Sustained Release for St. John’s Wort: A Rational Idea?

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Abstract

Purpose: Aim of this study was to evaluate St. John’s wort (SJW) extract for its suitability to be formulated in a sustained release dosage form using Ze 117 as example extract.

Methods: Hypericin as marker for naphthodianthrones and quercetin as marker for contained flavonoids in Ze 117 were evaluated for solubility through shake flask method and for in vitro permeability through Caco-2 monolayers. Furthermore, different intestinal segments were screened for absorption capacity of markers using an in situ rat model.

Results: Over a physiological pH range, naphthodianthrones exhibited pH-dependent solubility profiles with best solubility at pH 6.8. In contrast, solubility of flavonoids was pH-independent. In Caco-2 monolayer system, low permeation was evident for naphthodianthrone hypericin, while the flavonoid quercetin showed high permeation. Results of in situ rat model showed absorption of hypericin and quercetin mainly in jejunum.

Conclusion: SJW extract covers components with different physicochemical properties. Predominant absorption in rat intestinal segments indicated presence of an absorption window in small intestine. Furthermore, there is high drug concentration per single dose in combination with a complex extract mixture. Thus, development of a sustained release formulation for SJW extract is challenging.

Keywords: Hypericum perforatum, St. John’s wort, BCS, Caco-2 cells, Intestinal absorption, In situ rat model, Solubility, Bioavailability

Abbreviations: SJW: St. John’s wort; Ze 117: St. John’s wort extract of Max Zeller Söhne AG (Romanshorn, Switzerland); HRS: Herbal reference substance; EP: European Pharmacopoeia; USP: United States Pharmacopoeia; BCS: Biopharmaceutics classification system; ab: Apical-to-basolateral compartment; ba: Basolateral-to-apical compartment; Papp: Apparent permeability; R50: Recovery rate; TEER: Trans-epithelial electrical resistance; PK: Pharmacokinetics; Cmax: Maximum plasma concentration; tmax: Time to maximum plasma concentration; AUC: Area under the curve; AUC∞: Area under the curve related to length of intestinal segment and transit time; SEM: Standard error of the mean; RSD: Relative standard deviation; HPLC: High performance liquid chromatography; UPLC: Ultra-high performance liquid chromatography; UV/Vis: Ultraviolet/visible light (detector for HPLC); MS: Mass spectrometry; ES: MS ionization source in negative ion mode; m/z: Mass to charge ratio; SIR: Selected ion recording; SR: Sustained release.

Introduction

In order to improve patient compliance and to provide an ideal efficiency side effect profile, sustained release (SR) products came in the focus of pharmaceutical development [1]. However, there are hardly any products with sustained release kinetic among herbal medicinal drugs. To target a therapy improvement for St. John’s wort (SJW) products, the aim of this study was to evaluate SJW extract for sustained release suitability. Clinical relevance demonstrates oral administration of SJW as valuable alternative to conventional antidepressant drugs [2,3]. Committee of Herbal Medicinal Products (HPMC) of the European Medicines Agency (EMA) granted Hypericum perforatum with well-established use status for the treatment of mild to moderate depressive episodes as well as for short-term treatment of symptoms in mild depressive disorders [4].

The compounds in SJW cover a large range of substances of which numerous show biological activity. Flavonoids represent the greatest part of secondary metabolites with an amount of 2% up to 4%. Beneath the flavonol glycosides rutin, hyperoside, isorutin and quercetin as well as biflavonoids such as biochanin and amentoflavone [5]. Naphthodianthrones namely hypericin and pseudohypericin occur in small amounts up to 0.3% in the flowers and leaves mainly in the dark glands [6]. The occurrence of the phloroglucinol hyperforin in the above ground parts mainly in the translucent glands [7] is characteristic for SJW. Another phloroglucinol, adhyperforin which is a methyl analogous of hyperforin, occurs in flowers and stems in a concentration of 0.8-1.8%. Additionally, SJW contains the phenolcarbon acids caffeic acid and chlorogenic acid as well as proanthocyanidins, tannins, ethieric oils, fatty acids and chlorophyllin [8]. Pharmacological activity was evident for hypericin [9,10]. However, high doses of pure hypericin were necessary for reaching pharmacological activity, which can be barely achieved by administration of total SJW extract due to the low water solubility of hypericin. An explanation for this conflict in theory and praxis could be that solubility of pure hypericin in water is increased by some phenolic constituents typical for SJW extracts especially by procyanidins and hyperoside [11]. This synergistic effect may lead to an increase in the

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oral bioavailability of hypericin. Accordingly, the pharmacological activity of hypericin and pseudohypericin was enhanced in the forced swimming test when combined with the procyanidin fraction [12]. Also, the combination with rutin leads to a strong increase of the effect in the forced swimming test [13]. In addition, flavonoids such as hyperoside and rutin as well as the quercetin metabolite isorhamnetin were shown to have pharmacologic activity [14]. Correlation of hyperforin with the antidepressant effect of total SJW extract is not ensured [15]. In summary, no mechanism of action neither active compounds of SJW are clearly determined. Therefore, the whole extract is observed as active ingredient and European Pharmacopoeia monographed SJW as quantified extract (Type B1) (EP 9.3/1438).

For quality control of SJW extracts, there have been defined analytical markers in European Pharmacopoeia (EP 9.3/1874), e.g. naphthodianthrones calculated as hypericin, flavonoids and phloroglucinols calculated as hyperforin. However, hyperforin was identified as key component in SJW causing drug-drug interactions Moore [16]. This was the reason for selecting SJW extract Ze 117 for SR evaluation since it contains only low amounts of hyperforin (≤ 0.2%). SJW extracts with low dose hyperforin such as Ze 117 showed reduced or no relevant clinical interactions [17-20]. Hypericin and flavonoids such as quercetin were selected as analytical marker for extract characterization based on EP monograph (EP 9.3/1874). Plasma half-lives of flavonoids contained in SJW extract are about 4 hours [21]. For active substances with short half-lives, SR systems reduce peak plasma concentrations and prolong the presence of therapeutic plasma levels. Half-lives of hypericin in plasma was about between 19 hours [21] up to 29.1 hours [22] and demand no SR formulation. However, flavonoids may benefit from a sustained release delivery system.

To evaluate SR suitability of SJW extract, pH-dependent thermodynamic solubility was determined for analytical markers in Ze 117 using an in vitro shake flask method. Absorption properties were estimated by in vitro permeation through Caco-2 cell monolayers. Finally, different intestinal segments were screened for appropriate absorption capacity using an in situ rat model.

