Transport of paclitaxel (Taxol) across the blood-brain barrier in vitro and in vivo

Stephan Fellner,1 Björn Bauer,2 David S. Miller,3 Martina Schaffrik,1 Martina Fankhänel,1 Thilo Spruß,3 Günther Bernhardt,1 Claudia Graeff,2 Lothar Färber,4 Harald Gscchaimdeier,4 Armin Buschauer,1 and Gert Fricker2

1Institute of Pharmacy, University of Regensburg, Regensburg, Germany
2Institute of Pharmaceutics and Biopharmacy, University of Heidelberg, Heidelberg, Germany
3Laboratory of Pharmacology and Chemistry, National Institute of Environmental Health Sciences, NIH, Research Triangle Park, North Carolina, USA
4Novartis Pharma GmbH, Nürnberg, Germany

Paclitaxel concentrations in the brain are very low after intravenous injection. Since paclitaxel is excluded from some tumors by p-glycoprotein (p-gp), the same mechanism may prevent entry into the brain. In vitro, paclitaxel transport was examined in capillaries from rat brains by confocal microscopy using BODIPY Fl-paclitaxel. Western blots and immunostaining demonstrated apical expression of p-gp in isolated endothelial cells, vessels, and tissue. Secretion of BODIPY Fl-paclitaxel into capillary lumens was specific and energy-dependent. Steady state luminal fluorescence significantly exceeded cellular fluorescence and was reduced by NaCN, paclitaxel, and SDZ PSC-833 (valspodar), a p-gp blocker. Leukotriene C4 (LTC4), an Mrp2-substrate, had no effect. Luminal accumulation of NBDL-cyclosporin, a p-gp substrate, was inhibited by paclitaxel. In vivo, paclitaxel levels in the brain, liver, kidney, and plasma of nude mice were determined after intravenous injection. Co-administration of valspodar led to increased paclitaxel levels in brains compared to monotherapy. Therapeutic relevance was proven for nude mice with implanted intracerebral human U-118 MG glioblastoma. Whereas paclitaxel did not affect tumor volume, co-administration of paclitaxel (intravenous) and PSC833 (peroral) reduced tumor volume by 90%. Thus, p-gp is an important obstacle preventing paclitaxel entry into the brain, and inhibition of this transporter allows the drug to reach sensitive tumors within the CNS.


Introduction

Paclitaxel (Taxol) and derivatives are active against various tumors (1–6) and have also been used to treat malignant glioma and brain metastases (7–9). However, brain tumors constitute a difficult problem and the therapeutic benefit of paclitaxel has been variable and low. This could be attributed to its limited entry into the CNS. Although paclitaxel is very lipophilic, concentrations in the CNS are very low after intravenous administration (10, 11).

Paclitaxel appears to be a substrate of the multidrug resistance protein p-gp (12–14), and it is likely that this transporter contributes to its limited access to the brain. P-gp is expressed in high levels in cultured brain capillary endothelial cells and in intact brain capillaries (15, 16). It is localized at the luminal surface of the endothelium (17, 18), and therefore is in the correct location to restrict permeation of a variety of drugs into the CNS (19–21). Animals with reduced p-gp function show an accumulation of p-gp substrates in the brain as well as a markedly increased sensitivity to neurotoxic p-gp substrates, e.g., ivermectin (18, 22, 23).

The present study identifies in vitro a mechanism that limits paclitaxel access to the CNS and outlines a strategy to circumvent it by blocking p-gp. We demonstrated that the p-gp blocker valspodar enhances paclitaxel entry into the brains of mice after intravenous dosing and that valspodar dramatically increases paclitaxel effectiveness against a human glioblastoma implanted into the CNS of nude mice. These are the first data directly demonstrating the role of p-gp in limiting the therapeutic availability of paclitaxel to the CNS.

Methods

Chemicals. Valspodar (SDZ PSC 833) and the fluorescent cyclosporin derivative NBDL-CS were from Novartis Pharma GmbH (Nürnberg, Germany and Basel, Switzerland). Paclitaxel, BODIPY Fl-paclitaxel,
and calcein-AM were from Molecular Probes Europe BV (Leiden, The Netherlands).

Capillary isolation. Capillaries from male Wistar rat brains were isolated at 4°C in pregassed (95% O₂/5% CO₂) solutions as described (18). Briefly, pieces of cortical gray matter were homogenized in three volumes (vol/wt) of buffer A (103 mM NaCl, 4.7 mM KCl, 2.5 mM CaCl₂, 1.2 mM KH₂PO₄, 1.2 mM MgSO₄, 15 mM HEPES), and, after addition of dextran (final concentration 30%), the homogenate was centrifuged at low speed. The pellet was resuspended in buffer A supplemented with 25 mM NaHCO₃, 10 mM glucose, 1 mM Na-pyruvate, 0.5% (wt/vol) BSA, and filtered through a 200-µm nylon mesh. The filtrate was passed over a glass bead column, and after washing, capillaries adhering to the beads were collected by gentle agitation. Capillaries were centrifuged, the pellet was resuspended in ice-cold, BSA-free Krebs-Henseleit buffer (KHB) and immediately used for transport experiments.

