

UNIVERSITE LILLE NORD DE FRANCE
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**ORAL CONTROLLED DRUG DELIVERY SYSTEMS:
OPTIMIZATION OF RELEASE PATTERNS AND ELUCIDATION OF
RELEASE MECHANISMS**

**SYSTEMES ORAUX A LIBERATION CONTROLEE:
OPTIMISATION DES CINETIQUES DE LIBERATION ET ELUCIDATION
DES MECANISMES IMPLIQUES**

THESE

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Table of content

1. Introduction (English).....	1
1.1. General.....	2
1.2. Purposes of this work.....	5
1.3. Mathematical modeling of drug release and the advantages of lipid matrix formers... 6	
1.4. Application of terahertz pulsed imaging for film coating characterization (from Haaser et al., 2013).....	12
1.5. Ethanol-resistant polymeric film coatings.....	17
2.Introduction (Français).....	25
2.1 Généralité.....	26
2.2. Objectifs de la thèse.....	29
2.3. Modèle mathématique de libération de principe actif et avantages des matrices lipidiques.....	30
2.4. Imagerie terahertz pulsee pour la caracterisation de film d'enrobage.....	36
2.5. Ethanol-resistant polymeric film coatings.....	40
3. References.....	48
4. <i>In-silico</i> simulation of niacin release from lipid tablets:.....	64
Theoretical predictions and independent experiments.....	64
4.1. Materials and methods.....	65
4.1.1. Materials.....	65
4.1.2. Tablet preparation.....	65
4.1.3. Tablet characterization.....	66
4.1.4. Equilibrium solubility measurements.....	66
4.2. Results and discussion.....	67
4.2.1. Model development.....	67
4.2.2. Model fittings to experimental results.....	74

4.2.3. Deeper insight into vitamin release mechanisms	76
4.2.4. Model predictions and independent experiments	81
4.3. Conclusion	85
5. Investigation of the viscosity grade of guar gum in polymer blends to overcome ethanol sensitivity of ethylcellulose-based coated pellets.	86
5.1. Materials and Methods	87
5.1.1. Materials.....	87
5.1.2. Preparation and characterization of thin polymeric films	87
5.1.3. Pellet coating.....	88
5.1.4. Drug release measurements	88
5.1.5. SEM studies	89
5.2. Results and discussion.....	90
5.2.1. Ethylcellulose/guar gum ratio	90
5.2.2. Guar gum concentration in the total dispersion.....	95
5.2.3. Coating level.....	97
5.2.4. Storage stability.....	100
5.2.5. Guar gum viscosity	102
5.3. Conclusion	105
6. Elucidation of the underlying drug release mechanism of ethanol resistant coated pellets.....	106
6.1. Materials and methods	107
6.1.1. Materials.....	107
6.1.2. Preparation and characterization of thin polymeric films	107
6.1.3. Pellet coating.....	109
6.1.4. Drug release measurements	109
6.1.5. SEM studies	110

6.1.6 Diffusion cell studies	110
6.1.7. Determination of the drug solubility and of the partition coefficient of the drug ..	110
6.2. Results and discussion.....	111
6.2.1. Drug release from single pellets.....	111
6.2.2. Impact of the osmolality of the release medium.....	113
6.2.3. Morphology and mechanical properties of the film coatings.....	115
6.2.4. Drug mobility within the film coatings.....	119
6.2.5. Mathematical modeling of drug release.....	122
6.2.6. Intermediate ethanol concentrations	128
6. 4. Conclusion	128
7. Effects of film coating thickness on in-vitro drug release about sustained-release coated pellets: Using terahertz pulsed imaging.....	130
7.1. Materials and Methods	131
7.1 1. Materials.....	131
7.1 2. Preparation of the Pellets	131
7.1.3. Terahertz Pulsed Imaging (TPI)	132
7.1.4. Dissolution Testing	133
7.1.5. Scanning Electron Microscopy (SEM).....	134
7.2. Results and discussion.....	134
7.3. Conclusion	140
8. Summary.....	142
9. Résumé	146

1. Introduction (English)

1.1. General

Controlled drug delivery systems are frequently used in order to optimize the therapeutic effects of drug treatments. Importantly, drug levels below the minimal effective concentration at the site of action lead to failure of the medical treatment, whereas toxic drug levels can cause serious side-effects, potentially resulting in the termination of the treatment. Thus, it is of great practical importance to maintain the drug concentration within its therapeutic range, also called therapeutic window.

This can either be achieved by lowering the administered dose and shortening the time intervals between the administrations, or by using time-controlled drug delivery systems. Most often, increasing the application frequency to stay within the therapeutic range of the drug is not suitable, because of the reduced compliance of the patient, or simply for practical reasons (e.g., lack of administrations during the night). Hence, it can be highly desirable to develop controlled drug delivery systems, releasing the drug at predetermined rates to achieve optimal drug levels at the site of action. This is especially true for highly potent drugs with narrow therapeutic ranges, e.g., anticancer drugs. Comprehensive reviews of the developed strategies and investigated devices are given by Tanquary and Lacey (1974), Baker (1987), Fan and Singh (1989), and Siepmann and Siepmann (2012).

For example, matrix tablets can be used to release a drug in a pre-programmed manner. Several mechanisms such as dissolution, diffusion, erosion or osmotic effects can act to control the release rate. The determination of drug release mechanisms is important to apprehend the different ways to optimize the drug delivery system (DDS).

A major advantage of matrix systems is often the relative ease of preparation and low production cost. Another strong point is the wide range of drug release profiles that can be achieved by varying the matrix former, e.g. polymer composition: hydrophobic, hydrophilic, biodegradable, mineral or lipid matrix, or drug loading. Other industrial processes such as coating can allow achieving a large range of drug release profiles.

Coating is a process frequently used to taste masking, in the protection of the drug against exterior condition, and to ensure controlled drug delivery from pellets in example. In

a batch of coated pellets, some pellets may be in bad conformation or not coated regularly and as a consequence exhibit non desired release kinetics. So drug release profiles don't have to be in good correlation with expected kinetics. In the case of multiparticulates device only a fraction of the sub-units may presents this defect and thus, the drug release profiles of the entire population are less impacted by failed system.

Food and Drug Administration (FDA) initiated since several years, the concept of Process Analytical Technology (PAT) as a tool to analyze and control pharmaceutical process. The aims are to assure a better comprehension of fabrication and an optimization of galenic forms to obtain a high quality final product. In the case of coated multiparticulates forms, it's important to calculate the mass gain on the systems surface. Terahertz technology is accepted as new tool to determine the coating thickness on high size pellets and tablets surfaces' (Ho, 2008 and 2009).

One of the major concerns with single coated unit is "dose dumping". Upon contact with the release media, water diffuses in the device and can generate a hydrostatic pressure. Depending on the polymer properties and film coating properties and stability, this generated hydrostatic pressure can lead to crack formation, causing a failure of the system. With the use of matrix for sustained release, the content of drug can be relatively high. But, in the case of default in the coating for example, the entire drug dose can be released in a short time period. In the worst case, fatal side effects can happen. A solution is given by dividing the drug dose into multiple units. Attention has to be taken not only in product quality but also in external factor. Since several years the impact of gastric and intestinal content dependant of food intake is became a parameter non-negligible in the development of new formulations. More recently, the FDA reported the consummation of alcohol beverage in close time with DDS led to failure of the device and total drug release in short time. In these cases, dose dumping is observed for all types of drug delivery systems sensible to ethanol medium.

With respect to the major rate controlling mechanism, controlled drug delivery systems can be classified as follows:

- 1) Diffusion controlled devices
- 2) Swelling controlled devices
- 3) Erosion controlled devices
- 4) Osmotic controlled devices
- 5) Others (e.g., magnetic and/or electrostatic effects)

with diffusion, swelling and erosion being the most important features of the commercially available products. However, very often the entire process of drug release is not only determined by a single mechanism, but results from a combination of several different mechanisms.

Despite the steadily growing importance of time-controlled drug delivery systems, major challenges remain to be addressed, as explained in more detail in the following.

1.2. Purposes of this work

The major purposes of this PhD thesis were:

- (i) to decrease the ethanol resistance of polymer coated controlled release dosage forms
- (ii) to improve the characterization of controlled release polymeric film coatings
- (iii) to better understand the mechanisms controlling drug release from lipid-based matrix tablets

Various drugs niacin, theophylline and metoprolol succinate were used as highly soluble drugs

For lipid matrix: glyceryl behenate called Compritol 888 ATO.

For coating ethanol resistant: Aquacoat ECD 30: a dispersion of ethylcellulose as hydrophobic polymer and guar gum as pore former. This system was plasticized by Dibutylsebacate plasticizer.

