echinococcus multilocularis*, are generally confined to the Northern hemisphere (8). Dogs (*E. granulosus*) and foxes (*E. multilocularis*) are the major definitive hosts. After ingestion of eggs, each containing a single oncosphere, *E. granulosus* normally develops into a single fluid-filled unilocular metacestode, whereas *E. multilocularis* metacestodes are characterized by exogenous budding, and form multilocular conglomerates that exhibit typical features of tumor-like proliferation. Growth and/or proliferation of metacestodes over a long period of time leads to the development of space-occupying lesions, causes organ malfunction, and will eventually lead to death (15, 26).

The preferred treatment strategy for CE and AE, is radical resection of the parasitic mass (26). However, in inoperable cases, chemotherapy is the only option. Benzimidazole carbamate derivatives such as albendazole and mebendazole are currently the drugs of choice, being rather efficient for treatment of CE, but acting only parasitostatic and not parasitocidal in patients suffering from AE (15, 28). Another more recently exploited option includes amphotericin B for patients that develop hepatic complications with benzimidazoles (29). Spillage of protoscoleces during surgery can be a new source of infection, Thus, praziquantel, as a protoscolicidal agent, can be included during surgical drainage approaches (23, 24, 40). Nevertheless, especially for AE the recurrence rates after interruption of therapy are high (15, 26, 28, 29), and new options for chemothapeutical treatment are needed.

Flavonoids have been attracting considerable attention as valuable therapeutic options against a number of diseases. Besides their important role in the interactions between plants and...
microorganisms, flavonoids and isoflavones exhibit a range of mammalian health-promoting, antifungal, antimicrobial and antioxidant activities (reviewed in 1, 5, 7). Isoflavones represent by far the largest flavonoid subclass, with a 15-carbon (C6-C3-C6) backbone arranged as a 1,2-diphenylpropane skeleton (30). Isoflavones are commonly found in soya, with genistein being the most abundant one. There are very few studies that have examined the role of isoflavones against parasitic helminths (6, 18, 22). More recently, Gargala et al. (10) reported on the proliferation inhibitory efficacies of epidermal growth factor (EGF) receptor-targeted genistein-derivatives against the apicomplexan parasites *Sarcocystis neurona*, *Neospora caninum* and *Cryptosporidium parvum*. In *E. multilocularis*, an EGF-receptor orthologue (EmER) has been identified and characterized on the molecular level (34). EmER is expressed during infection of the intermediate host, and Egfd, a parasite EGF-like protein with significant homologies to mammalian EGF was shown to be highly upregulated in metacestodes upon incubation with host feeder cells (reviewed in 3). Engagement of EmER by EGF-like peptides is likely to play a crucial role in proliferation and differentiation, through activation of the intracellular tyrosine kinase domain, which then initiates downstream signaling pathways, of which best characterized is the mitogen-activated protein (MAP) kinase cascade (3). Recently, an ERK-like MAP kinase from *E. multilocularis* (EmMPK1) was shown to be functionally activated in response to human EGF (35). Thus, due to the obvious presence of an EGF-like signaling pathway in *Echinococcus*, the in vitro efficacies of genistein and of a limited number of synthetic EGF-receptor tyrosine kinase-targeted genistein-derivatives against *Echinococcus* larval stages and protoscoleces were assessed in this study.
MATERIALS AND METHODS

Biochemicals and drugs. If not otherwise stated, all tissue culture media were purchased from Gibco-BRL (Zurich, Switzerland), and biochemical reagents including genistein were from Sigma (St. Louis, Mo.). Nitazoxanide (NTZ) and tizoxanide (TIZ) were obtained from Romark Laboratories, Tampa, Florida, USA. The structures of genistein and of the synthetic isoflavones (genistein-derivatives) Rm6423, Rm6424, Rm6426 and Rm6427 used in this study have been previously described in Dixon and Ferreira (7) and Gargala et al. (10), respectively. The genistein-derivatives were synthesized at the Department of Chemistry, University of Liverpool.