Materials and Methods

SJW dry extract Ze 117 was provided by Max Zeller Söhne AG (Romanshorn, Switzerland). The used batch of Ze 117 (batch no. 142408) contained 0.1% of hyperforin. Hypericin and quercetin were purchased form PhytoLab (Vestenbergsgreuth, Germany). Glycerol was purchased from Merck (Darmstadt, Germany). Wild type male Sprague Dawley rats from Janvier Labs (Le Genest-Saint-Isle, France) were used.

Caco-2 cell culture

For transport assay, Caco-2 cells were seeded on 0.336 cm² PET (polyethylene terephthalate) filters and 24 well plates with a density of 1x10⁶ cells per cm². Caco-2 cells were used 21 days post seeding in passage 74. The cells were maintained in DMEM supplied with 10% fetal bovine serum (FBS), 1% L glutamine, 1% non-essential amino acids and penicillin/streptomycin (10 mg mL⁻¹) at 37°C in an atmosphere of 5% CO₂ and 90% relative humidity. Culture medium was changed every 23 days.

Animals

The in situ study was performed at the Department of Pharmacy and Molecular Biotechnology at University of Heidelberg. All procedures involving animals and their care were conducted according to European Union (EEC Council Directive 86/609, OF L 358, 1; 12 December 1987) and were approved by the local authorities. Wild type male Sprague Dawley rats of 200-300 g weight were acclimatized and fasted overnight with free access to water.

Methods

As herbal extracts contain a complex mixture of physicochemical different constituents, analytical markers were selected. Hypericin and pseudohypericin served as surrogates for contained naphthodianthrones in SJW extract. For flavonoids that represent the greatest part of secondary metabolities in SJW [5], Beneath flavonol glycosides rutin, hyperoside, isoquercetin and quercetin as major flavonoids, also flavonol aglycon quercetin as well as biflavonoid biapigenin served as surrogate. Figure 1 gives an overview about the structural formulas of selected markers. Solubility samples and dissolution samples were quantified for the naphthodianthrones hypericin and pseudohypericin as well as for the flavonoids rutin, hyperoside, isoquercetin, quercetin, quercetin and biapigenin both as single markers and as sum of naphthodianthrones and sum of flavonoids, respectively. In the scope of Caco-2 cell monolayer assay, hypericin was selected to serve as surrogate for naphthodianthrones and quercetin accordingly for flavonoids. In plasma samples, hypericin, pseudohypericin and quercetin were quantified. Additionally, plasma samples were screened for tamarixetin and isorhamnetin as metabolites of quercetin. For considering influence of other extract constituents on the performance of selected analytical marker, experiments were performed using Ze 117 in its entity.

pH-dependent thermodynamic solubility

To determine pH-dependent thermodynamic solubility, shake flask method according to Yalkowsky was used [23], where the test substance was shaken over a defined period of 48 hours in a flask containing the test medium whereby a balance between solid substance (solute) and dissolved substance (solution) appeared. Sodium phosphate buffers of pH 1.2, 2.2, 3.2, 4.5, 5.5, 6.8 and 7.5 were prepared according USP and EP recommendations. Amounts of 2.5 g SJW dry extract Ze 117 each were weighed out in 50 mL Erlenmeyer flasks. After adding 25 mL of the appropriate buffer medium, Erlenmeyer flasks were closed. Then, the flasks were shaken in a water bath Julabo SW 22 at 37°C (± 1°C) and 90 rpm for 48 hours. Samples of buffer solution were taken after 24 and 48 hours by using a 5 mL syringe. Samples were filtered through a syringe glass filter (0.45 μm) by discarding the first mL of the filtrate. Three replicate determinations of solubility for each buffer were performed. The samples were quantified for naphthodianthrones and flavonoids using an HPLC-UV/V is method, which is described below.
Caco-2 cell monolayer permeability assay

Permeation of hypericin and quercetin was measured from apical to basolateral side (ab) and from basolateral-to-apical side (ba) in six-fold determination. The analytical marker substances hypericin and quercetin were dissolved with concentrations of 2.5 µM as single substances, in presence of each other as well as in the matrix of SJW dry extract Ze 117 in HBSS supplemented with 10 mM HEPES (pH 7.4) and 1% DMSO as transport buffer. Caco-2 cells on PET filters in 24-well plates were used for the permeability assay 21 days post seeding. For testing ab permeability, the volume of test solution added to the apical side (donor) was 500 µL whereas the basolateral side (acceptor) was filled with 1000 µL of transport buffer. Accordingly, for testing ba permeability, the volume of test solution added to the basolateral side (donor) was 1100 µL whereas the apical side (acceptor) was filled with 400 µL of transport buffer. During the assay, the well plates were placed on a horizontal shaker at 37 °C and gently shaken for 180 minutes. Samples of the donor compartment of 100 µL each were taken at 0 and 180 minutes. Samples of the acceptor compartment (100 µL) were taken after 60, 120 and 180 minutes and replaced by 100 µL transport buffer. TEER (transepithelial electrical resistance) was measured using a Millicell ERS voltmeter with chopstick electrodes from Merck Millipore (Darmstadt, Germany) at room temperature before and after the Caco-2 cell assay. The substance flux dc/dt (µmol/mL/s) for hypericin and quercetin was calculated through linear regression of the accumulated concentration in the acceptor compartment as a function of time. The apparent permeability $\text{P}_{\text{app}}$ (cm/sec) was calculated according to Artursson [24], where $A$ is the surface area of the membrane (cm$^2$), $C_0$ is the concentration in donor compartment at start of the assay (µmol/mL) and $V_A$ is the volume of the acceptor compartment (cm$^3$):

$$\text{P}_{\text{app}} = \frac{dc/dt \times (1)}{(A \times C_0)} \times V_A$$

Efflux ratio $R_{e}$ was calculated as quotient of apparent permeability in ba direction and apparent permeability in ab direction:

$$R_{e} = \frac{(\text{P}_{\text{app}} \text{ ba})}{(\text{P}_{\text{app}} \text{ ab})}$$

The recovery rate $R_M$ was calculated by examining the amount of the analytical marker, respectively, in donor compartment $n_{D}$ and
acceptor compartment $n_a$ after 180 minutes in comparison with the amount of the analytical marker in donor compartment $n_d$ before start of the assay:

$$R_{\text{ml}} = \frac{n_d + n_{\text{ca}}}{n_d}$$

The TEER values ($Ω \times \text{cm}^2$) were calculated as product of measured resistance $R$ ($Ω$) in Caco-2 cell monolayers, corrected by measured resistance without cells $R_{\text{blank}}$, and the surface area of the used filters ($0.336 \text{ cm}^2$):

$$\text{TEER} = (R - R_{\text{blank}}) \times A$$

In order to evaluate determined $P_{\text{ml}}$ values and efflux ratios, permeation of reference substances propranolol, metoprolol, verapamil, digoxin, furosemide, hydrochlorothiazide and FITC dextran was determined analogously. (Figure A1)