For coating to characterize coating thickness: a dispersion of polyvinyl acetate : Kollicoat SR 30D, and PVA-PEG graft copolymer: Kollicoat IR plasticized by Triethyl Citrate.

1.3. Mathematical modeling of drug release and the advantages of lipid matrix formers

Mathematical modeling of mass transport in controlled drug delivery systems can be highly beneficial: On the one hand side, the underlying drug release mechanisms can be elucidated, on the other hand side, time-consuming and cost-intensive series of trial-and-error experiments can be replaced by rapid *in-silico* simulations (Siepmann, 2006-2012a-2013a ; Peppas, 2013). Compared to other scientific domains, such computer-assisted device design is yet rarely applied in pharmaceuticals.

A mechanistically realistic mathematical theory should always be based on a thorough physico-chemical characterization of the dosage form before and after exposure to the release medium. Based on this knowledge, appropriate model assumptions should be defined. Ideally, only the dominant mass transport phenomena should be included, whereas processes which have only a minor impact, should be neglected in order to keep the model straightforward and simple to use. Examples for mass transport phenomena, which might play a role in oral controlled drug delivery systems include water penetration into the system (Hariharan, 1993), drug particle dissolution (Siepmann, 2013b), drug diffusion through water-filled channels and/or polymeric networks (Siepmann, 2012 ; Grassi, 1999-2003 ; Frenning, 2011 ; Yin, 2011), polymer swelling, degradation and/or dissolution (Brazel, 2000 ; Borgquist, 2006 ; Sackett, 2011 ; Kaunisto, 2011), time- and position-dependent changes in the mobility of water and drug within the dosage form (Mallapragada, 1997), and system disintegration. In certain cases, only one of these phenomena might be dominant and the mathematical description of the delivery system might be straightforward (Siepmann, 2010). However, in other cases a multitude of processes might be decisive at the same time and an accurate mathematical treatment is complex (Siepmann, 2012b). Due to the large variety of controlled release dosage forms it can also not be expected that one single model could be valid for all types of systems. Instead, on a case-by-case basis, the validity of a model must be evaluated for each type of systems.

Some reports are available on lipid controlled drug delivery systems (Zaky, 2010 ; Siepmann, 2011), in which a drug is embedded within a lipid matrix former. In particular for

lipid *implants* different mathematical theories have been described so far (Siepmann, 2011 ; Guse, 2006). However, these models: (i) were either developed for relatively complex systems, in which for instance protein precipitation due to the presence of co-dissolved polyethylene glycol is of importance (Herrmann, 2007a ; Herrmann, 2007b ; Siepmann, 2008), or (ii) take only one single mass transport phenomenon into account, e.g. diffusion (Kreye, 2011a-2011b-2011c-2011d ; Gueres, 2012). Yet, there is a lack of appropriate, mechanistically realistic mathematical theories quantifying mass transport in lipid tablets and allowing for the prediction of the impact of key formulation and processing parameters, such as the initial drug loading, tablet height and tablet radius as well as the manufacturing procedure of the systems on the resulting drug release kinetics.

Lipids have recently been proposed as alternative material for the preparation of controlled drug delivery systems (Kreye et al., 2008, 2011a, 2011b; Maschke et al., 2004; Vogelhuber et al., 2003). The development of new drug delivery systems with these later can be explain by a broad spectrum of excipient type due to differences in fatty acid chain length, esterification or as lipid blends. They offer several advantages: being physiological substances, lipids show good biocompatibility, and they might be less expensive than polymeric materials. Being water-insoluble and non-swellable, lipid materials have major applications in sustained-release systems, especially for systems containing high loadings of freely water-soluble drugs. Drug release is also dependant of the diffusion coefficient and due to the degradation of the matrix induces by the lipases or an erosion of the device.

Glyceryl behenate is a lipid material, which was originally introduced as a lubricant for tablets (Jannin et al, 2003), and recently as a sustained-released excipient as matrix former (Obaidat, 2001). Several types of device are investigated with this polymer such as: tablets, suspension, beads, implants and microcapsules.

However, the production process with lipid powder is challenging due to poor flowability of the dry powder which can cause tablet size variability and potential de-mixing. Compritol 888 ATO is characterized by a low melting point, and is therefore used as a matrix former for sustained-release dosage forms (Barthelemy et al., 1999; Faham et al., 2000 ; Mirghani et al., 2000).

Impact of the technique for preparation of sustained lipophilic matrices is shown by several authors (Abd El-halim et al., 2010; Li et al., 2006). Tablets obtained by heat

treatment show a lower drug release rate than classical methods due to higher matrix tortuosity and lower porosity (Jagdale et al., 2002; Zhang et al., 2003). An advantage of the lipophilic matrix is the wide range of materials with broad spectrum of melt temperature.

Hot melt extrusion can be defined as a process to form extrudats, by forcing a powder through an orifice or die under controlled conditions: temperature, mixing... Against to classical extrusion, the use of heating leads to modification in drug release profiles with the same powder composition, because of a more intense embedding of the drug particle by the matrix former (Chokshi, 2004 ; Breitenbach, 2002 ; Repka, 2007)(figure 1).

The composition of extruder is one of the parameters which shows an impact on the quality and drug release profiles obtained. An increasing in kneading elements leads a higher torque value and can show an impact on the extrudats densities which imply a modification in drug release and quality of the product (Reitz, 2013; Vercruysse, 2012).

Other two crucial parameters are the Residence Time Distribution (RTD) and the temperature of extrusion (Verhoeven, 2008). These both affect the quality and characterization of the product. Gao et al. in 2012 elucidated the influence of the RTD on the drug and formulation properties. A high time of residence can lead a thermal degradation of the drug, but can be favorable in the case of DDS due to a slow dissolution of the drug in the blend. The impact of RTD is relevant with the involved kinetics formulation as well as the drug decomposition. This parameter is directly influenced by the screw speed and the powder feed rate.

The temperature affects the resulting drug release kinetics. An increasing in temperature increases the tortuosity and results in less porous systems and thus, decreases the release rate. So it's necessary to have a full understanding of the physical state of the drug in the extrudates formulations. A thermoanalytical technique, Differential Scanning Calorimetry (DSC), is a perfect tool to study the behavior of API or polymer to temperature variations (Qi et al., 2007) based on the T_g of amorphous samples and T_m of crystalline samples. Before HME process, several authors used hot stage microscopy to determine the HME temperature to work (determination of the temperature where the drug becomes to dissolve in the polymere, i.e.) (Agrawal et al., 2013). Several PAT are used to characterize these systems in line. Saerens and al in two differentes publications show the potential for

NIR and Raman spectroscopy to characterize polymer-drug interaction inside extrudates, and determine the solid state of the drug too (Saerens et al., 2011 ; 2012).

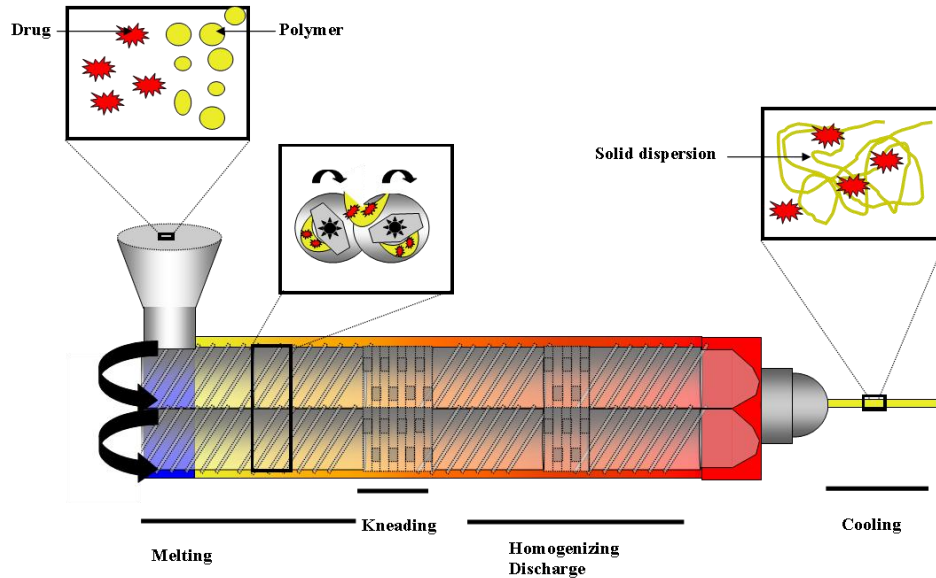


Figure 1 : Schematic hot melt extrusion process to obtain solid dispersion.