In vitro culture of *E. multilocularis* metacestodes. In vitro cultivation of *E. multilocularis* metacestodes was carried out as described previously (12, 14). Briefly, *Meriones unguiculatus* were infected intraperitoneally with *E. multilocularis* clone KF5 and isolate IM280, respectively. After 1 to 2 months, the animals were euthanized and the parasite tissue was recovered from the peritoneal cavity under aseptic conditions. The tissue pieces were cut into small tissue blocks (0.5 – 1 cm³), which were washed twice in Hanks balanced salt solution (HBSS). Two pieces of tissue were placed in 75 ml of culture medium (RPMI 1640 containing 12 mM HEPES, 2 mM glutamine, 100 U of penicillin/ml, 100 μg of streptomycin/ml, 0.50 μg of amphotericin B/ml) supplemented with 10% fetal calf serum (FCS) and phenol red. Tissue blocks were kept in tightly closed culture flasks (200 ml) placed in an upright position in an incubator at 37°C with 5% CO₂, with medium changes every 2 to 4 days. These metacestodes were used for in vitro drug assays as described below.
In vitro culture of *E. granulosus* protoscoleces. *E. granulosus* hydatid cysts containing protoscoleces were removed under aseptic conditions from infected sheep presented for routine slaughter in abattoirs in Spain or Kasachstan. In vitro culture of *E. granulosus* protoscoleces and metacestodes was carried out as previously described (41). Briefly, the hydatid cysts (2–5 cm in diameter) were cut open and vesicle fluid (containing protoscoleces) was separated from the metacestode tissue and host adventitia. Protoscoleces were allowed to settle in a 50 ml Falcon tube, were washed twice in HBSS and placed into culture medium (DMEM, 2mM glutamine, 100 U of penicillin/ml, 100 μg of streptomycin/ml and 0.50 μg of amphotericin B/ml) supplemented with 10% FCS and phenol red. Protoscoleces were maintained in culture flasks (200 ml) placed in an upright position in an incubator at 37°C, 5% CO₂, with medium changes every 4–8 days. These protoscoleces were used for (i) in vitro drug assays within 5-10 days of culture, (ii) for mouse infection experiments within 14 days of culture, or (iii) for long-term cultivation and in vitro formation of metacestode stage parasites.

Generation of *E. granulosus* metacestodes in mice. Balb/C mice were purchased from Harlan (Horst, Netherlands) at 6 weeks of age and were housed in a temperature-controlled, light-cycle room in animal facilities according to the Swiss federal animal protection guidelines, with food and water ad libitum. Mice were infected by intraperitoneal inoculation of 2000 viable protoscoleces /mouse, harvested from hydatid cysts and kept in culture for not longer than 14 days. After 3 months of infection, cysts were collected from the peritoneal cavity and maintained in vitro in DMEM with 10 % FCS as described for *E. multilocularis* metacestodes. These *E. granulosus* metacestodes were used for in vitro drug assays as described below.
Drug treatment of *E. granulosus* protoscoleces. All drugs were prepared as stock solutions of 10 mg/ml in dimethyl sulfoxide (DMSO). Treatments of protoscoleces were initiated latest within 10 days of in vitro culture. Initial screenings of compounds were performed in 24 well tissue culture plates containing 100 protoscoleces / well in 1 ml culture medium, which were supplemented with the drugs to concentrations of 1, 5 or 10 μg/ml. Control cultures were supplemented with an equal volume of DMSO alone. The viability of protoscoleces was assessed on a daily bases by microscopic observation of movements, flame cell activity and trypan blue exclusion test (41).