**In situ absorption study**

SJW dry extract Ze 117 was dissolved in a mixture of glycerol and purified water (1:1) in a concentration of 25 mg/mL in addition of hypericin (0.3 mg/mL). Male Sprague Dawley rats were fasted overnight with free access to water and then anesthetized by intraperitoneal injection of 50 mg/kg pentobarbital prior to the experiment. The peritoneum was opened by a midline incision and approximately 5 cm of the appropriate intestinal segment (duodenum, jejunum, ileum or colon) was ligated by using a medicinal suture. Ligation was performed in a way to avoid pass over of intestinal content but keeping an intact intestinal blood supply. The herbal extract solution of SJW dry extract was administered directly into the ligated intestinal segment in a concentration of 37.5 mg or 75.0 mg Ze 117 dry extract per kg rat body weight. Injection volumes were between 250 µL and 750 µL, dependently on dose and rat body weight. Blood samples of 200 µL were taken at 0, 30, 60, 120 and 180 minutes from the portal vein to minimize influence of hepatic first pass effect. Blood samples were filled in Li Heparin microvettes and centrifuged for 5 minutes at 16'000 g. The plasma was separated and stored in appropriate vials at – 20°C until analytical quantification via UPLC MS/MS as described below. A 4-fold determination was performed.

**Analytical methods**

For quantification of naphthodianthrones and flavonoids in solubility samples, dissolution samples, Caco-2 cell samples and plasma samples, HPLC-UV/Vis or UPLC MS/MS methods were used. Table 1 lists an overview.

Assay of naphthodianthrones and flavonoids in solubility and dissolution samples was performed via HPLC UV/Vis based on European Pharmacopeia. The system consisted of a Waters Acuity quaternary pump with autosampler coupled to a Waters diode array detector (Waters AG, Baden Daettwil, Switzerland) and connected to a PC running Waters Empower software. Liquid chromatography for naphthodianthrones was performed using 39 volumes of ethyl acetate mixed with 41 volumes of a 1.5 g/L solution of sodium dihydrogen phosphate adjusted to pH 2 with phosphoric acid and 160 volumes of methanol as mobile phase. An isotropic mode with a flow rate of 1.0 mL/min was used. Injection volume was set to 20 µL. Separation was performed on a Waters HPLC column Atlantis dC18, 150 x 4.6 mm, 5 µm (Waters AG, Baden Daettwil, Switzerland) at a column temperature of 40°C. Detection wavelength was 590 nm and as reference substance, SJW dry extract HRS was used. LC for flavonoids was performed using 0.3% phosphoric acid in water as aqueous phase (A) and 0.3% phosphoric acid in acetonitrile as organic phase (B) in a gradient mode of % A (t(min)), 82 (0.0) – 82 (8.0) – 47 (18.0) – 3 (18.1) – 3 (31.0) – 82 (31.1) – 82 (33.0). Injection volume was set to 10 µL and the pump flow rate to 1.0 mL/min. Separation was performed on a Phenomenex HPLC column Luna C18, 150 x 4.6 mm, 3 µm

### Table 1: Analytical methods for quantification of naphthodianthrones and flavonoids.

<table>
<thead>
<tr>
<th>Assay application</th>
<th>Analyte</th>
<th>System</th>
<th>Column</th>
<th>Mobile phase</th>
<th>Gradient</th>
<th>Oven temp.</th>
<th>Flow rate</th>
<th>Injection volume</th>
<th>Detection</th>
</tr>
</thead>
<tbody>
<tr>
<td>solubility, dissolution</td>
<td>naphthodianthrones</td>
<td>HPLC-UV/Vis$^*$</td>
<td>Atlantis dC18 150 x 4.6 mm 5 µm</td>
<td>Ethyl acetate, NaNH$_2$PO$_4$, 15.6 g/L pH 2, methanol 39:41:160 V/V</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>40</td>
<td>1.0</td>
</tr>
<tr>
<td>solubility, dissolution</td>
<td>flavonoids</td>
<td>HPLC-UV/Vis$^*$</td>
<td>Luna C18 150 x 4.6 mm 3 µm</td>
<td>Water, phosphoric acid 1000:3 V/V</td>
<td>Acetonitrile, phosphoric acid 1000:3 V/V</td>
<td>0.0</td>
<td>8.0</td>
<td>82</td>
<td>20</td>
</tr>
<tr>
<td>in vitro Caco-2 assay, in situ studies in rat</td>
<td>hypericin</td>
<td>UPLC-MS/MS$^*$</td>
<td>Acquity BEH C18 100 x 2.1 mm 1.7 µm</td>
<td>Ammonium acetate in water 10 mM</td>
<td>Acetonitrile</td>
<td>0.0</td>
<td>2.0</td>
<td>40</td>
<td>50</td>
</tr>
<tr>
<td>in vitro Caco-2 assay, in situ studies in rat</td>
<td>quercetin</td>
<td>UPLC-MS/MS$^*$</td>
<td>Acquity BEH C18 100 x 2.1 mm 1.7 µm</td>
<td>Purified water, 0.1 % Formic acid</td>
<td>Acetonitrile, 0.1 % Formic acid</td>
<td>0.0</td>
<td>4.0</td>
<td>90</td>
<td>50</td>
</tr>
</tbody>
</table>

$^*$HPLC-UV/Vis on a Waters Acuity quaternary pump with autosampler coupled to a Waters diode array detector (Waters AG, Baden Daettwil, Switzerland) and connected to a PC running Waters Empower software

$^*$UPLC-MS/MS on a Waters H-class system with an autosampler and a quaternary pump coupled to a Waters Xevo TQ-S tandem (triple) quadrupole spectrometer and Waters MassLynx software. Detection in multiple reaction monitoring (MRM) with an atmospheric pressure ionization source in negative ion mode (ES-), capillary voltage of 2.5 kV, source temperature of 450°C and a sampling rate of 2 Hz, transition of 503.10 to 405.07 for hypericin and 519.22 to 486.96 for pseudohypericin, transition of 301.16 to 151.05 for quercetin and 315.16 to 151.00 for tamarixin and isorhamnetin.


(Phenomenex, Torrance, CA, USA) at a column temperature of 20 °C. Detection wavelength was 360 nm and as reference substance, rutin trihydrate was used. Both naphthodianthrones hypericin and pseudohypericin were calculated as hypericin whereas flavonoids were calculated as rutin.