Fluorescence microscopy. Capillaries were transferred to a Teflon incubation chamber, containing 0.5–1.0 ml of Krebs-Ringer medium with fluorescent compounds and transport effectors. Fluorescent compounds and inhibitors were added to the incubation medium in DMSO stock solutions. Earlier experiments showed that DMSO concentrations less than or equal to 0.5% had no significant effect on transport. The chamber floor was a 2 × 2–cm glass coverslip through which the tissue could be observed by a confocal laser scanning microscope (LSM 5 Pascal, 488-nm laser line, 515-nm long-pass emission filter; Carl Zeiss Jena GmbH, Jena, Germany). A low laser intensity (<3% of maximum) was used to avoid photobleaching. Tissue autofluorescence was not detectable. The images were analyzed with the Scion Image software (Beta 3; Scion Corp., Frederick, Maryland, USA). Epifluorescent images of capillaries were acquired by averaging four to eight frames. Earlier, using fluorescein as reference compound, it was shown that fluorescence intensity and fluorescein concentration had an approximately linear relationship (24). Here, the data are given as average of measured pixel intensity rather than estimated concentration of fluorescent compounds. Intensities were measured as described for a similar system of kidney proximal tubules (25). Several cellular and luminal areas were selected, and the average pixel intensity for each area was calculated. Analysis by high performance thin layer chromatography of BODIPY Fl-paclitaxel and NBDL-CS after extraction out of tissue with ethanol gave no hint for degradation or transformation of the fluorescent compounds for at least the time of incubation (≤ 1 hour).

Isolation of endothelial cells and primary culture. Porcine brain capillary endothelial cells were isolated as described (26). Primary cultures were prepared according to ref. 27. Cells were collected and washed twice in medium containing 10% horse serum. The final suspension was kept in medium with 20% horse serum and 10% DMSO and stored in liquid nitrogen until use.

For uptake experiments, cells were seeded onto collagen-coated (Hoffman-LaRoche Ltd., Basel, Switzerland) 96-well plates. They were cultured under standard conditions using isolation medium without gentamycin, but with 10% horse serum. One day before transport experiments medium was replaced with 45% MEM, 45% F12-HAM, 100 µg/ml streptomycin, 100 µg/ml penicillin G, 10 mM HEPES, 2 mM glucose, and 550 mM hydrocortisone. For the calcein assay, cells were washed with KRB buffer (37°C) and incubated with the test compounds for 5–15 minutes. Calcein-AM (1 µM) was added, and cells were incubated 15–45 minutes. Then they were washed three times with ice-cold KRB and lysed with 1% Triton X-100. Fluorescence was measured using a Fluoroskan Ascent reader (LabSystems GmbH, Frankfurt, Germany).

Western blots. P-gp was detected by Western blot analysis using the mAb C219 (Alexis Deutschland GmbH, Grünberg, Germany). The p-gp–overexpressing cell line (MDR+)P388 was used as control. SDS PAGE was performed with a Mini-Protean II apparatus (Bio-Rad Laboratories GmbH, Munich, Germany). Proteins were transferred to nitrocellulose membranes using a Mini Trans-Blot cell for 2.5 hours at 250 mA. The membranes were incubated with Ab (1:20 dilution) in PBS containing 0.05% Tween 20, 1% BSA, and 1% milk powder for 2 hours at 37°C. After washing, p-gp was detected using enhanced chemiluminescence (ECL) reagent (Amersham International, Buckinghamshire, United Kingdom).

Tumor cell lines and culture conditions. Human U-87 MG (HTB 47) and U-118 MG (HTB 15) glioblastoma/astrocytoma cell lines were obtained from American Type Culture Collection (Rockville, Maryland, USA). Cell banking and quality control were performed according to the “seed stock concept.” U-87 MG cells were grown in Eagle’s MEM containing 1-glutamine, 2.2 g/l NaHCO₃, 110 mg/l sodium pyruvate, and 10% FCS. U-118 MG cells were maintained in DMEM supplemented with 10% FCS. Cells were cultured in a carbonated atmosphere at 37°C and were passaged following treatment with trypsin (0.05%)/EDTA (0.02%). For tumor cell implantation into nude mice, cells were harvested mechanically, spun down, and washed twice with serum-free medium.