Drug treatment of *Echinococcus* metacestodes and recovery of medium supernatants. Free-floating metacestodes with diameters between 1 and 5 mm were harvested from *E. granulosus* and *E. multilocularis* cultures, respectively. The time of vesicle collection was selected in order to obtain actively growing and culture-adapted metacestodes. The metacestodes were pooled, washed three times in serum free medium, and again divided into separate cultures with approximately 50 vesicles in 15 ml of RPMI culture medium without FCS and phenol red. The drugs were added to the cultures, yielding final concentrations between 1 and 10 μg/ml. For each experiment, control cultures were performed with equal amount of DMSO alone. The parasites were incubated at 37°C with 5% CO₂. Every day, 300 μl of culture medium supernatants were collected and centrifuged at 10,000 x g for 30 min at 4°C. The supernatants were recovered and stored at -20°C for subsequent measurements of *E. multilocularis* alkaline phosphatase (EmAP) activity for *E. multilocularis* or assessment of leakage of proteins by SDS-PAGE for both *E. granulosus* and *E. multilocularis*. 
Determination of EmAP (*E. multilocularis* alkaline phophatase) activity. The procedure described by Stettler et al. (36, 37) was used for the quantitative assessment of EmAP in culture supernatant activity. Briefly, 30 μl from each culture supernatant was mixed with 170 μl of alkaline phosphatase substrate buffer (0.5 M ethanolamine, 0.5 mM MgCl₂ [pH 9.8]) containing *p*-nitrophenyl phosphate (1 mg/ml). A total of 200 μl of each sample was transferred into the wells of a 96-well ELISA plate, and the plate contents were incubated for 30 min at 37°C. *A*₄₀₅ values were read on a Dynatech MRXII ELISA reader.

**SDS-PAGE and immunoblotting.** Proteins from culture supernatants were precipitated in methanol-chloroform (42), and fractions corresponding to the same number of metacestodes were separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) under reducing conditions. Subsequently proteins were transferred onto nitrocellulose membrane, and nonspecific binding sites were blocked in 3% bovine serum albumin (BSA) in TBS-Tween (20 mM Tris-HCl, 150 mM NaCl, 0.3% Tween 20, pH 7.6) for 2 h at room temperature. Blots were labeled with anti-*E. granulosus* hydatid fluid antiserum diluted 1:1'000 in TBS-Tween / 0.3% BSA at 4°C overnight. Bound antibodies were visualized using goat anti-rabbit–alkaline phosphatase conjugates (Promega) according to the instructions provided by the manufacturer.

**Zymography.** Drug treated culture supernatants were precipitated with 80% cold acetone and zymography was performed with 0.1% gelatin substrate SDS-PAGE. Gels were washed with 2.5% Triton-X-100 two times each within 30 min to remove the SDS and to renature the proteins. After a brief wash in water, gels were incubated overnight at 37°C in an incubation buffer (50mM Tris, pH 7.6 containing 50mM NaCl and 10mM CaCl₂) with gentle shaking. In some experiments, the metalloprotease inhibitor 1,10 phenanthroline (1 mM) or the serine protease
inhibitor PMSF (1 mM) were added into the incubation buffer. Following overnight incubation at room temperature, gelatinolytic activity was visualized with Coomassie brilliant blue G250 stain as clear bands in blue background.

**Scanning (SEM) and transmission electron microscopy (TEM).** At day 7 of drug treatment, metacestodes and protoscoleces cultured *in vitro* were processed as described by Hemphill and Croft (13) for SEM and TEM. Briefly, metacestodes were gently opened with a scalpel, and metacestodes and protoscoleces were fixed in 2.5% glutaraldehyde in 100 mM sodium cacodylate buffer (pH 7.2) for 2 h at room temperature, followed by postfixation in 2% OsO₄ in 100 mM sodium cacodylate buffer (pH 7.2) for 2 h at room temperature. Then samples were washed in distilled water and treated with 1% uranyl acetate for 30 min. Subsequently, the specimens were extensively washed in distilled water and dehydrated by sequential incubations in increasing concentrations of ethanol.

For SEM analysis, dehydrated specimens were finally immersed in hexamethyl-disilazane and air-dried under a fume hood. They were then sputter-coated with gold, and inspected on a JEOL 840 scanning electron microscope operating at 25 kV.

For TEM, the specimens were fixed and dehydrated as described above and subsequently embedded in Epon 812 resin (13). Polymerization of the resin was carried out at 65°C overnight. Sections were cut on a Reichert and Jung ultramicrotome and were loaded onto 300-mesh copper grids (Plano GmbH, Marburg, Germany). Ultra thin sections of 80 – 100 nm were made for transmission electron microscopy. Staining with uranyl acetate and lead citrate was performed as described previously (13).
RESULTS