Assay for hypericin and quercetin in Caco-2 cell samples was performed via UPLC/MS/MS. The system consisted of a Waters Acquity quaternary pump with autosampler coupled to a Waters Xevo TQ S spectrometer (Waters AG, Baden Daettwil, Switzerland) and connected to a PC running Waters MassLynx software. Separation was done on an Acquity UPLC column BEH C18, 150 x 2.1 mm, 1.7 µm (Waters AG, Baden Daettwil, Switzerland). LC for hypericin was performed using 10 mM ammonium acetate in water as aqueous phase (A) and acetonitrile as organic phase (B) in a gradient mode of % A (t(min)), 90 (0.0) – 40 (4.0) – 5 (4.1) – 5 (5.0) – 90 (5.1) – 90 (6.5). Injection volume was set to 10 µL and the pump flow rate to 0.7 mL/min. Transition of (m/z) 503.10 to the monitoring ion (m/z) 405.07 for hypericin and (m/z) 519.22 to the monitoring ion (m/z) 486.96 for pseudohypericin was detected. An atmospheric pressure ionization source in negative ion mode (ESI-) with capillary voltage of 2.5 kV, source temperature of 450°C and a sampling rate of 2 Hz was used. LC for quercetin and its metabolites tamarixetin and isorhamnetin was performed using 0.1 % formic acid in water as aqueous phase (A) and acetonitrile as organic phase (B) in a gradient mode of % A (t(min)), 90 (0.0) – 40 (4.0) – 5 (4.1) – 5 (5.0) – 90 (5.1) – 90 (6.5). Injection volume was set to 10 µL and the pump flow rate to 0.5 mL/min. Transition of (m/z) 301.16 to the monitoring ion (m/z) 151.05 for quercetin in an atmospheric pressure ionization source in negative ion mode (ESI-) with capillary voltage of 2.5 kV, source temperature of 450°C and a sampling rate of 2 Hz was used.

Plasma samples from rat taken after intestinal administration of Ze 117 were pooled for each sampling point to enable sample work up for naphthodianthrones and additionally for quercetin. For determination of hypericin and pseudohypericin, 50 µL of each plasma sample were pipetted into an Eppendorf tube. 1.5 mL ethylacetate were added and the samples were shaken overnight for 10 minutes. The samples were frozen out overnight using a refrigerator and then decanted into a test tube. Then, the samples were evaporated to dryness at 40 °C under a gentle stream of nitrogen. The residue was dissolved in 100 µL methanol. Supernatant was transferred into a glass vial. For sample work up of quercetin, 50 µL of each plasma sample were pipetted into an Eppendorf tube. 5 µL of internal standard solution (0.05 ng/µL quercetin d3 in methanol/0.5 % ascorbic acid 1:1 (V/V)) were added and the samples were briefly shaken. Then, 10 µL sodium acetate solution 0.25 % in methanol and 5 µL glucuronidase solution (1:100 in water) were added. Samples were briefly shaken again and incubated for 5 hours at 40°C. Subsequently, 10 µL 1N HCl was added and samples were shaken briefly. After addition of 1.25 mL ethyl acetate, samples were shaken over night using a refrigerator. After, the supernatants were decanted into a test tube and evaporated to dryness at 40°C under a gentle stream of nitrogen. The residue was dissolved in 100 µL methanol. Supernatant was transferred into a glass vial. To each sample, 50 µL of 0.1% formic acid were added. Concentration of herbal marker in plasma was measured via UPLC MS/MS using same analytical methods as described for Caco-2 cell samples.

Results

Solubility of naphthodianthrones was pH-dependent

The pH-dependent thermodynamic solubility of naphthodianthrones in SJW dry extract Ze 117 is outlined in Figure 2 and supplemental data A2. Hypericin exhibited pH dependence in its solubility profile with the lowest measured solubility at pH 1.2. With a value of 23.78 ± 0.43 µg/mL, saturation solubility of hypericin at pH 6.8 was about 5-fold higher than the one at pH 1.2, which showed a value of 4.61 ± 0.03 µg/mL. Same findings were evident for pseudohypericin, where the highest measured concentration of 9.84 ± 0.07 µg/mL at pH 5.5 was about 2.5-fold higher than saturation solubility at pH 1.2. In contrast to hypericin, the lowest concentration for pseudohypericin (2.97 ± 0.14 µg/mL) was measured at pH 7.5. Determined values after 48 hours were slightly lower than that measured after 24 hours except for maximum concentrations where no time dependent difference was evident.

Solubility of flavonoids was not pH-dependent

Main flavonoids contained in SJW extract Ze 117 showed no pH dependence in their solubility profiles (Figure 2 and supplemental data Table A1, A2). Saturation concentration was between 4.56 ± 0.06 and 5.90 ± 0.37 mg/mL for total flavonoids expressed as sum of rutin, hyperoside, isoquercitrinose, quercitrin and biapigenin. Single flavonoids exhibited saturation concentrations correlating to their amount in SJW extract. Main constituents were hyperoside and rutin followed by isoquercitrinose and quercitin. Determined values after 48 hours were comparable to that measured after 24 hours.

Hypericin showed low permeation through Caco-2 cell monolayers

For permeation assessment of Ze 117, hypericin was used as surrogate for naphthodianthrones. Results are shown in Table 2. Hypericin as pure substance showed a P app in ab direction of 3.19 ± 0.72 x 10-6 cm/s confirming literature data [25,26]. Hypericin in the presence of quercetin showed an enhanced permeation with a P app of 5.23 ± 1.13 x 10-6 cm/s. A permeation enhancement of hypericin through Caco-2 cell monolayers induced by the presence of flavonoids such as quercitin was also found by Verjee [25]. Indeed, permeation of hypericin was observed to be lower when measured in the matrix of Ze 117. Here, P app value for ab permeation was 1.48 ± 0.51 x 10-6 cm/s. Hypericin as pure substance showed for permeation in ab and in ba direction similar P app values. P app value for ba permeation decreased in the presence of quercitin and additionally in a much higher extent, in the presence of Ze 117. Overall, determined P app values for hypericin were comparable to that of low permeable reference substances (Figure 3). Efflux ratio of 0.94 for pure hypericin decreased markedly in presence of quercitin and Ze 117, respectively. Comparing TEER values measured before and after performed Caco-2 cell monolayer assay, presence of hypericin determined a significant decrease in cell monolayer integrity. A reduction in TEER values of approximately 25% was evident in Caco-2 cells treated with pure hypericin. In presence of hypericin and quercitin, there was a 45% reduction and in presence of Ze 117, TEER values declined even by 63% (Figure 4). In contrast, TEER values were comparable before and after performed Caco-2 cell monolayer assay for Caco-2 cells treated with reference substances or quercitin as pure substance, respectively.