Chemosensitivity assay. The in vitro antitumor activity of paclitaxel and the modulating effect of valsopdar were determined by a kinetic assay based on crystal violet staining. Processing procedure and data analysis were essentially performed as described (28). Paclitaxel and valsopdar were added as 2,000-fold concentrated feed solutions in 70% ethanol. Doubling-time analysis of the growth curves of untreated controls was accomplished by a polynomial regression fit applying the least-squares method as earlier reported (29).
Immunocytochemistry. Subconfluent tumor cells, cultured on chamber slides, were fixed with 4% phosphate buffered paraformaldehyde, pH 7.4, for 30 minutes and washed three times with PBS. Endogenous peroxidase was blocked by incubation in PBS containing 3% H2O2 and 10% methanol for 30 minutes, followed by two incubations in PBS for 5 minutes. Nonspecific binding was prevented by 30-minute incubation in 10% goat serum (Dianova, Hamburg, Germany), 2.5% skimmed milk powder (Milupa, Friedrichsdorf, Germany), and 0.3% Triton X-100 in PBS. Subsequently, cells were washed three times with PBS containing 0.04% Triton X-100 (PBST).

Primary Ab’s AB-1 (PC03) (polyclonal anti-human/mouse p-gp, mdr; Dianova) and JSB-1/IgG-1 (monoclonal anti-human p-gp; Alexis Deutschland GmbH) were diluted 1:50 with washing buffer consisting of 10 ml PBST, 100 µl NGS, and 0.3% Triton X-100. After incubation with primary Ab overnight at room temperature (RT), the slides were washed three times with PBS containing 0.34% Triton X-100 and treated with secondary Ab (for Ab-1, biotinylated goat anti-rabbit, 1:200; Sigma-Aldrich, Deisenhofen, Germany; for JSB-1, biotinylated donkey anti-mouse 1:800; Dianova) for 2 hours at RT in a humid chamber. Then, cells were washed three times in PBST and incubated for 1 hour in horseradish-conjugated avidin-biotin complex (Vectastain Elite Kit ABC; Camon Labor Service GmbH, Wiesbaden, Germany) in PBST. Slides were washed three times for 10 minutes in PBST before incubation in a diaminobenzidine (DAB) solution for 10 minutes (0.05% DAB, 0.02% ammonium nickel sulfate in PBST). H2O2 was added to a final concentration of 0.006%, and slides were incubated for 10 minutes. Subsequently, they were washed three times in PBS, processed in an ascending alcohol series, and mounted in DePex.

Immunohistochemistry. Isolated brain capillaries were transferred onto slides and fixed for 20 minutes with 3% paraformaldehyde, 0.1% glutardialdehyde, and 3.4% sucrose in PBS. Capillaries were permeabilized for 15 minutes with 1% (vol/vol) Triton X-100 in PBS. Then, the capillaries were incubated for 1 hour at 37°C in a humid chamber with primary Ab (150 µl murine mAbB C219, 1:20 dilution). After washing the corresponding fluorochrome-conjugated secondary Ab (150 µl FITC-conjugated rabbit anti-mouse IgG, 1:20 dilution; Hoffmann-LaRoche Ltd.) was added for 1 hour in a humid chamber in the dark.

Murine tissues and intracerebrally growing human tumors were fixed in Bouin’s solution for 3 days, then were dehydrated and embedded into paraffin. Sections of 5-µm thickness were prepared, deparaffinized with xylene, and rehydrated in a descending alcohol series. P-gp was detected with the polyclonal Ab-1 (PC03) Ab (anti-human and murine p-gp; Dianova) with nuclear fast-red counter staining. The protocol was the same as for immunocytochemistry except that the primary Ab was diluted 1:10 with washing buffer.

Pharmacokinetic studies in nude mice. NMRI nu/nu mice were bred under specified pathogen-free conditions at 26°C, 70% relative humidity, and a 12-hour light/12-hour dark cycle at the University of Regensburg (30). They were fed ad libitum with a breeding/maintenance diet (Altromin GmbH, Lage, Germany) and water containing 1.3 g/l potassium sorbate and 2 g/l chloramphenicol; pH was adjusted to 2.5 with HCl. The use of animals in this study complies with the Guide for the Care and Use of Laboratory Animals (NIH publication no. 86-23, revised 1985) and the current German law on the protection of animals.

Prior to drug administration, 6- to 8-week-old female NMRI (nu/nu) mice were assigned to groups of three to four animals per time point. Drinking solution containing 100 mg/ml valsodar was diluted to a concentration of 5 mg/ml with its vehicle, a mixture of Cremophor RH40, ethanol, d,l-α-tocopherol, propylene glycol, and Labrafil M2125CS. A volume of 0.2 ml was administered by gavage per 20 g of body weight, resulting in a dosage of 50 mg valsodar per kilogram of body weight. In the same way, the solvent was applied to the control animals.

For intravenous injection the commercial drug formulation Taxol (6 mg/ml paclitaxel; Bristol-Myers Squibb) was diluted with 0.9% NaCl to a paclitaxel concentration of 1.6 or 0.8 mg/ml. The drug was injected via the retrobulbar plexus in a volume of 0.1 ml per 20 g of body weight at a dosage of 8 or 4 mg paclitaxel per kg body weight within approximately 10 seconds. Four hours before the injection of paclitaxel, either a dose of 50 mg/kg valsodar (valsodar group) or a corresponding volume of the vehicle of valsodar (control) was administered to the mice by mouth.