For lipid, major phenomena in drug release about matrices are the diffusion of the drug through the network (Guse, 2006). Several authors worked on different devices (Kreye, 2011a – Siepmann, 2008a). In practice drug release is not linear but becomes progressively slower with time. The drug release profiles can be illustrated as figure 1.

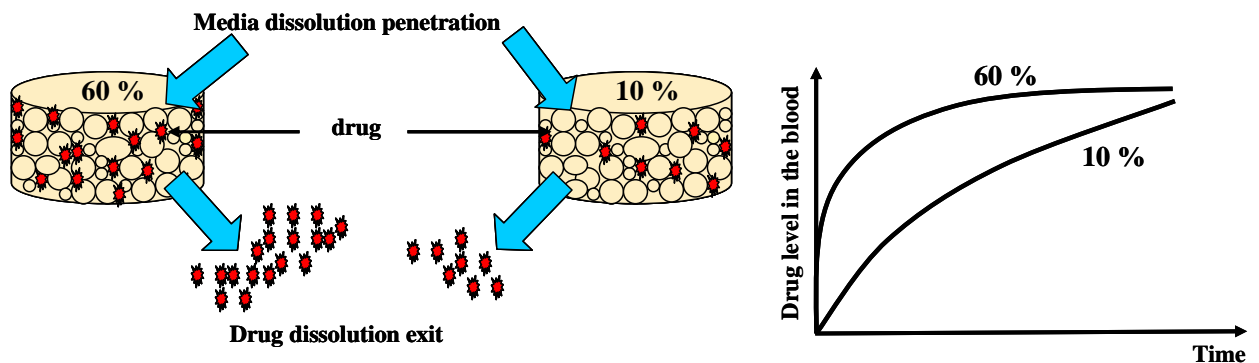


Figure 2 : Representation of matrix tablet obtained by direct compression and the impact of drug loading on drug release kinetics.

Certain authors explain drug release as follows:

- Phase 1: Relatively fast drug release obtain upon contact of matrix into medium dissolution which causes drug release from drug in surface and superficial layer.
- Phase 2: with time, drug in the superficial layer is leach out and new porous network is created. The dissolution medium diffuses through the empty porous network and dissolves the drug deeper. The dissolved drug has to travel through more tortuous network to reach the external dissolution medium. This phase characteristic by lower drug release is explained by an increasing in distance inside the matrix tablet.

The phenomenon of diffusion is based on the drug transport through polymeric network. Drug release profiles can be modified by variation in drug content or change in the size structure. The increasing in drug content increases the number of pathways by dissolution/diffusion of the drug, so higher drug release kinetics can be observed.

Certain authors use the percolation theory to explain drug release, e.g. the Leuenberger team. At low drug loading levels, clusters of drugs are completely surrounded by the polymer and the non-degradability of this latter lead to an inaccessibility of the drug to the release medium. So an incomplete drug release is observed (figure 3). With an increasing of drug content level, the connection between drug particles is more important and the dissolution of drug creates a network through which the drug can be released (Caraballo, 1996). This limit, where cluster of drug are embedded and cannot be release, is given by percolation threshold.

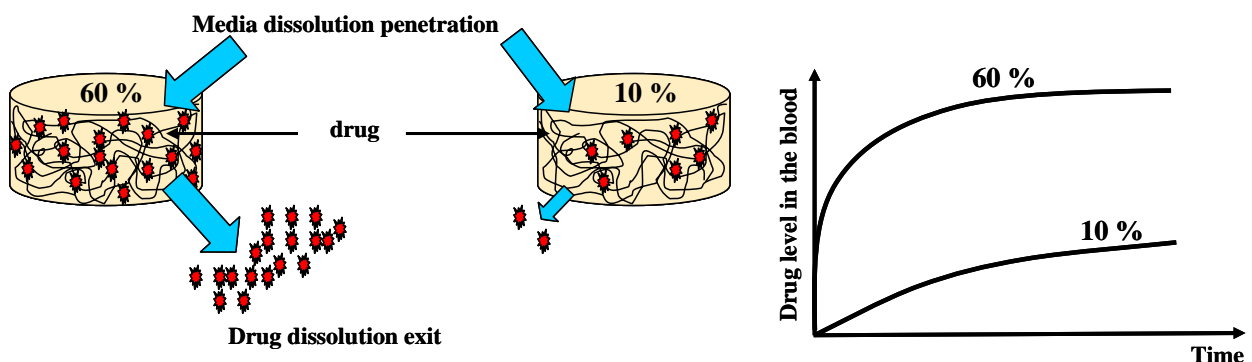


Figure 3: Representation of matrix tablet obtained by hot melt extrusion/direct compression and the impact of drug loading on drug release kinetics.

The aims of this work were to: (i) identify such a mathematical theory for glyceryl dibehenate-based tablets prepared either by direct compression or via hot-melt extrusion/grinding/compression, (ii) use this theory to better understand the relative importance of the involved mass transport processes, (iii) use this theory to quantitatively predict the impact of the tablet design (namely, of the composition, dimensions and type of preparation method) on the resulting release kinetics, and (iv) to evaluate the validity of these model predictions using several sets of independent experimental results. The vitamin niacin was used as “model drug”.

1.4. Application of terahertz pulsed imaging for film coating characterization (from Haaser et al., 2013)

Multiparticulate dosage forms are desirable drug delivery systems owing to a number of advantages over single unit dosage forms, such as better control of the gastric transit time and associated drug absorption, and a lower susceptibility to dose dumping (Bechgaard and Nielsen, 1978). Frequently, the particles are coated to modify drug release kinetics. Thus, the product performance directly correlates with critical film coating quality attributes, including the coating thickness and uniformity (Haddish-Berhane et al., 2006).

Routinely, indirect monitoring methods such as the product weight-gain and the amount of coating polymer applied are used to infer the pellet film coating thickness (and thus the coating quality) (Ringqvist et al., 2003). For complex systems, e.g. drug-layered sugar starter cores coated with a sustained-release coating, the non-specific character of the weight-gain measurements as well as the fact that coating thickness uniformity can be related to the drug layer surface morphology, render weight-gain as a sole indication of the coating quality insufficient (Ho et al., 2008; Ho et al., 2010).

First point with multiple-unit system is the elucidation of properties of each subunit separately. An idea of release characteristic can be applied by the studies of variation in kinetics between subunits. Thus, a number of studies using mechanical analysis, e.g. in vitro drug release testing, have been used to obtain more insight into pellet coating structures and their effects on drug release (Siepmann et al., 2007; Siepmann et al., 2008; Muschert et al., 2009). Those mechanistic methods provide deeper understanding of the drug release mechanism from the coated dosage form. But there is still a lack of detailed information on critical film coating quality attributes such as coating thickness, uniformity and morphology. Information on the coating thickness, uniformity and morphology may be obtained with other analytical techniques including scanning electron microscopy (SEM)(Heinicke and Schwartz, 2007), fluorescence microscopy (Andersson et al., 2000), atomic force microscopy (AFM)(Ringqvist et al., 2003), optical coherence tomography (OCT)(Zhong et al., 2011), confocal Raman microimaging (Ringqvist et al., 2003), energy dispersive X-ray imaging (EDX)(Ensslin et al., 2008), nuclear magnetic resonance spectroscopy (NMR)(Ensslin et al., 2008), electron paramagnetic resonance spectroscopy (EPR)(Ensslin et al., 2009) and

confocal laser scanning microscopy (CLSM)(Depypere et al., 2009).Cahyadi et al. in 2010 make an interesting comparison between some non-destructive techniques used to characterise the coating thickness. Direct observation in Direct Optical Microscopy seems the best technique, with SEM observations, to be the model references to validate others analytical techniques. However, both need time and money consuming, and destruction of the sample. Second technique which allows rapid and on-line characterisation is using the micrometer to measure the difference between the thickness coated and non coated pellets (Römer et al., 2008), but it's again time-consuming technique. Based on these observations Cahyadi and al., supposed Raman spectroscopy seems to be the best method for characterisation. Problems with this technique is the interpretation of signals which needs some calculations. In-line NIR (Near Infra Red) can be used to detect quantitative film coating too (Andersson et al.,2000 ; Lee et al., 2011 ; Moes et al., 2008). Cogdill et al., in 2007 compare NIR with Terahertz technology and bienque this last shows a problem of sensitivity, Terahertz technology seems to be the directe technique to characterise coating thickness. Indeed, like Raman spectroscopy, lots of noise are read in the diffractograms for NIR interpretation and a bad estimation of the results are observed. Recently, Möltgen combined NIR with science based calibration to determine noise in the diffractograms and by calculation with this value, coating thickness can be approximate on-line (Möltgen et al., 2013).