Quercetin showed high permeation though Caco-2 cell monolayers

Quercetin as marker for flavonoids in Ze 117 permeated through Caco-2 cell monolayers with a P app value of 33.76 ± 5.35 x 10-6 cm/s in ab direction similar P app values. P app value for ba permeation decreased in the presence of quercitin and additionally in a much higher extent, in the presence of Ze 117. Overall, determined P app values for hypericin were comparable to that of low permeable reference substances (Figure 3). Efflux ratio of 0.94 for pure hypericin decreased markedly in presence of quercitin and Ze 117, respectively. Comparing TEER values measured before and after performed Caco-2 cell monolayer assay, presence of hypericin determined a significant decrease in cell monolayer integrity. A reduction in TEER values of approximately 25% was evident in Caco-2 cells treated with pure hypericin. In presence of hypericin and quercitin, there was a 45% reduction and in presence of Ze 117, TEER values declined even by 63% (Figure 4). In contrast, TEER values were comparable before and after performed Caco-2 cell monolayer assay for Caco-2 cells treated with reference substances or quercitin as pure substance, respectively.

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Figure 2: pH-dependent solubility profile of hypericin and pseudohypericin calculated as hypericin as well as pH-dependent solubility profile of rutin, hyperoside, isoquercitrin, quercitrin, quercetin and bioapigenin calculated as rutin in buffer media of different pH values after 24 hours and after 48 hours, values are given as average of saturation concentration ± SEM (n=3).
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Direction (Figure 3 and Table 2). Permeation differed not significantly in the presence of hypericin and Ze 117, respectively. Permeation in ba direction was similar to Papp values of ab permeation and independent of quercetin and St. John’s wort extract Ze 117. Overall, determined Papp values for quercetin were comparable to that of high permeable reference substances. Efflux ratio was found to be between 0.99 and 1.15 (Figure A3).

Hypericin was absorbed in situ mainly in jejunum

Absorption of naphthodianthrones expressed as sum of hypericin and pseudohypericin exhibited great differences between duodenum, jejunum, ileum and colon (Table 3 and Figure 5). There was an evident absorption in upper intestine that decreased significantly the lower the intestinal region. Administration of Ze 117 extract solution into duodenum resulted in the highest plasma levels of hypericin and pseudohypericin. Plasma concentration showed delayed absorption with maxima after two hours, what correlates with information in HMPC monograph [4]. Plasma AUC of the two naphthodianthrones after jejunal absorption amounted approximately 60-70% of that of duodenal absorption. In contrast to duodenal absorption, in jejunum there was no delayed absorption evident. Even after 30 minutes there were naphtodianthrone concentrations detectable in plasma samples. In ileum and colon, there was nearly no absorption of naphtodianthrones. Plasma profiles varied proportional to the dose. Determined plasma concentrations were correlated to 3 hours measurement in intestinal segment each ligated for the same length. As both length of intestinal segments and transit times differ in vivo, measured absorption capacity had to be corrected by taking into account physiological conditions. Table 4 lists human intestinal transit times and length of single intestinal segments according GastroPlus™ software. Calculated AUC for each intestinal segment was related to appropriate human length and transit time and termed as AUCphys (Figure A3). With correlation

Table 2: Apparent permeability Papp, recovery rate RM, efflux ratio RE and amount of permeated substance through Caco-2 cell monolayer after 180 min in ab and ba directions (n=6), values are given as average ± SEM.

<table>
<thead>
<tr>
<th>Substance</th>
<th>Papp [10⁻⁶ cm/s] ab</th>
<th>Papp [10⁻⁶ cm/s] ba</th>
<th>RM ab</th>
<th>RM ba</th>
<th>RE</th>
<th>Permeated ab [%]</th>
<th>Permeated ba [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td>FITC-dextran</td>
<td>0.64 ± 0.29</td>
<td>-</td>
<td>1.13 ± 0.06</td>
<td>-</td>
<td>-</td>
<td>0.8 ± 0.05</td>
<td>-</td>
</tr>
<tr>
<td>Propranolol</td>
<td>24.23 ± 3.60</td>
<td>34.13 ± 1.62</td>
<td>0.69 ± 0.06</td>
<td>1.12 ± 0.04</td>
<td>1.41 ± 0.18</td>
<td>27.4 ± 5.96</td>
<td>13.6 ± 0.60</td>
</tr>
<tr>
<td>Metoprolol</td>
<td>30.12 ± 2.35</td>
<td>33.64 ± 1.74</td>
<td>0.63 ± 0.08</td>
<td>1.08 ± 0.03</td>
<td>1.12 ± 0.1</td>
<td>30.6 ± 1.45</td>
<td>12.1 ± 0.78</td>
</tr>
<tr>
<td>Verapamil</td>
<td>26.54 ± 1.24</td>
<td>36.30 ± 1.58</td>
<td>0.66 ± 0.05</td>
<td>1.16 ± 0.04</td>
<td>1.37 ± 0.11</td>
<td>28.4 ± 0.01</td>
<td>13.0 ± 0.01</td>
</tr>
<tr>
<td>Digoxin</td>
<td>1.48 ± 0.43</td>
<td>21.28 ± 2.16</td>
<td>0.97 ± 0.05</td>
<td>0.93 ± 0.03</td>
<td>1.40 ± 4.76</td>
<td>1.3 ± 0.23</td>
<td>5.6 ± 0.38</td>
</tr>
<tr>
<td>Furosemide</td>
<td>0.99 ± 0.23</td>
<td>6.67 ± 0.34</td>
<td>0.59 ± 0.01</td>
<td>0.84 ± 0.01</td>
<td>0.73 ± 1.27</td>
<td>0.8 ± 0.00</td>
<td>2.1 ± 0.00</td>
</tr>
<tr>
<td>Hydrochlorothiazide</td>
<td>1.16 ± 0.24</td>
<td>1.29 ± 0.02</td>
<td>0.67 ± 0.02</td>
<td>0.88 ± 0.01</td>
<td>1.11 ± 0.04</td>
<td>0.7 ± 0.00</td>
<td>0.4 ± 0.00</td>
</tr>
<tr>
<td>Hypericin pure</td>
<td>3.19 ± 0.72</td>
<td>3.00 ± 1.47</td>
<td>0.89 ± 0.08</td>
<td>0.88 ± 0.04</td>
<td>0.94 ± 1.28</td>
<td>8.3 ± 1.67</td>
<td>0.8 ± 0.31</td>
</tr>
<tr>
<td>Hypericin in presence of quercetin</td>
<td>5.23 ± 1.13</td>
<td>2.80 ± 1.65</td>
<td>0.85 ± 0.06</td>
<td>0.56 ± 0.12</td>
<td>0.54 ± 0.22</td>
<td>4.8 ± 0.91</td>
<td>0.7 ± 0.38</td>
</tr>
<tr>
<td>Hypericin in matrix of Ze 117</td>
<td>1.48 ± 0.51</td>
<td>0.35 ± 0.07</td>
<td>0.63 ± 0.05</td>
<td>1.10 ± 0.10</td>
<td>0.24 ± 0.27</td>
<td>1.2 ± 0.32</td>
<td>0.1 ± 0.02</td>
</tr>
<tr>
<td>Quercetin pure</td>
<td>33.76 ± 5.35</td>
<td>33.29 ± 6.76</td>
<td>0.83 ± 0.18</td>
<td>0.90 ± 0.16</td>
<td>0.99 ± 0.26</td>
<td>21.0 ± 2.97</td>
<td>11.6 ± 2.06</td>
</tr>
<tr>
<td>Quercetin in presence of hypericin</td>
<td>29.38 ± 11.93</td>
<td>33.73 ± 5.32</td>
<td>0.87 ± 0.05</td>
<td>1.12 ± 0.34</td>
<td>1.15 ± 0.89</td>
<td>23.6 ± 5.18</td>
<td>12.8 ± 2.43</td>
</tr>
<tr>
<td>Quercetin in matrix of Ze 117</td>
<td>26.30 ± 8.95</td>
<td>27.62 ± 6.70</td>
<td>0.86 ± 0.17</td>
<td>0.88 ± 0.13</td>
<td>1.05 ± 0.14</td>
<td>28.1 ± 13.46</td>
<td>7.7 ± 1.76</td>
</tr>
</tbody>
</table>