To minimize adulterations of the analytical results by drug in the vasculature, ketamine/xylazine–anesthetized mice were killed by cardiac puncture, 1.5, 3, and 24 hours after intravenous injection, and heparin-plasma, brain, liver, and kidneys were collected. Tissues were homogenized in 4% BSA within 15 minutes and shock-frozen in liquid nitrogen. For homogenization, each organ was minced into small pieces. One milliliter 4% BSA solution was added per 150 mg tissue, and homogenization was carried out with a Potter-Elvehjem homogenizer by 15 strokes. All samples were stored at −78°C.

Paclitaxel analysis. Tissue extraction was by a modification of the method of Sparreboom et al. (31). Thawed tissue homogenate (1-2 ml) was pipetted into a glass centrifugation tube with a ground joint. Samples less than 1 ml were filled up to 1 ml with 4% BSA solution. Tissue samples underwent diethyl ether extraction and solid-phase extraction (SPE). Plasma samples consisted of 250 µl of thawed murine plasma. Samples with less than 250 µl volume were filled up to 250 µl with human citrate-phosphate-dextrose/plasma. Only plasma samples underwent SPE.

To each sample 10 µl of 50 µM docetaxel solution was added as internal standard. For docetaxel stock...
solutions Taxotere (80 mg docetaxel per 2 ml of polysorbate 80; Rhone-Poulenc Rorer, Antony Cedex, France) was diluted with absolute ethanol. HPLC analysis was performed on a Thermo Separation Products liquid chromatograph (Thermo Quest; Egelsbach, Germany). A stainless steel (125 x 4 mm) analytical column equipped with a guard column (4 x 4 mm), both packed with 5 μm LiChroSpher 100 RP-18 material (Merck KGaA, Darmstadt, Germany), was cooled to 33°C. The mobile phase consisted of acetonitrile/methanol/0.2 M ammonium acetate buffer, pH 5.0, 38:10.5:51.5 (vol/vol). Ultraviolet detection was performed at 227 nm, and a washing gradient was applied after each run (additional 35% vol/vol acetonitrile). Injection volumes were either 50 or 100 μl. For calibration, paclitaxel and docetaxel stock solutions in anhydrous ethanol, stored at –20°C, were diluted with mobile phase (1:100) on the day of analysis.

The method of analysis was validated for murine plasma, brain, liver, and kidney samples (32). Correlation coefficients for paclitaxel calibration curves in the concentration range 50–1,000 nM were greater than 0.999. Recovery rates for paclitaxel ranged from 50 to 85%, depending on paclitaxel concentration and the type of tissue. The method’s accuracy for tissue samples was below 2% for paclitaxel concentrations between 0.667 and 6.67 nmol/g of tissue and better than 8% for 0.133 nmol/g. The precision for these determinations was better than 10%.

Tumor implantation, treatment, and evaluation. In the intracerebral model, the parietal bone of anaesthetized male nude mice (mean body weight 35 g) was drilled through with a 1-mm-diameter bit 3 mm on the right-hand side of the sagittal line and 3 mm rostral of the coronal line. Human U-87 MG cells (10⁶; glioblastoma/astrocytoma grade III) or 300,000 U-118 MG (glioblastoma) tumor cells suspended in 10 μl of serum-free Eagle’s minimum essential medium were injected 3 mm deep into the brain tissue, and the wound was closed with a surgical clamp.

Mice were assigned to three groups (n = 7). The control group received no medication. Two additional groups were treated as described for the pharmacokinetic experiment either with a combination of valspodar and paclitaxel or the vehicle of valspodar and paclitaxel. Valspodar (50 mg/kg) and vehicle were given to the animals inoculated with the U-87 MG tumors by mouth 4 hours before the intravenous injection of two doses of paclitaxel, i.e., 4 mg/kg on day 7 and another 2 mg/kg on day 14 after tumor implantation. Animals were killed at day 20, and brains were fixed in Bouin’s solution for histological examination. Animals bearing intracerebral U-118 MG tumors were treated analogously, except drugs were administered on days 8 and 15, the first paclitaxel dose was reduced to 3 mg/kg, and brains were removed for morphometric analysis on day 35.

Brains were fixed in Bouin’s solution for 3 days, dehydrated, and embedded into paraffin. Serial 5-μm coronal sections were prepared every 200 μm (U-87 MG) or 300 μm (U-118 MG) starting and ending 1.5 mm rostral and caudal to the region of tumor cell implantation, deparaffinated, and stained according to Masson and Goldner (33). Tumor areas on nine central sections were determined per brain morphometrically with the software Metamorph (Visitron Systems GmbH, Puchheim, Germany), and the tumor volume in the investigated brain region was calculated.