Culture of *Echinococcus* metacestodes in the presence of genistein results in profound morphological alterations. We found that the addition of genistein (5 or 10 μg/ml) into cultures of both *E. multilocularis* and *E. granulosus* metacestodes induced profound morphological and ultrastructural alterations within 7 days (Fig. 1). This was evident by SEM (Fig. 1A, B), with genistein-treated parasite tissue clearly showing decreased numbers of viable cells compared to control (DMSO-treated) parasites. Upon addition of 1 μg/ml of genistein, metacestodes remained unaffected (data not shown). TEM confirmed these findings. Untreated parasites (Fig. 1C) exhibited the typical appearance of the metacestode compartments, including the outer, acellular laminated layer, tegument, and the germinal layer. The tegument lies adjacent to the laminated layer, with distinct microtriches protruding well into the laminated layer. The interior parasite tissue or germinal layer is composed of muscle cells, glycogen storage cells, connective tissue, and undifferentiated cells with a large nucleus. In metacestodes treated with genistein (10 μg/ml) for 7 days, the microtriches were largely shortened or absent, some nuclei exhibited a high degree of chromatin condensation, the cytoplasm in many cells was largely vacuolized, large numbers of lipid droplets were visible, mitochondria appeared electron dense and rounded, and in some areas the laminated layer had separated from the tegumental tissue (Fig. 1 D, E). In addition, the matrix of the laminated layer contained an increased number of small, vesiculated structures of 50-100 nm in diameter (Fig. 1F). Identical alterations in *E. granulosus* metacestodes were noted after treatment with 5 μg/ml genistein, but genistein-treatment at 1μg/ml did not produce any notable alterations in metacestode ultrastructure as seen by TEM. Identical observations were made when *E. multilocularis* metacestodes were treated with genistein (data not shown). Taken together,
genistein-treatment, applied at 5 and 10 μg/ml, resulted in significant morphological and structural alterations in *Echinococcus* metacestodes that marked considerable metabolic stress.

**In vitro efficacy of synthetic EGF-receptor tyrosine kinase-targeted isoflavones (genistein-derivatives) against *E. granulosus* and *E. multilocularis* metacestodes.** For *E. multilocularis* metacestodes, EmAP activity has been earlier used as an indicator assay to demonstrate the loss of viability of drug treated vesicles (36, 37). Thus, we investigated whether in vitro maintenance of *E. multilocularis* metacestodes in the presence of synthetic isoflavones (at 10 μg/ml) had an adverse effect on parasite viability. As positive control the metabolic derivative TIZ of the nitrothiazole analogue NTZ (10 μg/ml) was used. A dramatic increase in EmAP activity in the culture supernatants within 8 days was observed only with Rm6423 and TIZ, while the other isoflavones did not show any effect (Fig. 2A). These experiments were repeated four times and provided identical results. Rm6423 treatment was subsequently done at 1, 5, 2.5 and 10 μg/ml for a period of 14 days (Fig. 2B), demonstrating that the effect of Rm6423 was dose-dependent: 5 and 10 μg/ml lead to strongly increased alkaline phosphatase activity in the medium supernatant, reaching a plateau at around day 7, while at 2.5 μg/ml EmAP levels were increasing continuously until days 14. Addition of 1 μg/ml of genistein was least effective (Fig. 2B).

The alkaline phosphatase activity in culture supernatants of *E. granulosus* metacestodes treated identically was also assessed, but no rise in activity could be seen in any of the culture supernatant samples (data not shown). One possible explanation for the lack of activity could be that the enzyme is trapped within the laminated layer of *E. granulosus* metacestodes, which is much thicker compared to *E. multilocularis*. However, *E. granulosus* metacestodes exhibited loss of turgor and profound morphological alterations already after 3-4 days when treated with TIZ and Rm6423 (data not shown). Thus, *E. granulosus* culture supernatants were separated by SDS-
PAGE, and following immunoblotting, a polyclonal rabbit anti-hydatid fluid antiserum revealed that the amounts of hydatid fluid proteins progressively increased with time in Rm6423-treated-culture supernatants in comparison to cysts treated with the corresponding amounts of DMSO (Fig. 2C). Similar effects could be noted in TIZ-treated cultures, but no appearance of vesicle fluid components could be observed in *E. granulosus* cysts treated with the other isoflavones (data not shown). Similar leakage of vesicle fluid content was seen with *E. multilocularis* culture supernatants collected during drug treatments and probed with a polyclonal anti-*E. multilocularis* metacestode hyperimmune serum (data not shown). Thus, taken together, treatment of *Echinococcus* metacestodes with the isoflavone (genistein-derivative) Rm6423 induces leakage of vesicle fluid content into the medium.