Scanning electron microscopy (SEM) is a widely used and complete technique for pellets characterization. Characterization of the surface and cross section of coated pellets give valuable information on the layer thickness, structure and the eventual presence of pores or defects. Surface pellets' observations after release in dissolution medium; allow an understanding of solvent effect and mass transport due to the leaching out of soluble polymer in this medium. Studies of pellet surface and presence of cracks at the end of release indicate a mechanical failure of the coating. To testify the acts of these mechanisms other studies have been performed such as diffusion cell studies to calculate the permeability of the coating; diffusion coefficient of the drug or determination of the partition coefficient (Marucci, 2007). Diffusion cells experiments performed under different media temperature can approach the polymer state in the polymeric membrane and explain the change in drug release due to the impact of Tg. During swelling, mass accumulation

inside the pellets can be measured by comparison in water uptake before and after drug release and microscopic measurement.

Curing, another important parameter, necessity to be characterize too. Determination of the curing end points to testify long term stability is actually based on dissolution method. In this goal, X-ray micro-computed tomography (X μ CT), Near Infra Red (NIR) (Gendre, 2011), Raman spectroscopy (De Beer, 2011) and more recently X-ray micro diffraction were employed to perform change in the microstructure during curing process (Gendre et al., 2012).

However, the applicability of some of those characterisation methods, e.g. SEM, fluorescence microscopy, atomic force microscopy and confocal Raman microimaging, is restricted to the coating surface of the sample, or the methods, e.g. SEM, fluorescence microscopy and EDX, require the samples to be cut to determine coating thickness information, , which leads to the irreversible destruction of the sample. Moreover, in NMR and EPR spectroscopy information on critical coating quality attributes can only be determined indirectly, i.e. signals are obtained during drug release testing, and CLSM needs the aid of chemometric models to evaluate coating quality characteristics.

A recently established non-destructive technique to gain deeper understanding on film coating characteristics (including coating thickness and uniformity) is terahertz pulsed imaging (TPI) (Ho et al., 2007; Ho et al., 2009; Ho et al., 2010). Terahertz radiation is part of the far infrared region of the electromagnetic spectrum (2 cm^{-1} and 120 cm^{-1}) and most of the well-established polymer formulations used in film coatings are transparent or semitransparent to the pulsed coherent light used in TPI (Zeitler et al., 2007b). Hence, the generated terahertz pulse can propagate through the sample and reflections caused by interfaces within the sample structure, due to refractive index changes, can be measured against time. Thus, single or multiple layer thicknesses (at depth) can be derived from the peak-to-peak distance in the time-domain signal (time delay of the terahertz pulse) (Fitzgerald et al., 2005) (figure 4).

Depending on the single or multiple layer thicknesses, calculation can be derived from the peak-to-peak distance in the time-domain signal (time delay of the terahertz pulse) (Fitzgerald et al., 2005). When a chemical or structural change in the device is measured, the refractive index and absorption coefficient change, leading to pulse reflections and more

signals in the terahertz waveform First change in index refraction is related to the interface between coating surface and environment media. A part of the signal is reflected and a decrease of the peak is observed during the penetration depth of the remaining terahertz pulse. The following signal correspond to a change in device nature, in this case different polymer coated, and the time delay between the both signals characterize the layer thickness between each layer composition. Coating thickness was derived directly using the time-of-flight equation:

$$2 d_{(coating)} = \Delta t c/n,$$

Where $d_{(coating)}$ is the coating thickness, Δt is the time delay between the terahertz reflections, c is the speed of light and n the refractive index of the coating matrix (Ho et al., 2007).

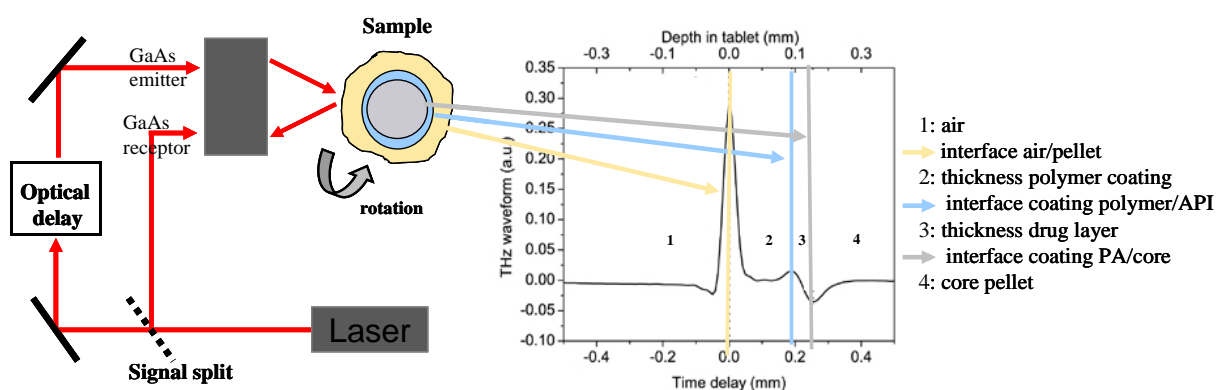


Figure 4: Schematic presentation of the terahertz pulsed imaging system to examine film coatings of solid oral dosage form. Terahertz pulse reflections occur at each interface whenever there is a change in the refractive index (Schema adapted from Shen, 2011).

Spencer et al, in 2008, correlated the TPI, coating thickness and the Mean Dissolution Time (MDT). The drug polymer layer on the pellet surface is measured and mapped into 3D model. In other study the use of TPI for sustained release of coated pellets of 6mm diameter is described. The influence of pellets preparation: starter core, drug layer, film coating and curing is analyzed. The results show a relation between size and density of layer thickness with in vitro drug release (Ho et al, 2009b). Results are confirmed by SEM observation.

This technology is a new good PAT for on-line study and quality by design (QbD). Several default types are related for coated forms; uniformity of coating, shape of coated pellets (function or not of the initial shape of the core), presence of cracks which influence drug release kinetics. An investigation of Haddish in 2006 correlate the irregular shape or default pellets with variation in drug release pattern.

For biconvex tablets, a difference in coating uniformity is observed: top and bottom surface have a coating surface more important than the tablet central domain (Ho, 2009a). The region of the smallest coating thickness is reported as an important factor influencing drug release. The consequence is an undesired large range of release pattern, and in worst case dose dumping. An index to determine crack initiation by TPI is given by Momose in 2012. This index is dependant of the density of layer coating. Coating uniformity has been shown to affect drug release behaviour from coated dosage forms (Ho et al., 2008; Ho et al., 2010). Although most analyses of film coatings using TPI have involved tablets, recently, TPI has also been shown suitable for the analysis of coating and internal drug layer thicknesses and uniformities in large sustained-release coated pellets (6 mm in diameter) (Ho et al., 2010).

Importantly, not only the average coating thickness, but also the critical coating quality attribute coating uniformity, can be accessed by TPI.

New parameter has been elucidated by Ho et al. in 2008, TEFPS (*Terahertz Electric Field Peak Strength*) which can be obtain by extraction from the obtained signal. This parameter is in correlation with physico-chemical properties of the film coating and solid dosage form under investigation (Shen and today, 2008). It is exprimed in percentage and determined from the reflected signal on the coated surface by calculation of the normalization to amplitude with the impulsion reference (Ho *et al.*, 2008).TEFPS provides information on the degree of the surface roughness and relative density of the film coating. Indeed, TEFPS was determined for each pixel and an average value over the sampled area.

In this study, TPI was employed for the first time to analyse film coating and internal drug layer thicknesses and uniformity in standard size pellets (1 mm in diameter). Furthermore, the effect of coating characteristics on the subsequent drug release behaviour of the pellets was investigated.

1.5. Ethanol-resistant polymeric film coatings

In a multiparticulate drug delivery system, the dosage of the drug is divided among several discrete delivery entities, in contrast to a classical single-unit dosage form. In case of failure of coated systems, the consequences are much more limited if only a few subunits fail compared to single unit dosage forms which are more prone to dose dumping. Multiparticulates take higher importance than single unit matrix tablets in the development of drug delivery systems, especially pellets. One advantage over single units is that gastric emptying time is less variable and higher reproducibility in GI tract, minimization of inter and intra subject variability of plasma profile (Varum, 2010-2013).

Controlled drug delivery systems generally contain higher drug doses than immediate release dosage forms, since the drug is intended to enter the human body at a pre-programmed rate over a prolonged period of time. If the release rate controlling mechanism is based on barrier functions of substances, which are insoluble in water, but soluble in ethanol, the co-consumption of alcoholic beverages can lead to “dose dumping”: “Unintended, rapid drug release in a short period of time of the entire amount or a significant fraction of the drug contained in a modified release dosage form” (Meyer and Hussain, 2005; Varum et al., *in press*). The consequences can be severe, because: (i) toxic drug concentrations might be achieved, with eventually fatal side effects, and (ii) the therapeutic effect is no more guaranteed during the envisaged time period. This is particularly true for highly potent drugs with narrow therapeutic windows, such as many opioid drugs.