Figure 3: Apparent permeability Papp of reference substances as well as hypericin and quercetin as analytical marker substances of Ze 117 through Caco-2 cell monolayers in ab and ba directions, values are given as average ± SEM (n=6-9), significance is indicated by asterisk p < 0.05.
Figure 4: TEER values of Caco-2 cell monolayers after treatment with reference substances, hypericin and quercetin as pure substances and Ze 117, values given as mean ± SEM (n=6-9), significance is indicated by asterisk p < 0.001.

Figure 5: Plasma profiles as sum of hypericin and pseudohypericin calculated as hypericin (overhead) as well as sum of quercetin (beneath) after intestinal administration of Ze 117 in doses of 37.5 mg/kg rat body weight in addition of hypericin corresponding to 5.0 mg/kg (left) and administration of Ze 117 in doses of 75.0 mg/kg rat body weight in addition of hypericin corresponding to 10.0 mg/kg (right), values given as mean ± SEM (n=3-4).
of measured AUC for hypericin in rat plasma with human intestinal length and transit times, duodenum showed lowest absorption capacity as length and transit time are relatively short compared to other intestinal segments. Highest absorption capacity was calculated for ileum. Also for ileum and colon, absorption capacity was evident that was calculated to be a third of that in jejunum.

**Quercetin was absorbed in situ mainly in small intestine**

Quercetin was measured in plasma samples as surrogate for flavonoids contained in Ze 117. Highest AUC of quercetin was found after administration of Ze 117 extract solution into jejunum followed by that in duodenum (Figure 5 and Table 3). Administration into ileum resulted in markedly lower plasma levels whereas nearly no absorption of quercetin was found in colon. Absorption maxima were between 30 and 60 minutes. Plasma profiles varied proportional to the dose. Metabolites of quercetin such as tamarixetin and isorhamnetin were not found in plasma samples. Taking into account both length of intestinal segments and transit times, duodenum and colon showed very low absorption capacity for quercetin. Best performance regarding quercetin absorption was calculated for jejunum followed by ileum (Table 4 and Figure A3).

### Discussion

For a solid drug substance, release out of the dosage form (liberation) is crucial prior to absorption in human gastro intestinal tract. Sustained release dosage forms exhibit slow drug release with controlled dissolution profiles. Here, dissolution is the rate-determining step in absorption. Nevertheless, drug solubility in aqueous media presents an important physicochemical parameter for selection of a suitable sustained release system. Over a physiological pH range, pH-dependent thermodynamic solubility of naphthodianthrones and flavonoids contained in SJW extract Ze 117 was determined. Both regarded naphthodianthrones hypericin and pseudohypericin showed pH-dependence in their solubility profiles with highest measured concentration at pH 6.8 for hypericin and pH 5.5 for pseudohypericin. Solubility at acidic pH values was much lower for both naphthodianthrones. High concentration of protons in acidic media may determine protonation of hypericin at a carbonyl group and thus, change of the chromophoric properties [26,27]. At basic pH values, a deprotonation of a hydroxyl group may lead to an analogous effect. pKa values of hypericin are 2 and 11 [27]. The large conjugated system of hypericin leads to good stabilization of positive or negative charging. An accompanied change in absorption maximum may result in lower concentrations measured at predetermined detection wavelength [28]. Some considerations apply to pseudohypericin. Indeed, lowest solubility for pseudohypericin was at pH 7.5. Corresponding to pKa value of 7.2 [29]. Furthermore, naphthodianthrones form with the large conjugated system insoluble aggregates in aqueous solutions [30]. These stacking aggregates may result in low aqueous solubility. Solubility of pure hypericin in water is increased by some phenolic constituents typical for SJW extracts [11]. As pH-dependent thermodynamic solubility was determined for hypericin and pseudohypericin in the matrix of Ze 117, other constituents may influence solubility of hypericin and pseudohypericin in a positive way. Lowest measured saturation solubility for hypericin was 4.61 ± 0.03 µg/mL at pH 1.2, whereas for pseudohypericin lowest value was 2.97 ± 0.14 µg/mL at pH 7.5. Overall, determined saturation solubility of hypericin was higher than that of pseudohypericin. SJW extracts of European origin exhibit characteristic pseudohypericin ratios of 1.5-2:1 as pseudohypericin exhibiting higher hydrophilicity than hypericin and is extracted in a higher quantity

### Table 3: \(T_{\text{max}}\), \(C_{\text{max}}\) and AUC of hypericin and quercetin in plasma samples obtained from in-vivo study, values are given as average ± SEM (n=4).