**Rm6423 and TIZ differentially affect the expression of secreted metacestode metalloproteases.** Gelatin-zymography was performed with the culture supernatants from Rm6423-, TIZ-, and DMSO-treated metacestodes. Despite a continuous increase of protein concentration during drug treatment (see Fig. 2C), we noted a clear loss of protease activity in the supernatants of Rm6423 treated cultures compared to those from the DMSO-control and TIZ-treated cultures (Fig. 3A). In contrast, direct addition of Rm6423 during processing of the zymography gels of separated control medium supernatants did not have any impact on protease activities (data not shown). This indicated that the observed effects were due to impaired expression of proteases, associated with the culture in the presence of Rm6423, rather than due to a direct functional inhibition of protease activity by Rm6423. Incubation of zymography gels in the presence of 1 mM 1,10-phenanthroline, a metalloprotease inhibitor, resulted in complete inhibition of all protease activities, (Fig. 3B), while 1mM of the serine protease inhibitor PMSF, did not interfere in the activities (Fig. 3C).
In vitro treatment of metacestodes with Rm6423 induces profound morphological and ultrastructural alterations. SEM of Rm6423-treated *E. multilocularis* metacestodes was performed at day 7 of treatment. SEM showed that, in comparison to control metacestodes, Rm6423 exhibited a devastating impact, with a major portion of the germinal layer being largely distorted by the drug, and only tissue residues were present (Fig. 4A, B). TEM confirmed these findings, and demonstrated, that Rm6423 exhibited effects that were very different from, and actually much more dramatic compared to, genistein (Fig. 4D, E). Although microtriches were still discernable in many areas and still protruding well into the laminated layer, the matrix of the laminated layer exhibited a completely different, more condensed and electron dense, texture compared to the controls (Fig. 4C). All that remained of the tegument and the germinal layer was cellular debris, membrane fragments, nuclear residues and electron dense bodies, thus largely non-viable tissue. Essentially identical results were obtained during Rm6423-treatment of *E. granulosus* metacestodes (data not shown).

Isoflavones affect the viability of *E. granulosus* protoscoleces. The in vitro effects of the same isoflavones on freshly isolated protoscoleces were assessed in comparison to NTZ and TIZ. Only Rm6423 and Rm6426 exhibited profound activities on protoscoleces at 10 μg/ml, in that they reduced the number of viable protoscoleces by 60% within a period of 4 days (Fig. 6A). Fig. 6C demonstrates the degree of morphological alterations and desintegration of the protoscoleces upon treatment with the drugs at 10 μg/ml. At 5 μg/ml, Rm6426 had a clearly decreased efficacy, with 65% of protoscoleces still viable, compared to Rm6423 (45% viable protoscoleces) after 4 days of treatment (Fig. 6B). However, Rm6423 and Rm6426 were clearly less efficient than NTZ and TIZ. Especially treatments with TIZ at both 5 and 10 μg/ml lead to death of all protoscoleces latest at day 7 of culture without the addition of fresh drug during the incubation period.
The number of viable protoscoleces in cultures treated with Rm6423 and Rm6426 at 5 and 10 μg/ml was not further reduced after prolonged treatment with the same drug-containing medium (up to 7 days). In turn, when fresh drug was added at day 7, all parasites were eliminated shortly thereafter (data not shown). At 1 μg/ml, both Rm6423 and Rm6426 exhibited no notable in vitro protoscolicidal activity (data not shown).

**DISCUSSION**

Genistein (4’,5,7-trihydroxyisoflavone) is the most abundant isoflavone, and implicated in prevention of numerous types of cancer and cardiovascular disease. Genistein and a number of other isoflavones have been shown to mediate anti-plasmodial and anti-coccidial activities (5, 10, 20). The compound exhibits micromolar potency in inhibiting protein tyrosine kinases, which probably accounts for most of its effects (2, 39).