Impact of food and ethanol beverages on gastric functions

Since several years, the quality of process and device show more and more development. Physiological processes are a parameter to identify relevant in vitro dissolution testing. Oral digestion should be estimated and has to be taken into account for in-vitro drug release profiles. Physiological differences in saliva secretion, gastric emptying, pH value (Singer et al., 1987 ; Van Aken et al., 2007), GIT motility and fast or fed state will be revealed in dissolution kinetics. For example, in 2006, the FDA recommends for all new orally drug products to make food effect bioavailability studies.

The bioavailability of drug from tablets is sometimes lower after administration in close time with food due the impact on small intestinal transit (Fadda, 2008). Indeed, Freire et al., (2011) show the importance of a multitude of factors which influence the gastrointestinal transit. But several studies show the importance of the tablet composition too. For immediate or extended drug release tablet, the composition has an impact on the drug release profiles obtain after administration of food. The rate of drug absorption is often slower, sometimes “dose dumping” and nothing effect in several cases (Abrahamsson et al., 2004; Schug et al., 2002). Drug absorption rate for DDS might also be regulated by the gastric emptying which influence the residence time of the DDS in the stomach especially with larger particles such as tablet. Because of their large size, tablets are also retained in the stomach where they might be ground and hydrated in a sufficient state to pass the pyloric sphincter, whereas small particles such as pellets can pass through this barrier (Meyer et al., 1981). So, an increasing time in the stomach leads to an increasing exposition to the gastric fluid which might affect the dissolution time. A study about the absorption of caffeine from enteric coated pellets in the intestine shows a correlation with the time delay in drug release and the gastric emptying rate. An increasing concentration due to retention in the stomach gastric can lead to a dose dumping of the drug (Weitschies et al., 2005).

It is well known that ethanol and alcohol beverages have several impacts on gastric functions and the major might be the inhibition of gastric emptying (Franke et al., 2004-2005). Park and al, in several studies, make a distinction between ethanol/alcoholic beverages composition and impact on gastric emptying. First study in 2004 has shown that the gastric emptying rate of pure ethanol solution is significantly slower than that of water. In 2005, they have shown for different types of alcohol beverages (beer or red wine, based on different ethanol and caloric content) diverging effects on gastric emptying of solid meals. These effects can't be assigned on the difference in ethanol or caloric content, but suggest the role of other non-alcoholic ingredients. Lennernas et al., (2009) go deeper in the observation: the inhibition of gastric emptying can be related to the caloric content of the beverage, but it's independent of the caloric content of meal consumed in the same time. Another impact of ethanol on biological function is the increasing of permeability of the intestinal mucosa (Lavo et al., 1992).

Impact of ethanol beverages on drug release devices

Indeed, dose dumping may be due to the delay of gastric emptying but also in the case of an accidental or intentional change in the release rate of the device. Many formulations for multiparticulates devices are based on polymeric coating. However, major polymers are soluble in organic solvents like ethanol. So an exposition of the device to high ethanol concentration leads to a premature dissolution of the polymeric coating (Weathermon et al., 1999).. A combination of higher permeability of the drug and higher dissolution of devices upon ethanol contact result in a higher C_{max} and in the worst case dose dumping. Because the resulting side effects caused by dose dumping of opioid drug, the marketing of the Palladone was suspended (FDA Alert, July 2005).

One attempt to reduce the risk of dose dumping is to add warning labels on the drug product. However, it has been reported that heavy drinkers suffering from chronic low back pain did not reduce their opiate use, despite such warnings about concomitant use of alcohol and opiates (Booker et al., 2003). Furthermore, the Behavioral Risk Factor Surveillance System reported that 1 of 3 drinkers in the U.S. are “binge drinking”: They consume 4/5 drinks (women/men) in a short period of time (Serdula et al., 2004). Thus, controlled drug delivery systems might be exposed to high ethanol concentrations in the stomach in practice. In 2009, Lennernaes published a very interesting review article on ethanol vulnerable formulations, pointing out the risk of dosage form failure due to the co-consumption of ethanol. Both, reservoir devices as well as matrix systems might be affected. The oral controlled drug release product “Palladone” is an example for a system, which exhibits a risk of dose dumping when alcoholic beverages are co-administered. The drug is hydromorphone HCl in this case, the release of which is controlled from pellets comprising ethylcellulose, ammonia methacrylate copolymer type B and stearyl alcohol (Fadda et al., 2008). The pellets are administered once daily in capsules. A clinical trial with healthy subjects revealed that the co-administration of 240 mL of 40 % alcohol together with a 12-mg Palladone capsule resulted in an average peak hydromorphone concentration, which was about 6 times greater than when the capsules were taken with water (HMP1013). One subject even showed a 16-fold increase in the maximal plasma concentration. As this opioid drug is highly potent and its side effects are severe, the manufacturers decided to suspend

marketing this product. The ethanol sensitivity of Palladone SR capsules was also demonstrated *in vitro* (Walden et al., 2007).

Fadda et al. (2008) compared 3 commercially available controlled drug delivery products containing 5-aminosalicylic acid: Pentasa, Asacol, Salofalk. They showed that the addition of up to 40 % ethanol to the release medium significantly affected the resulting drug release profiles. Interestingly, the observed changes strongly depended on the type of formulation. In 2010, Smith et al. reported on *in vitro* studies, which were conducted with 27 oral modified release products. Different types of drugs were considered, including opioids, calcium channel blockers, antidepressants, and antiarrhythmics. Importantly, 9 of 10 capsule formulations and 2 of 17 tablet formulations showed accelerated drug release in media containing 40 % ethanol. Also, Traynor et al. (2008) studied the impact of the addition of high amounts of ethanol to the release medium on drug release. They showed that the release rate of tramadol from the 24 h controlled release formulation "T-long" significantly increased. In the case of hydroxypropyl methylcellulose (HPMC) matrix formulations, Levina et al. (2007) reported only moderate effects of ethanol on drug release *in vitro*. In contrast, Roberts et al. (2007) showed a significantly increased aspirin release rate from HPMC matrix tablets in the presence of high ethanol concentrations. This might at least partially be explained by slower tablet swelling and increased drug solubility. Furthermore, Roth et al. (2009) reported that verapamil release from controlled release matrix systems was significantly faster upon addition of 40 % ethanol to the release medium. Another very interesting study on the potential impact of ethanol in the release medium on the performance of polymeric controlled drug delivery systems was published by Larsson et al. (2010). They prepared free films based on ethylcellulose and hydroxypropyl cellulose (HPC) by spraying organic polymer solutions and studied the water permeability of the systems in the presence and absence of ethanol. Importantly, the water permeability of films with *low* HPC contents increased with increasing ethanol concentration, whereas the water permeability of films with *high* HPC contents decreased with increasing ethanol concentration. The increase in water permeability at low HPC contents is likely due to more pronounced ethylcellulose swelling in the presence of ethanol. The decrease in water permeability at high HPC contents was explained as follows: In these systems a continuous HPC network is formed, through which water can be relatively rapidly transported. Adding

ethanol to the release medium leads to more important ethylcellulose swelling, which probably at least partially closes the “HPC pores”. These various *in vitro* and *in vivo* examples clearly demonstrate that the consumption of alcoholic beverages can be very dangerous for patients treated with highly potent drugs in the form of oral controlled delivery systems. There is a clear need to reduce this risk.

Blend coating polymer to ethanol-insensitivity of sustained oral devices

Recently, a novel type of polymeric film coatings has been proposed providing ethanol-resistant drug release from coated dosage forms: blends of ethylcellulose and guar gum (Rosiaux et al., 2013). The use of polymer blends can indeed offer major benefits compared to the use of single polymers (Lecomte et al., 2003, 2004a,b, 2005a,b; Siepmann et al., 2007a,b). In the present case, the presence of ethanol insoluble guar gum avoids the undesired dissolution of ethylcellulose in ethanol-rich media. Vice-versa, the presence of water insoluble ethylcellulose avoids the undesired dissolution of guar gum in water.

Ethylcellulose is a hydrophobic coating material often used for controlled release, taste masking and moisture protection applications. It is generally used due to its non toxic, non allergenic and non irritant properties. Ethylcellulose can be applied from organic solutions but to avoid the use of solvent like methanol, dichloromethane or ethanol for example, aqueous dispersion of ethylcellulose have been marketed. Due to the fact ethylcellulose is soluble in ethanol, this one make a good candidates to develop a new DDS resistant to alcohol beverages.