<table>
<thead>
<tr>
<th>Analytical markers</th>
<th>Dose administered</th>
<th>Intestinal segment</th>
<th>PK parameters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hypericin, pseudo- hypericin</td>
<td>37.5 mg/kg Ze 117</td>
<td>duodenum 120</td>
<td>(T_{\text{max}}) [min]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>jejenum 120</td>
<td>59.99 ± 4.86</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ileum 120</td>
<td>2.79 ± 0.29</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Colon 60</td>
<td>0.52 ± 0.07</td>
</tr>
<tr>
<td></td>
<td>75.0 mg/kg Ze 117</td>
<td>duodenum 120</td>
<td>(C_{\text{max}}) [ng/mL]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>jejenum 120</td>
<td>24.40 ± 1.84</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ileum 120</td>
<td>3.90 ± 0.56</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Colon 60</td>
<td>2.40 ± 0.05</td>
</tr>
<tr>
<td></td>
<td>37.5 mg/kg Ze 117</td>
<td>duodenum 120</td>
<td>AUC [µg min/mL]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>jejenum 120</td>
<td>47.8 ± 0.46</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ileum 120</td>
<td>2.25 ± 0.01</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Colon 60</td>
<td>0.59 ± 0.03</td>
</tr>
<tr>
<td></td>
<td>75.0 mg/kg Ze 117</td>
<td>duodenum 120</td>
<td>(T_{\text{max}}) [h]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>jejenum 120</td>
<td>30 ± 1.684 ± 30.88</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ileum 120</td>
<td>185.56 ± 2.20</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Colon 60</td>
<td>27.13 ± 1.54</td>
</tr>
<tr>
<td>Quercetin</td>
<td>37.5 mg/kg Ze 117</td>
<td>duodenum 30</td>
<td>(C_{\text{max}}) [ng/mL]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>jejenum 30</td>
<td>1'581.70 ± 79.77</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ileum 60</td>
<td>121.09 ± 2.21</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Colon 60</td>
<td>185.56 ± 2.20</td>
</tr>
<tr>
<td></td>
<td>75.0 mg/kg Ze 117</td>
<td>duodenum 30</td>
<td>AUC [µg min/mL]</td>
</tr>
<tr>
<td></td>
<td></td>
<td>jejenum 30</td>
<td>83.40 ± 11.01</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ileum 60</td>
<td>5.51 ± 0.63</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Colon 120</td>
<td>6.40 ± 1.25</td>
</tr>
</tbody>
</table>

### Table 4: AUC of naphthodianthrones hypericin and pseudohypericin in plasma and AUC of quercetin in plasma related to human transit time and intestinal length.
by using extraction solvent of ethanol water mixtures [31]. However, results of pH-dependent solubility study did not reflect this pseudohypericin hypericin ratio. Obviously, constituents of Ze 117 influence solubility of hypericin and that of pseudohypericin in a different manner. When evaluated according to class boundaries of FDA Guidance for Industry “Waiver of in vivo bioavailability and bioequivalence studies for immediate-release solid oral dosage forms based on a Biopharmaceutics Classification System (BCS)” [32], criteria “highly soluble” is fulfilled if the highest dose strength is soluble in less than 250 mL buffer over a pH range of 1 to 7.5. With regard to the lowest determined saturation concentration, hypericin can be regarded as “highly soluble” up to maximal dose strength of 1.15 mg according BCS. The “highly soluble” limit for pseudohypericin was calculated to be 0.74 mg. Compared to the drug product Remotiv 250 (Max Zeller Söhne AG, Romanshorn, Switzerland) containing 250 mg SJW dry extract Ze 117 and a total content of hypericin and pseudohypericin in the range of 0.1-0.3% (corresponding to 0.25-0.75 mg), both naphthodianthrones can be regarded as highly soluble substances according BCS. However, there is not a large margin between determined “highly soluble” limit and single dose concentration of naphthodianthrones. If compared to the drug product Remotiv® 500 (Max Zeller Söhne AG, Romanshorn, Switzerland) containing 500 mg SJW dry extract Ze 117, pseudohypericin has to be classified as “low soluble” substance according BCS criteria. However, low solubility resulting of a closed in vitro system has a merely limited prediction for in vivo performance. As in vivo, concentration of solute drug is decreased continuously by absorption, reaching of saturation solubility in the intestinal tract is not presumable. Based on the results for pH-dependent thermodynamic solubility of naphthodianthrones, there was no indication for difference in in vivo performance of Remotiv 250 and Remotiv® 500. Hypericin as marker for naphthodianthrones in Ze 117 showed a low permeation when investigated using Caco-2 cell monolayers. Presence of quercetin resulted in an enhanced ab permeation and a slightly decreased ba permeation of hypericin. Ze 117 seemed to influence both ab and ba permeation of hypericin in a negative way. Interaction with efflux transporter proteins such as P-glycoprotein (P- gp) was not indicated as permeation in ab direction was higher than that in ba direction. As P app values observed for hypericin were comparable to that of reference substances with low permeation, hypericin can be regarded as low permeable substance and was classified as BCS II substance. Comparison of pure hypericin with performance of quercetin and extract matrix of SJW extract Ze 117, respectively, was determined within the same concentration ranges for hypericin to ensure comparability of results. Considering measured TEER values of Caco-2 cell monolayers treated with hypericin, a decrease was evident. Observed reduction in TEER values indicated a loss of Caco-2 cell monolayer integrity. Probably, used hypericin concentration compromised cell viability and Caco-2 cell monolayers treated with lower hypericin concentrations may show more descriptive results. TEER measurement of Caco-2 cell monolayers treated with Ze 117 showed lowest values after the assay, matrix of Ze 117 seemed to intensify cell viability impairment caused by hypericin. In this study, hypericin concentration was selected based on the analytical limit of quantification. For lower hypericin concentrations than used, there was no permeation detectable especially in acceptor compartments at the beginning of the assay. Low TEER values indicate impairment of cell monolayers. A non-sufficient monolayer integrity may result in overestimation of permeability. In fact, determined P app values were higher than they should be. Mass transfer of hypericin over 3 hours showed linear progression, so significant increase of hypericin permeation during the assay was not evident. This finding encourages sufficient Caco-2 monolayer integrity despite of decreasing TEER values. A general minimum limit of TEER indicating a non-sufficient cell monolayer integrity is hard to determine as TEER values vary depending on used equipment [33] and other factors such as temperature and used media [34]. In addition, decrease in TEER is proportional with an increased electron flux through cell monolayers. As electrons are much smaller than drug molecules, decreasing TEER values do not correlate proportionally to cell monolayer permeability for drug molecules and drug substance flux. However, an overestimation of permeation would not change BCS classification. Hypericin is still classified as low permeable substance even when exhibiting lower P app values.