Our results show that in vitro treatment with genistein induces a number of significant alterations on *E. multilocularis* and *E. granulosus* metacestodes that could eventually impair parasite viability and lead to parasite death (see Fig. 1). Genistein, like other isoflavones, exhibits the basic structure of estrogen, and thus can exert estrogenic effects through binding to the estrogen receptor. This represents a serious health concern in terms of the use of isoflavones as therapeutic agent, especially for long-term treatments as for echinococcosis (4, 31, 32, 38). Crystal structure analysis of ligand-receptor complexes involving estrogen receptor-β and genistein showed that the phenolic C-ring interacts with estrogen receptor-β (27). Thus, it was demonstrated that the C-ring, and more precisely the respective 4’-OH-group, is responsible for the estrogenic effects.

In this study, we used synthetic isoflavones, in which the crucial 4’ OH-group on the C-ring was removed, and other functional steric groups were added at different positions. Of the drugs tested (see reference 10), only Rm6423 exhibited a profound anti-parasitic effect towards metacestodes
of both *E. multilocularis* and *E. granulosus*, leading to leakage of parasite proteins into the medium supernatants (see Fig. 2). Rm6423 is almost identical to genistein, but lacks the OH-group on the 4’-position of the C-ring, and a bromo-group is added at position 2’. Interestingly, if the bromo-group is added to position 3’ of the C-ring, as in Rm6424 (10), the efficacy of the compound is completely lost. As evidenced by SEM and TEM, the damage induced by Rm6423 (see Fig. 4) is comparable to what had been previously observed for NTZ and TIZ, with no retraction of microtriches but efficient and complete desintegration and necrosis of germinal layer (37), but clearly different from the changes identified in genistein-treated metacestodes (see Fig. 1). However, a similar enrichment of small vesicles within the matrix of the laminated layer has also been previously observed in NTZ-treated *E. multilocularis* metacestodes (37), and retraction of microtriches had been previously observed in *E. multilocularis* metacestodes treated with albendazolesulphoxide and albendazolesulphone, respectively (17).

All these features probably reflect different mechanisms of action of these drugs. Benzimidazoles such as albendazole have been shown to bind to tubulin and inhibit its polymerization into microtubules, (21). In contrast, the mode of action of thiazolides such as NTZ and TIZ in helminths has not been elucidated so far. Possibly, the drugs interfere in the functional activity of enzymes that are similar to pyruvate ferredoxin oxidoreductase (PFOR) in anaerobic bacteria, but other mechanisms of action are currently being discussed (19, 16, 25, 33).

Gelatin zymography clearly demonstrated that the activity of some metalloprotease bands was impaired in medium supernatants of Rm6423-treated *E. granulosus* metacestodes, despite of the fact that the overall protein concentration of hydatid fluid components has increased by drug treatment. This suggests that Rm6423 has a negative influence on metalloprotease expression in *Echinococcus* metacestodes. In mammalian cells, metalloprotease expression is regulated through selective activation or inhibition of a number of signaling systems, including the EGF-
receptor-regulated p38 MAP kinase in cancer cells (11, 19). Whether any member of the recently discovered *Echinococcus* EGF signaling pathway and MAP kinase cascade (reviewed in 3) is affected by Rm6423 needs to be investigated in future studies.

Treatment of freshly isolated *E. granulosus* protoscoleces with the small panel of isoflavones used in this study resulted in the identification of Rm6423 and Rm6426 as two compounds with limited protoscolicidal activity. This effect was dose-dependent, and not evident anymore at 1 μg/ml. Both isoflavones were not as efficient as NTZ and TIZ, respectively. In addition, the isoflavones had basically lost their efficacy after 4 days, in contrast to NTZ and TIZ, which continued to exert the anti-parasitic activity until day 7, when all protoscoleces were non-viable (Fig. 5). However, when the medium containing Rm6423 and Rm6426 was replaced with fresh drug containing medium after 7 days, all protoscoleces had lost viability within the next 24 h. It is therefore possible that, in contrast to the thiazolides, during the first 3-4 days isoflavones are metabolized or converted into inactive compounds, and loose antiparasitic efficacy.