In the subcutaneous model, approximately 5 × 10⁶ U-118 MG cells (100 μl) were injected into the right flank of 6-week-old female NMRI (nu/nu) mice. Solid tumors developed within 8 weeks. For serial passage, the tumors were cut into 2-mm³ pieces and transplanted subcutaneously into the thoracic region with a trocar (30).

Mice were assigned to four groups of ten animals. The control group received no medication, another group received the complex vehicle of valspodar, whereas valspodar was administered to the third and fourth groups at a dosage of 10 and 50 mg/kg, respectively. Valspodar and its vehicle were administered orally twice per week, resulting in cumulative doses of 120 and 600 mg valspodar per kilogram. On day 46 after tumor transplantation the animals were killed by cervical dislocation.

Tumor growth kinetics was recorded by measuring tumor diameters with an electronic caliper. Tumor areas were calculated as the product of two perpendicular diameters, of which one represents the largest possible diameter of the tumor. Tumor size and the weight of the animals were measured twice a week.

Statistical analysis. Data are presented as mean values plus or minus SEM and were evaluated by one-way ANOVA using SPSS 10.0 for Windows. Except for the pharmacokinetic results, which were analysed non-parametrically by the U test according to Mann and Whitney, the unpaired Student t test for inhomogeneous variances was applied. P values less than or equal to 0.05 were considered statistically significant.

Results

Isolated brain capillaries. Integrity of isolated capillaries was checked by microscopy. After a 60- to 80-minute incubation with FITC-dextran 4000 as a marker for paracellular permeation, no fluorescence was detected inside the capillary lumina by confocal scanning microscopy, indicating that tight junction integrity was maintained and that the endothelium of isolated capillaries is a significant barrier to diffusion of large molecules, which is consistent with in vivo observations.

Immunohistochemical assessment of p-gp in intact rat capillaries (Figure 1a) showed the primary localization of the export protein in the luminal membrane, with only weak staining at the outer membrane. Presence of p-gp was confirmed by Western blots of porcine brain capillary endothelial cells that were cultured for 7 days prior to transport studies (Figure 1b). In the human brain, p-gp (MDR1) was found in capillaries in the cortex, the hippocampus, the cerebellum, and the spinal cord, but not in submeningeal blood.
vessels and the vessels of the choroid plexus (34). In coronal sections of the brains of NMRI nude mice, p-gp (AB-1 Ab’s) was not only detected in capillaries of the cortex but also in deeper medullar regions. Figure 1c shows that not only small capillaries and larger blood vessels in the cortex stained positive, but also that p-gp is expressed in submeningeal arteries and veins. The immunopositive small capillaries were unevenly distributed in the cerebral parenchyma, reflected in a somewhat lower density between the outer band of Baillarger and Kaes-Bechterew’s band (results not shown). Hepatic bile canaliculi, renal proximal tubules, and small intestinal brush borders also stained positive (not shown).

Figure 1
(a) Immunostaining of p-gp in a freshly isolated rat brain capillary (primary Ab: MAB C219; secondary Ab: FITC-conjugated rabbit antibody to mouse IgG). (b) P-gp detection by Western blot using the MAB C219. Left lane: brain capillaries; right lane: p-gp overexpressing cell line (MDR+)P388 as positive control. (c) Immunostaining of coronal sections of the brains of nude mice, using the polyclonal AB-1 Ab. Healthy animal, with P-gp expression in capillaries and larger blood vessels in the cerebrum (bottom) and the submeningeal arteries and veins (top). (d) Intracerebral human glioblastoma (U-87 MG) with immunopositive capillary endothelia and negative tumor cells. The arrow points to the ependyma.

Figure 2a shows the time course of BODIPY Fl-paclitaxel accumulation in intact rat brain capillaries. Fluorescence in cells and capillary lumina rapidly increased, reaching steady-state values after 30–45 minutes. At steady state, luminal fluorescence was about three times cellular fluorescence. Medium fluorescence was barely detectable, so BODIPY Fl-paclitaxel clearly accumulated in the cells and lumina. In capillaries exposed to 1 mM NaCN, steady-state luminal fluorescence was significantly reduced compared with controls, but cellular fluorescence was unchanged (Figure 2a). With NaCN, luminal fluorescence approached cellular levels. Thus, accumulation of BODIPY Fl-paclitaxel within the cells was not energy dependent, probably reflecting passive diffusion, binding to cellular elements, and partitioning into membranes. In contrast, movement of BODIPY Fl-paclitaxel from cell to lumen was energy dependent, suggesting uphill transport. Luminal accumulation of BODIPY Fl-paclitaxel was reduced in a concentration-dependent manner by unlabeled paclitaxel, suggesting that both compounds interact with the same transporter. Luminal accumulation of BODIPY Fl-paclitaxel was also reduced by the p-gp inhibitor, valspodar (Figure 2b). Neither paclitaxel nor valspodar reduced cellular fluorescence. In contrast, the Mrp