Assessing flavonoids in Ze 117, thermodynamic solubility for rutin, hyperoside, isoorcitsinose, isoorcitsidine, quercetin and baipigenin exhibited no pH- dependence. According BCS criteria, flavonoids can be regarded as “highly soluble” up to maximal dose strength of 1.14 g. As products of SJW such as Remotiv® 250 and Remotiv® 500 contain single doses of SJW dry extract Ze 117 of 250 mg and 500 mg, respectively, flavonoids can be classified generally as “highly soluble”. Quercetin as marker for flavonoids in Ze 117 showed a high permeation when investigated using Caco-2 cell monolayers. Based on the findings, quercetin could be classified into BCS class I. Interaction with efflux transporter proteins such as P glycoprotein (P-gp) was not indicated. However, for highly permeable substances identification of efflux transporter substrates is hard to determine as high passive diffusion would not be altered significantly by active efflux transporters. To identify efflux transporter substrates, presence of an efflux transporter inhibitor is needed during Caco-2 cell monolayer assay.

Permeation of several reference substances measured through Caco-2 cell monolayers exhibited a good in vitro in vivo correlation [35]. However, Caco-2 cell assay results of hypericin and quercetin in SJW extract Ze 117 admit just an estimation about the extent of absorption. No prediction of absorption performance of different intestinal segments could be done. For sustained release dosage forms, a continuous drug absorption along the whole intestine is necessary for most systems. To evaluate the absorption performance of hypericin and quercetin in Ze 117 in dependence of intestinal regions, an in situ study was performed in rat. Hence, the aim was to compare local absorption capacity of different intestinal segments in order to identify possible absorption windows and determine a rational timeline for drug release. Results showed the highest absorption capacity of hypericin after administration of Ze 117 in jejunum. Significantly lower performance was found after administration in duodenum, ileum and colon. Furthermore, a delayed absorption with maximal plasma concentration of hypericin after 2 hours was obvious in rat duodenum. As human transit time in duodenum is assumed to be 0.3 hours at the mean, duodenal absorption processes are rather irrelevant for naphthodianthrones. For most SR formulations, drug absorption along the whole intestinal tract is an important requirement for proper performance of the delivery system. If there is sufficient in vivo absorption capacity for hypericin along human ileum and colon when drug is released out of a SR system, could be answered best by a pharmacokinetic in vivo study in humans. Quercetin was found to be absorbed mainly in small intestine, whereas highest absorption capacity was found after administration of Ze 117 into jejunum followed by that into ileum. Duodenum showed a low absorption capacity as length and transit time are relatively short compared to other intestinal segments. In colon, there was nearly no absorption of quercetin. With regard to the large microbial system present in colon [36], there is the possibility that quercetin underwent microbial metabolism in a faster way than
absorption processes took place. Metabolites were assumed to be small unspecific molecules such as phloroglucinol and acid derivatives [37] or carbon dioxide [38]. As plasma samples were not screened for these metabolites, low absorption capacity of quercetin could also have traced back to the comparably low surface area in colon. If so, quercetin is not suitable as analytical marker for sustained release formulations with drug release over the whole intestinal tract including colon. Without the assumption of metabolism, for quercetin as marker for flavonoids, results indicate an absorption window in small intestine, mainly in jejunum. Considering human intestinal transit times for small intestinal tract, dissolution profile target of 80% drug release within 8 hours is reasonable. This target ensures reproducible release and absorption of the majority of drug substance.

Dissolution target profiles with drug release longer than 8 hours demand for hydrodynamic systems (floating systems) that float over the gastric content for a prolonged period of time [39] while drug release. The active substance is secreted this way into small intestine continuously, where absorption processes can take place. As a high concentration of excipients is needed for hydrodynamically systems, the technology is suitable just for low drug loads. For SJW with a daily dose of 500 mg Ze 117, hydrodynamically systems are considered to be not suitable. Therefore, dissolution target of sustained drug release within 8 hours is rational.

Depending on components in SJW extract exhibiting physicochemical different properties and the high drug load of 500 mg Ze 117 per daily dose, selection of a suitable sustained system is challenging. Type and amount of excipients are limited. Synchronous release of all extract components is preferable as changing the relative ratio of components may result in an alteration of pharmacological activity. Sustained release products are desired to prolong drug release without alteration of pharmacological properties. For achieving synchronous release, excipients must not interact physicochemically with any extract compound. When selecting hydrophobic matrix delivery systems, there is the possibility for low soluble lipophilic extract compounds to remain in the hydrophobic matrix excipient. This results in insufficient release of these compounds and a non-synchronous release compared to high soluble hydrophilic extract compounds. Same risk applies for SR reservoir delivery systems, where lipophilic substances may permeate insufficiently through the SR coating polymer but cumulate on the coating material. Solution of drug inside of reservoir delivery systems is key for permeation through functional coating materials. That is a requirement what is critical especially for low soluble extract compounds. Best option to achieve synchronous release is to formulate SJW extract into a two-chamber osmotic push pull system. But as these systems are high in cost of production, it would not be first choice. Another possibility provides a hydrophobic matrix delivery system. Here, drug release relies mainly on erosion of matrix excipients what ensures a complete drug release covering all extract compounds. Both systems need a certain amount of excipients what is restricted by the high drug load of SJW extract. Pharmaceutical formulation development is needed for reliable feasibility evaluation.

For development of a SJW sustained release formulation, dissolution profiles would be optimized on analytical marker to represent the whole extract. As SJW extract is classified as quantified extract according EP, focus on analytical marker is just an approximation of sustained release properties of the dosage form. In vitro testing of analytical marker provides just an assumption on how the whole extract performs in vivo. Development of a sustained release drug product of SJW extract is due to the discussed points a very challenging assignment and has to be carefully considered.

Conclusion

Development of a sustained release formulation for SJW extract is challenging due to the high drug load and the complex mixture of SJW extract covering a range of substances with different physicochemical properties and pharmacological activities. Additionally, presence of an absorption window in small intestine, mainly in jejunum was found for analytical marker. Selection of a suitable sustained release formulation is time and cost consuming. As SJW extract is classified as quantified extract according EP, in vitro testing of selected analytical marker provides just an assumption on how the whole extract performs in vivo. Suitability of SJW extract to be formulated into a SR delivery system is restricted. Further evaluation by formulation studies and clinical studies are necessary.

Acknowledgement

We gratefully thank Pia Müller, Christin Jensbach and Peter Zimmermann for the opportunity to use LC-MS/IMS equipment of Takeda GmbH in Singen (Germany) and their assistance in data acquisition. In addition, we thank Thomas Lange for his support in method set up and data interpretation. The studies were supported financially by Max Zeller Söhne AG (Romanshorn, Switzerland).

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