Polysaccharide based formulations are good candidates for coating DDS. Polysaccharides might be not expensive and show a broad range in structure and a large variety of properties. (Hovgaard and Brondsted, 1996). Moreover, they might be highly stable, safe, non-toxic, and biodegradable and are hydrophilic. Due to this hydrophilic property, polysaccharides can often form a gel by swelling upon exposure to the dissolution medium and might become highly soluble, leading to a rapid drug delivery (Milojevic et al., 1996). To control this solubility characteristic and limit drug release, hydrophobic polymers can be added. A family of polysaccharides naturally occurring in plant: comprising guar gum show good properties in coated technologies. It might be used like a pore former, for colon targeting (Krishnaiah et al, 1998; Prabakaran, 2011), and in our case its insolubility in ethanol makes it a good candidate to achieve ethanol insensitive coatings.

For this thesis, pellets have been coated by application of a polymeric formulation where particles are fluidized and sprayed onto the pellets surfaces. Coating process can be explain to have better aspect, taste masking, protection against environmental conditions and particularly suitable for controlled release. Application of a film composed by insoluble agent, the ethylcellulose dispersion: Aquacoat, is necessary to create a barrier which holds the drug modifying drug release. The used of Wurster apparatus where the spray nozzle is fitted in the base plate allow the spray pattern inconcurrent with the air feed. Therefore, the use of Wurster cylinder and a base plate with different perforations allows acceleration by the air flow of pellets inside the tube (figure 5). Beads pass through the spray cone to be coated by solution/suspension; and due the fact that the nozzle is in immersion within the air flow (in center of Wurster cylinder), the coating solution travels only short distance before striking the pellet surface's (Wheatley et al., 1997). Important process parameters for application of aqueous polymer formulation with Wurster coating process have impact on coating steps such as the solid content of the dispersion (Wesseling et al., 1999 ; Pearnchob et al., 2003), the spray rate (Larsen, 2003), the nozzle diameter and atomization air pressure (Ronsse et al., 2008) and the product temperature (Bodmeier, 1997).

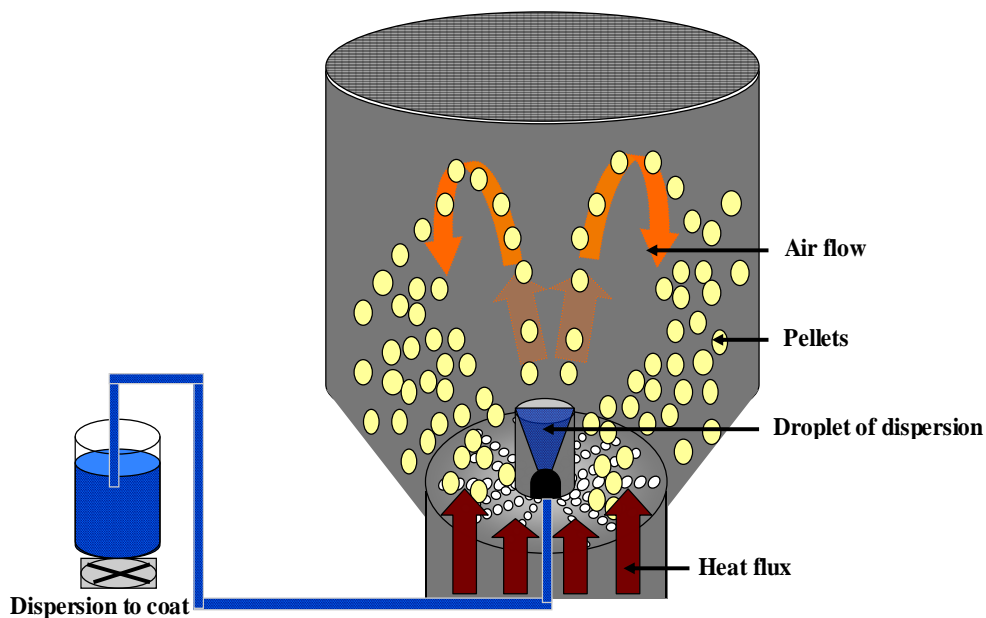


Figure 5: Schematic representation of Wurster apparatus.

To keep a good conformation and activity of the device, drug release profiles (time and concentration) should not be influenced by intrinsic or extrinsic factors. So, a challenge after bioavailability studies of the drug upon meal or alcohol substance is to have an insensitive device. Blends of polymers with different properties are interesting substances to coat solid dosage forms. A variation between polymers blend ratios allow a coating with broad physico-chemical properties and controlled drug release (Lecomte et al, 2003; Siepmann et al, 2008d).

Polymer blends have been used to have specific application in several conditions:

- pH resistance (Siepmann et al., 2008b, 2008d) ;
- colon targeting (Karrout et al., 2009) ;
- improve the storage stability (Siepmann et al., 2008c).

In this work, coated pellets have been studied as ethanol resistant drug delivery systems. Interestingly, theophylline release from pellets coated with blends of the aqueous ethylcellulose dispersion Aquacoat® ECD 30 and guar gum was shown to be virtually unaffected by the addition of 40 % ethanol to the release medium.

However, so far it is unclear what the crucial formulation parameters of these novel ethylcellulose:guar gum film coatings are. The aim of this work was to identify them. Specifically, the minimum amount of guar gum to be added in order to avoid ethylcellulose dissolution in ethanol-rich media was to be determined as well as the critical guar gum viscosity, required to provide ethanol-resistance. Also, the impact of the degree of coating dispersion dilution on the system's performance was to be studied.

Furthermore, different types of mass transport mechanisms can be involved in the control of drug release from coated dosage forms (Lecomte et al., 2005a,b; Siepmann et al., 2005, 2006), such as water penetration into the system, polymer swelling, drug dissolution, drug diffusion through the polymeric network and/or through water-filled pores/cracks, polymer dissolution, limited drug solubility and so on, to mention just a few (Marucci et al., 2011; Kaunisto et al., 2011). Appropriate mathematical equations can be used to quantitatively describe these physico-chemical phenomena (Muschert et al., 2009a,b; Siepmann and Siepmann, *in press*). Fitting such models to sets of experimentally determined drug release kinetics and other key features of the dosage form (e.g. water uptake behavior

and changes in the mechanical strength upon exposure to aqueous media) can allow determination of system-specific parameters, such as the apparent diffusion coefficient of the drug within the film coating. Knowing these values, the importance of the involved mass transport phenomena in the respective type of drug delivery system can be estimated and the dominant release mechanism can be identified. In addition, mechanistically realistic mathematical theories allow for the quantitative prediction of the effects of the device design (e.g. composition, geometry and dimensions) on the resulting drug release kinetics (Siepmann and Peppas, 2001; Siepmann and Goepferich, 2001; Borgquist et al., 2004; Marucci et al., 2008).

The aim of this PhD thesis was also to elucidate the mass transport mechanisms controlling drug release from pellets coated with the novel ethanol-resistant polymeric film coatings. Being multiple unit dosage forms, pellets provide an additional advantage for this type of applications: In case accidental film damage and dose dumping would occur in a pellet, the overall release from the ensemble of pellets would hardly be affected. Based on a comprehensive experimental characterization of the devices before and after exposure to different release media, appropriate mathematical equations were to be identified and a mathematical model to be developed allowing for facilitated drug product optimization: *In silico* simulations should be able to predict the effects of formulation parameters on the resulting drug release kinetics.

2.Introduction (Français)

2.1 Généralité

Les systèmes à libération contrôlée de principes actifs sont fréquemment utilisés pour optimiser les effets thérapeutiques lors de traitements médicamenteux. Une concentration de principe actif inférieure à la concentration minimale effective montre un défaut du traitement au niveau du site d'action, tandis qu'un niveau trop élevé dépassant le seuil de toxicité pourra occasionner de sérieux effets secondaires néfastes, pouvant entraîner l'arrêt du traitement. Ainsi, il est d'un grand intérêt de maintenir une concentration en principe actif à l'intérieur de l'intervalle thérapeutique, également appelée fenêtre thérapeutique.