**Figure 2**
(a) Time-dependent accumulation of BODIPY Fl-paclitaxel in cells and lumens of isolated rat brain capillaries in the absence and presence of 1 mM NaCN. Means ± SEM, n = 10–12, P < 0.05. (b) Accumulation of BODIPY Fl-paclitaxel and NBDL-CS (c), respectively, in the lumens of isolated capillaries in the absence and presence of unlabeled paclitaxel, the p-gp blocker valspodar, and the mrp2 substrate LTC4. The fluorescent compounds were added at concentrations of 1 µM. Means ± SEM, n = 10–12. *Significantly different from controls; P < 0.05.
Table 1
Effect of valspodar coadministration on paclitaxel levels (mean values ± SEM) in liver and kidney of nude mice (n = 3) after intravenous injection of 8 mg/kg paclitaxel

<table>
<thead>
<tr>
<th>Time (h)</th>
<th>Liver</th>
<th>Mean paclitaxel level ± SEM (nmol/g)</th>
<th>Kidney</th>
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<tr>
<td></td>
<td>Paclitaxel</td>
<td>Valspodar plus paclitaxel</td>
<td>Paclitaxel</td>
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<tr>
<td>1.5</td>
<td>33.49 ± 1.87</td>
<td>18.50 ± 1.18&lt;sup&gt;a&lt;/sup&gt;</td>
<td>12.25 ± 0.63</td>
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<tr>
<td>3</td>
<td>18.03 ± 2.98</td>
<td>20.17 ± 0.74</td>
<td>7.87 ± 0.65</td>
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<td>24</td>
<td>0.91 ± 0.24</td>
<td>2.28 ± 0.34&lt;sup&gt;a&lt;/sup&gt;</td>
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Four hours before intravenous application the animals were treated with the vehicle of valspodar by mouth (paclitaxel) or 50 mg/kg valspodar by mouth (valspodar plus paclitaxel). <sup>a</sup>Statistically significant differences (P ≤ 0.05).
KB-V1 cells stained intensely with both Ab’s, U-87 MG and U-118 MG glioma cells did not.

When inoculated intracerebrally, the human glioblastoma cell lines U-87 MG and U-118 MG formed solid tumors as shown in the Figure 1d. Immunostaining with the AB-1 Ab revealed p-gp expression in the neovascularure, which is consistent with the clinical situation (43–45). On the contrary, in agreement with the results of immunocytochemistry the tumor cells did not exhibit the MDR1 phenotype and did not stain with the monoclonal anti-human p-gp 170 JSB-1 Ab (not shown).

Within the tumor, blood vessels of varying diameters were immunopositive. Staining intensity of the tumor endothelia depended on tumor size and localization. Highest p-gp expression was observed in small neo- plasms, especially in the proliferating peripheral region. In this area, density and staining intensity of the newly formed capillaries were higher than of the blood vessels in the surrounding brain parenchyma. In contrast, when the glioma was implanted into nude mice subcutaneously, the tumor vessels failed to express p-gp (46).

To determine the sensitivity of U-87 MG and U-118 MG glioblastoma cells against paclitaxel and valsparodar, the cells were exposed to the drugs during the whole incubation period of up to 300 hours. Doubling times substantially differed between the two cell lines. For U-87 MG a mean minimal doubling time (t0) of 27.5 hours was determined, whereas proliferation of U-118 MG cells with a t0 of 50 hours was significantly slower (47). At a concentration of 100 nM (a typical peak plasma concentration), paclitaxel was cytostatic on U-87 MG cells, whereas proliferation of the U-118 MG cells had already ceased at 5 nM. At a concentration of 1 µM, valsparodar did not affect proliferation of either cell line. Combining 1 µM valsparodar with weakly cytotoxic concentrations of doxorubicin, etoposide, and paclitaxel, known substrates of p-gp 170, failed to enhance the cytotoxicity of these p-gp substrates. Therefore, both cell lines appear to be well suited to investigate the effect of valsparodar paclitaxel coadministration at the blood-brain barrier.

**Paclitaxel effects on intracerebral growth of human glioma.** Two experiments were carried out to determine whether the rise in the brain levels of paclitaxel in animals pretreated with valsparodar could provide a therapeutic benefit. In the first experiment, human U-87 MG glioblastoma cells were implanted intracerebrally into nude mice. This procedure results in aggressively growing intracerebral neoplasms (48). As described above, this cell line is relatively insensitive to paclitaxel and certain other chemotherapeutic agents in vitro. The animals (n = 7–8) were dosed with 4 mg/kg paclitaxel on day 7 after implantation and with 2 mg/kg paclitaxel on day 14, without and with 50 mg/kg valsparodar 4 hours before each paclitaxel dose. Brains were collected on day 20, and the intracerebral tumor tissue was quantified by morphometry. In this experiment, there were no statistically significant differences in the mean tumor volumes within the three groups: 7.4 ± 1.3 mm³ in the control, 8.6 ± 1.2 mm³ in the paclitaxel, and 7.8 ± 1.3 mm³ in the paclitaxel-valsparodar group. Whereas the mean body weight of the control animals increased by 5%, there was a significant reduction of mean body weight in both treatment groups with a decrease by 9% in the paclitaxel group and a weight loss by 17% in the paclitaxel-valsparodar group during the course of the experiment.