Taken together, we have shown that genistein and the genistein derivative Rm6423 exhibit profound activity against *Echinococcus* metacestodes. Rm6423 is an extremely interesting compound, as it lacks a functional estrogen-receptor binding domain, and the expected toxicity of the drug is low. Therefore, animal experimentation will be required to provide the proof of concept that Rm6423 could be useful for in vivo treatment of *Echinococcus* infection.
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Figure legends

Fig. 1. In vitro treatment of *Echinococcus* metacestodes with genistein induces distinct morphological and structural changes. (A, B): SEM. *E. granulosus* metacestodes were exposed to the solvent DMSO (A) or 10 μg/ml genistein (B), and the parasite tissue was visualized by SEM. Note the loss of cellular integrity of the germinal layer in (B). Bars = 280 μm for both A and B. Similar results were obtained for *E. multilocularis* (data not shown). (C-F): TEM of control and genistein-treated metacestodes. (C) control tissue, showing a section through the *E. granulosus* vesicle wall with laminated layer (LL), tegument (Te) with microtriches (Mt), and germinal layer (Gl). (D-F) treatment with genistein (10 μg/ml) for 7 days results in loss of microtriches, partial separation from tegument and laminated layer (arrowhead in (D)), formation of lipid droplets, and increased occurrence of small lipid vesicles (ld) in the laminated layer matrix (arrows in (F)). Bars in C = 2.4μm; D = 2.8μm; E = 1.9μm; F = 0.5μm. Similar results were obtained for *E. multilocularis* (data not shown).

Fig. 2. Assays for the detection of drug-induced metacestode damage. (A) EmAP assay demonstrating the increased release of alkaline phosphatase activity from *E. multilocularis* metacestodes during in vitro treatment with nitazoxanide (NTZ: pos. control), and synthetic isoflavonoids Rm6423, Rm6424, Rm6426, Rm6427. Note the increased efficacy of Rm6423. (B) Dose response EmAP assay with Rm6423, showing a clear relationship between drug concentration and presence of EmAP activity in medium supernatant of *E. multilocularis* metacestode cultures. (C) Measurement of release of hydatid fluid compounds from *E. granulosus* metacestodes following treatment with Rm6423 by immunoblotting of medium supernatants after SDS-PAGE at different timepoints and labeling with a polyclonal antiserum.
directed against *E. granulosus* vesicle fluid. Note the time-dependent increase of signal. As a negative control, vesicles were incubated with equivalent concentration of DMSO.

Fig. 3. Gelatin Zymography reveals differential protease expression pattern in *E. granulosus* metacestode culture supernatants of Rm6423-, TIZ- and DMSO-treated parasites. (A) Both TIZ- and Rm6423-treated fractions exhibit profound differences compared to DMSO-treated fraction. Those protease bands marked by arrowheads are completely absent in Rm6423-treated medium supernatants. (B) Note the complete inhibition of all protease in the presence of phenanthroline, indicating that these are all metalloproteases. (C) No inhibition of protease activity in the presence of PMSF.

Fig. 4. In vitro treatment of *Echinococcus* metacestodes with Rm6423. SEM (A, B) and TEM (C-E) showing the alterations induced by Rm6423 on *E. multilocularis* metacestodes. Similar findings were obtained with *E. granulosus*. (A) and (C) are control treated parasites exhibiting intact parasite tissue. In (B), (D) and (E), the damage induced by Rm6423 is shown. LL = laminated layer, Te = tegument, Gl = germinal layer, arrows point towards microtriches. Bars in A = 280μm; B = 280μm; C = 1.9μm; D = 2 μm; E = 2μm.

Fig. 5. Effects of synthetic isoflavones on *E. granulosus* protoscoleces. Protoscoleces of *E. granulosus* were exposed to Rm6423, Rm6426, NTZ and TIZ for 7 days, and the viability of the parasites was assessed by trypan blue staining and light microscopical inspection. In (A), all drugs are added at 10 μg/ml, in (B) at 5 μg/ml. The percentage of still viable protoscoleces is indicated at different timepoints. In (C), the effects of the different drug treatments (10 μg/ml) on the morphology and structural integrity of protoscoleces are visualized at day 3 of treatment.
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