Pour une délivrance sur une longue période, la concentration efficace doit donc être maintenue par administrations répétées de doses précises, à intervalles de temps régulier. Néanmoins, ces administrations successives comme moyen de maintien d'une concentration efficace ne sont pas une solution. A long terme, un problème d'observance chez le patient peut apparaître ou encore un non-suivi du traitement thérapeutique (lors d'administrations de nuit par exemple). L'objectif est donc de trouver pour chaque molécule et chaque pathologie, la forme galénique la mieux adaptée, par l'étude des différentes formes galéniques, de l'état physique du principe actif, des différents excipients et des procédés de fabrication. Ainsi, le dispositif obtenu sera au maximum de son utilité comme traitement avec une délivrance optimale de la quantité de principe actif sur le lieu d'action, mais aussi un traitement optimal pour le patient (Nicoli et Colombo, 2001). Cela est encore plus vrai pour des molécules hautement actives avec une fenêtre thérapeutique restreintes comme c'est le cas des molécules anti-cancéreuses. De nombreuses revues sur le développement des stratégies et nouveaux systèmes à libérations sont données par Tanquary and Lacey (1974), Baker (1987), Fan and Singh (1989), and Siepmann and Siepmann (2012).

Les systèmes matriciels permettent une délivrance du principe actif de manière continue. Plusieurs mécanismes comme la dissolution, l'érosion ou les effets osmotiques entrent en ligne de compte pour contrôler les taux de libération. Il est d'une grande importance de déterminer les mécanismes intervenant dans la libération du principe actif afin d'appréhender les différents moyens à mettre en œuvre pour optimiser le Système de Délivrance du Médicament (SDM).

L'avantage principal des systèmes matriciels est lié à leur facilité de préparation ainsi que leurs faibles coûts de production. Un autre point fort est lié à la haute variabilité des cinétiques de libération qui peuvent être observées par variation de la composition polymérique : hydrophobe, hydrophile, biodégradable, minéral ou lipidique, mais encore selon le taux de chargement initial. Des processus industriels comme l'enrobage permettent d'élargir encore l'éventail des profils obtenus.

L'enrobage est une technique fréquemment utilisée dans le masquage de goût, dans la protection du principe actif contre les conditions extérieures mais aussi pour permettre à des systèmes d'avoir des cinétiques de libération prolongée. Dans le cas des systèmes multiparticulaires, au sein d'un même lot de granules certaines unités peuvent être de mauvaises conformations (asymétrie du noyau par exemple) ou enrobées de manière irrégulière. Les profils de libération obtenus ne devraient pas, alors, être en adéquation avec ceux escomptés. Le fait d'avoir repartitionné le principe actif sur plusieurs sous-unités permet d'avoir non seulement une seule fraction qui se retrouve défectueuse, mais aussi un profil de libération de la population entière qui se retrouve moins impacté par la défaillance du système.

Le choix de la forme galénique est donc dépendante des profils de libération souhaités mais aussi des risques encourus lors d'une défaillance possible du système. Ainsi, on comprend l'importance que porte la FDA (Food and Drug Administration) dans la démarche PAT (Process Analytical Technology). Cette approche a pour but d'avoir une meilleure compréhension des procédés de fabrication et une optimisation des formes galéniques afin d'assurer un produit final de haute qualité. Cette démarche se définit par l'utilisation d'outils analytiques permettant un contrôle du produit. Le développement de nouvelles technologies pour assurer le contrôle au cœur même de la manipulation et en temps réel est donc primordial. Par exemple, la technologie terahertz a été acceptée comme nouvel outil dans la détermination de l'épaisseur pour des granules de grandes tailles (Ho, 2008 et 2009).

Depuis plusieurs années des études de plus en plus poussées ont montrées les fluctuations qu'occasionne le contenu gastrique et intestinal sur les profils de libération des systèmes médicamenteux. Plus récemment la FDA a reporté les dramatiques conséquences de l'absorption concomitante d'alcool et de systèmes réservoir à libération contrôlée.

L'emploi des formes à libération prolongée implique un contenu en principe actif relativement plus important que dans les formes conventionnelles. En cas de survenue d'un défaut, la totalité de la molécule active se retrouve disséminée sur une courte période. C'est le phénomène de « dose dumping ». Ce pic de concentration est à l'origine d'effets secondaires néfastes, voire létaux dans le pire des scénarios. L'utilisation des formes multiparticulaires peut sembler une solution par la division de la quantité initiale de principe actif en plusieurs fractions et non plus en une seule, évitant ainsi une libération totale et massive du contenu. Toutefois dans le cas d'intervention d'une source externe ayant un impact négatif, comme par exemple la présence d'éthanol, la division en plusieurs sous particules de la molécule active ne peut être efficace, chaque fraction étant susceptible de subir le phénomène de « dose dumping ».

Selon les mécanismes de libération contrôlant la fuite du principe actif, les systèmes à libération contrôlés peuvent être classifiés comme tel :

- 1) Systèmes contrôlés par diffusion
- 2) Systèmes contrôlés par gonflement
- 3) Systèmes contrôlés par érosion
- 4) Systèmes contrôlés osmotiquement
- 5) Autres (i.e., effets magnétiques et/ou électrostatiques).

Les phénomènes de diffusion, de gonflement ou d'érosion sont les mécanismes les plus retrouvés au niveau des produits commercialisés. Toutefois, il est rare lors de l'élucidation des mécanismes impliqués durant les phénomènes de libération de n'avoir qu'un seul mécanisme. La majorité des cas impliquent, en effet, une combinaison de plusieurs mécanismes.

Bien que l'importance accordée aux systèmes à libération contrôlée est croissante, le challenge reste néanmoins d'actualité comme expliqué plus en détail dans les paragraphes suivants.

2.2. Objectifs de la thèse

Les buts majeurs de cette thèse ont été:

- (i) De mieux comprendre les mécanismes impliqués dans des comprimés matriciels à base de lipides.
- (ii) La diminution de l'éthanol résistance des formes à libération contrôlée enrobés par des films polymériques
- (iii) D'augmenter la caractérisation de ces systèmes polymériques d'enrobages.

2.3. Modèle mathématique de libération de principe actif et avantages des matrices lipidiques

Les modèles mathématiques impliqués dans les transports de masses pour les systèmes à libération contrôlée sont hautement bénéfiques: D'un côté les mécanismes de libération peuvent être élucidés, et d'un autre côté la perte de temps et d'argent lié à la mise en place des séries expérimentales pour arriver au but peuvent être remplacés par des simulations *in-silico* rapides (Siepmann, 2006-2012a-2013a ; Peppas, 2013). Comparé aux autres domaines scientifiques, ces modèles sont rarement appliqués dans le domaine pharmaceutique.

Un modèle mathématique mécanistique et réaliste doit toujours être basé sur des caractérisations physico-chimiques de la forme galénique avant et après exposition au milieu de dissolution. Basés sur ces connaissances, des prédictions de modèles mathématiques peuvent être définis. Idéalement, seulement les phénomènes dominant durant le transport de masse devront être inclus, tandis que les procédés qui ont un impact mineur sur les phénomènes de libération devront être négligés dans l'optique de garder un modèle direct et simple d'utilisation. Plusieurs phénomènes pour le transport de masse, jouant un rôle crucial dans les formes orales à libération contrôlée inclus la pénétration de l'eau dans le système (Hariharan, 1993), la dissolution des particules actives (Siepmann, 2013b), la diffusion des particules actives via les canaux occupés par l'eau et/ou les réseaux polymériques (Siepmann, 2012 ; Grassi, 1999-2003 ; Frenning, 2011 ; Yin, 2011), les phénomènes de gonflement, dégradation et/ou dissolution impactant le polymère (Brazel, 2000 ; Borgquist, 2006 ; Sackett, 2011 ; Kaunisto, 2011), les changements de mobilité des molécules d'eau et de principe actif au sein du système, en fonction du temps et de la localisation (Mallapragada, 1997), et la désintégration du système. Dans certains cas seulement un de ces phénomènes va être dominant et la description d'un modèle mathématique sera directe et simple (Siepmann, 2010). Toutefois, dans d'autres cas une multitude de procédés seront décisifs dans le même temps et le traitement mathématique s'en trouvera beaucoup plus complexe (Siepmann, 2012b). Due à la large variété des formes à libération contrôlée, il est inutile d'espérer d'avoir un seul modèle mathématique qui serait valide pour l'ensemble des systèmes. Ainsi, sur la base du cas par cas, la validité d'un modèle devra être évaluée pour chaque type de systèmes.