In the second study with intracerebral U-118 MG tumors, paclitaxel dosage was reduced to avoid weight loss. A control group received no medication; a second group received paclitaxel 8 days (3 mg/kg) and 15 days (2 mg/kg) after tumor implantation; a third group received the same regimen of paclitaxel dosing, but in combination with 50 mg/kg valsparodar 4 hours before each paclitaxel dose. Mice were killed 35 days after

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**Figure 4**
Application scheme of PSC 833 and paclitaxel to nude mice. Each group consisted of three to four nude mice per observation time after intravenous injection.

**Figure 5**
Effect of valsparodar coadministration on the paclitaxel brain levels in nude mice after intravenous injection of 8 mg/kg paclitaxel. Four hours before intravenous application the animals were treated with the vehicle of valsparodar by mouth (control, open squares) or 50 mg/kg valsparodar by mouth (valsparodar group, filled squares). Brain levels for a valsparodar group with reduced paclitaxel dose (4 mg/kg, open circles) were investigated 24 hours after intravenous injection. Means ± SEM, n = 3–4. Compared with the respective controls, all values in the valsparodar groups were significantly (P ≤ 0.05) higher.
implantation, and the tumor volume in each brain was determined morphometrically (n = 5). Treatment with paclitaxel alone did not reduce the tumor volume significantly, whereas valspodar plus paclitaxel decreased the tumor volume by 90% (P < 0.05; Figure 6). As shown in Figure 6 (inset), the regimen was not compromised by intolerable toxicity. This result demonstrates that coadministration of valspodar can improve the therapeutic benefit of paclitaxel for brain tumor treatment. Clearly, in vivo the solid U-118 MG tumors were more sensitive to paclitaxel than the solid U-87 MG tumors, a result that agrees with proliferation kinetics and drug sensitivity of these cell lines in vitro (46, 47).

**Effect of valspodar on subcutaneous growth of implanted human glioma.** Recently, valspodar was reexamined as an antitumor agent in vitro (49). We considered the possible contribution of valsodarp to glioma cytotoxicity in an experiment with nude mice bearing U-118 MG glioblastoma subcutaneously. In a 46-day experiment, mice with implanted tumors were given intravenously 120 or 600 mg/kg valspodar, vehicle alone, or no treatment. At the end of the experiment, there were no statistically significant effects on tumor growth among the untreated control group, the vehicle group, and the two valspodar groups (Table 2), nor was there a hint of systemic or CNS toxicity.

**Discussion**

Malignant brain tumors (higher-grade gliomas) are rarely cured by surgery and/or radiotherapy, and chemotherapy has been of limited value, especially as long as the blood-brain barrier remains intact. In addition, the brain is sanctuary for metastases in cancer patients otherwise responding to cytostatic drugs such as paclitaxel (50).

One aim of this study was to find a mechanistic explanation for the low permeability of paclitaxel using in vitro and in vivo studies. Since paclitaxel is excluded from many tumor cells by p-gp, it is likely that this drug efflux pump also limits the access of the drug to the brain. Brain capillaries express p-gp and other ATP binding cassette (ABC) transporters at their luminal membrane (16, 51). To study the impact of ABC transporters on paclitaxel transport we applied a newly developed approach using isolated rat brain capillaries, fluorescent substrates, and confocal imaging. Isolated rat brain capillaries accumulated fluorescently labeled paclitaxel within the lumina. This accumulation was up to 3.5-fold above the cellular accumulation and showed all the hallmarks of specific active transport. It was blocked when metabolism was inhibited by NaCN and when unlabeled drug or the p-gp modifier, valspodar, were added. Unlabeled paclitaxel inhibited the secretion of NBDL-cyclosporin, a p-gp substrate. In cultured endothelial cells from porcine brain, paclitaxel enhanced in a dose-dependent manner the intracellular accumulation of calcine derived from calcein-AM, a p-gp substrate that is otherwise excluded from the cells. All results point to p-gp as a major obstacle to paclitaxel entry into the CNS.

Valspodar is a potent inhibitor of p-gp that alters drug transport across barrier and excretory tissues in vitro and in vivo. As a result, it has the potential to alter the pharmacokinetics of the many therapeutic agents

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**Table 2**

Effect of valspodar on the growth of subcutaneously implanted U-118 MG glioma

<table>
<thead>
<tr>
<th>Treatment group (n = 10)</th>
<th>Mean tumor area ± SEM (mm²)</th>
<th>Day 11</th>
<th>Day 25</th>
<th>Day 36</th>
<th>Day 46</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>3.5 ± 0.47</td>
<td>6.64 ± 0.82</td>
<td>20.2 ± 4.06</td>
<td>24.5 ± 5.62</td>
<td>36.5 ± 10.05</td>
</tr>
<tr>
<td>10 mg/kg valspodar</td>
<td>2.9 ± 0.48</td>
<td>9.1 ± 1.22</td>
<td>18.0 ± 3.60</td>
<td>41.0 ± 9.96</td>
<td>53.7 ± 27.80</td>
</tr>
<tr>
<td>P</td>
<td>(0.423)</td>
<td>(0.104)</td>
<td>(0.699)</td>
<td>(0.141)</td>
<td>(0.550)</td>
</tr>
<tr>
<td>50 mg/kg valspodar</td>
<td>3.4 ± 0.43</td>
<td>7.2 ± 0.88</td>
<td>18.3 ± 2.53</td>
<td>28.3 ± 4.48</td>
<td>35.2 ± 12.94</td>
</tr>
<tr>
<td>P</td>
<td>(0.933)</td>
<td>(0.645)</td>
<td>(0.719)</td>
<td>(0.608)</td>
<td>(0.948)</td>
</tr>
</tbody>
</table>

Valspodar was administered orally twice per week resulting in cumulative doses of 120 and 600 mg valspodar per kilogram. Treatment was started on day 1.

* * *
that are P-gp substrates, leading to longer terminal half lives and lower excretion rates. Such effects have been observed for paclitaxel and other P-gp substrates in clinical and preclinical investigations (23, 36, 37).

Other groups have shown that transporter ablation/inhibition increases drug delivery to the CNS. For example, van Asperen et al. (52) observed the tissue distribution of vinblastine in mice with homozygously disrupted mdr1a genes (mdr1a encodes for P-gp in mice predominantly expressed at the blood-brain barrier) and in the wild-type species. As in our work, they found the highest increase in tissue levels of mdr1a-deficient mice compared with the wild-type species for the brain, where 7–46 times higher vinblastine levels were found, depending on time and dose. Also consistent with our results, vinblastine levels in liver and kidney were significantly higher in knockout mice compared with wild-type animals no earlier than 24 hours after drug administration. The area under the plasma-time curve was found to be 1.4–1.6 times higher in mdr1a-deficient mice. Our results agree in principle with an increased plasma area under the curve. However, this fact cannot explain the six- to eightfold higher paclitaxel concentrations in the brain of valspodar pretreated animals within the first observation period of 1.5 hours, since the plasma levels were only slightly elevated compared with paclitaxel application alone. This supports the interpretation that the inhibition of P-gp by valspodar reduces paclitaxel efflux from the brain. With the available analytical techniques it is impossible to determine whether the drug is homogeneously distributed or accumulated by endothelial or other cells.

Drion et al. (53) perfused brains of rats for 20 seconds with radiolabeled vinblastine and analyzed brain sections. For rats that received 10 mg/kg valspodar intravenously 5 minutes before infusion, nine times higher vinblastine levels were found in all sections of gray brain tissue compared with rats that received no valspodar. These experiments are especially interesting, because the administration of valspodar has no influence on drug concentration in the perfusate and the influence of valspodar on the pharmacokinetics of the cytostatic is of no relevance.

The present results also show that valspodar pre-treatment not only increased brain levels of paclitaxel, but that it also produced a therapeutic effect on a paclitaxel-sensitive transplanted tumor. Although paclitaxel therapy by itself did not affect the volume of intracerebrally implanted U-118 MG tumors, combined valspodar-paclitaxel therapy resulted in a significant decrease in tumor volume (Figure 6). In contrast, neither paclitaxel nor valspodar-paclitaxel therapy affected the volume of implanted U-87 MG tumors derived from a cell line that is not paclitaxel sensitive. Neither glioblastoma cell line exhibited a multidrug-resistance phenotype. Thus, the statistically significant reduction in tumor volume caused by valspodar coadministration was due to increased paclitaxel levels in the brain, a result of inhibition of P-gp-mediated transport at the blood-brain barrier. Circumvention of multidrug resistance at the tumor cell was not a factor. These findings suggest that coadministration of valspodar can be of clinical benefit for the chemotherapeutical treatment of brain tumors sensitive to cytostatics, which are substrates of P-gp. The chemotherapy of other cancers expressing the MDR phenotype (e.g., colon carcinoma and kidney cancer) could also benefit from the coadministration of P-gp inhibitors such as valspodar. In the clinics, the risk of increased (CNS) toxicity, which cannot be totally excluded by our preclinical studies, might require an adjusted paclitaxel dose when used in combination with valspodar or other P-gp inhibitors such as elacridar.

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suchungen zur Optimierung neuer Zytostatica-beladener bioabbaubar.
er Implantate für die interstitielle Chemothepie maligner Hirntum.