Les lipides offrent une alternative aux polymères hydrophobiques et dégradables dont les produits de dégradation entraîne une série d'effets secondaires néfastes pour la préparation de système à visée contrôlée (Kreye et al., 2008, 2011a, 2011b; Maschke et al., 2004; Vogelhuber et al., 2003). Le développement de nouvelles formes galéniques à partir des lipides est expliqué par le large spectre des types d'excipients qui peuvent être obtenus par variation de la longueur de la chaîne d'acides gras, par des procédés d'estérifications qui peuvent y être rattachés ou encore par l'utilisation d'un mélange de lipides induisant de nouvelles cinétiques de libération. La libération à partir de ce type de matrice est gouvernée par le type d'acide gras composant la matrice ainsi que par la composition enzymatique des liquides digestifs. Les différents types de lipides retrouvés dans la littérature sont les acides et alcools gras initialement utilisés comme produits de lubrification. Leurs avantages sont multiples : en plus d'être des substances physiologiques, ils montrent une bonne biocompatibilité et sont moins chers que les excipients polymériques classiques. Du fait de leur insolubilité dans l'eau, de leur caractères non gonflant, les lipides sont largement applicables pour des systèmes à libération prolongée, cela est d'autant plus vrai pour les systèmes à haut chargement en principes actifs hautement solubles. La libération sera également dépendante des coefficients de diffusion de la molécule active par rapport à la matrice et de la dégradation de cette matrice induite par les lipases ou par l'érosion.

Le glyceryl behenate est un matériel lipidique qui a été introduit pour la première fois en tant que lubrifiant pour comprimés (Jannin et al, 2003), et plus récemment comme excipient à libération prolongée (Obaidat, 2001). Plusieurs types de systèmes sont alors obtenus comme par exemple les comprimés, les suspensions, les granules, les implants ou encore les microcapsules.

Toutefois, les procédés de fabrication avec les lipides sous forme de poudre est un véritable challenge au vue de leur faible fluidité qui peut causer des variations de taille et potentiellement de dé-mélange. Compritol 888 ATO est caractérisé par un faible point de fusion (74 °C) d'où son utilisation autant en tant que poudre que par des procédés thermiques (Barthelemy et al., 1999; Faham et al., 2000 ; Mirghani et al., 2000).

L'impact de la technique de préparation des matrices lipophiles à libération prolongée est montré par plusieurs auteurs (Abd El-halim et al., 2010; Li et al., 2006). Les comprimés obtenus par un traitement thermique montrent un faible taux de libération

comparé à ceux issus de méthodes classiques, ceci étant lié à l'augmentation de la tortuosité matricielle et la faible porosité (Jagdale et al., 2002; Zhang et al., 2003). Un avantage pour les formes matricielles est, comme dit plus haut, la haute variété de matériels montrant un large spectre au niveau des températures de fusion. Une technique thermodynamique, la Calorimétrie différentielle à balayage (DSC), est un outil parfait dans l'étude du comportement du principe actif ou du polymère aux variations de température.

L'extrusion en phase chauffante (HME) peut être définie comme un procédé pour former des extrudats, en forçant une poudre à passer par un orifice qui lui donnera sa forme, et cela sous différentes conditions : température, mélange... L'avantage des systèmes double vis est grand, il permet un meilleur mélange mais surtout un emprisonnement plus intense des particules actives au sein d'une matrice. Un extrudeur est typiquement constitué d'un fourreau renfermant des vis, une unité contrôlant la rotation des vis, un système de chauffage/refroidissement et un orifice de sortie (figure 6). De la zone de remplissage à la sortie, 4 étapes sont décrites : le remplissage ; la fusion et plastification ; le transport et mélange ; la mise en forme.

La composition au sein de l'extrudeur est un paramètre clé pour la formulation. Une augmentation du nombre d'éléments de cisaillement entraîne une augmentation de la densité des extrudats, impactant alors la qualité du produit et les cinétiques de libération (Vercruyse, 2012). Deux autres paramètres cruciaux sont la température et le temps de résidence de la poudre au sein de l'appareillage (Verhoeven, 2008). Il est montré qu'une augmentation de la température augmente la tortuosité, donnant un système moins poreux, donc une diminution des cinétiques de libération. L'avantage des polymères lipidiques contrairement aux autres excipients est de montrer une faible température de fusion, appréciable pour des manipulations d'extrusion en phase chauffante à faible température (principe actif thermosensible).

Une exposition trop longue d'un principe actif thermosensible entrainera la dégradation de ce dernier. Il est important d'effectuer un temps de passage rapide pour éviter la décomposition du principe actif, mais suffisamment long pour permettre un bon mélange des produits ainsi qu'une dissolution du principe actif dans le polymère pour permettre la rétention du principe actif. Cette valeur de temps de résidence est directement liée à la vitesse de rotation des vis, ainsi qu'au taux d'alimentation.

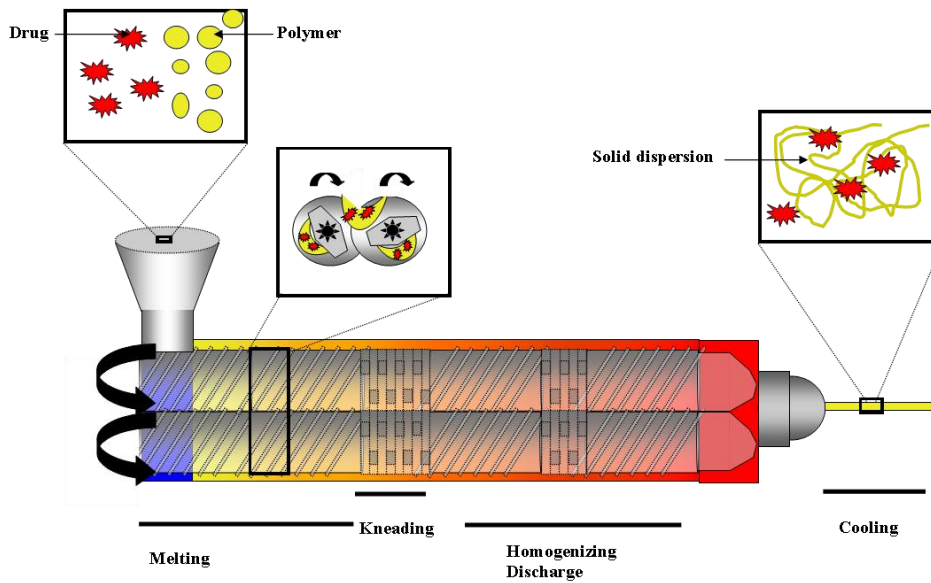


Figure 6 : Schéma du procédé d'extrusion en phase chauffante.

Pour les lipides, le phénomène majeur dans la libération du principe actif est la diffusion via un réseau poreux (Guse, 2006). Plusieurs auteurs ont travaillé sur ces systèmes (Kreye, 2011a – Siepmann, 2008a). En pratique, il est observé une libération non linéaire de la molécule, qui devient progressivement plus lente avec le temps. Les profils de libération sont illustrés comme ci-dessous.

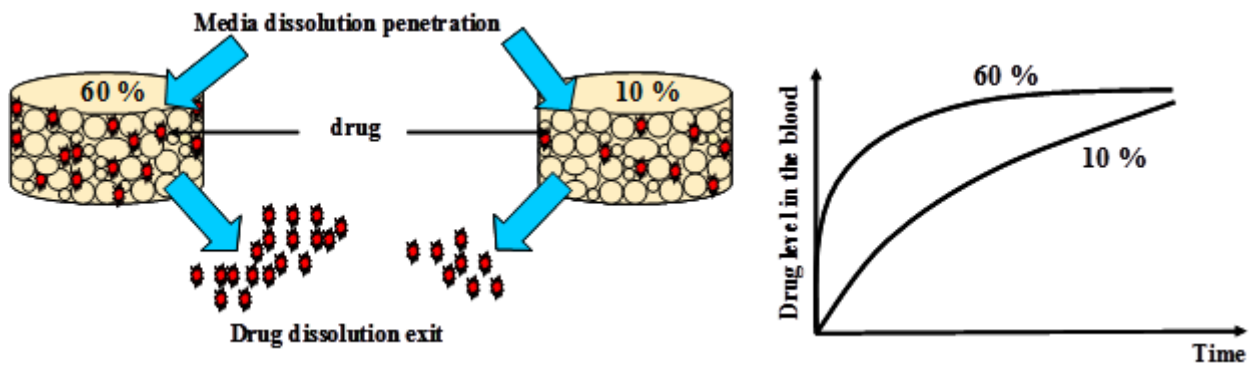


Figure 7 : Représentation de comprimés matriciels obtenus par compression directe et impact du taux de chargement sur les cinétiques de libération.

Certains auteurs expliquent la libération comme telle:

- Phase 1: Une relative rapide libération de PA obtenue après contact de la matrice au milieu de dissolution qui cause une fuite du PA de la surface et des couches superficielles.
- Phase 2: Avec le temps, le PA des couches superficielles est libéré et un nouveau réseau poreux est créé. Le milieu de dissolution diffuse alors via le réseau poreux et dissout le PA plus profondément. Le PA dissout va alors voyager au sein du même réseau poreux vide et ainsi être libéré au sein du milieu de dissolution externe. Cette phase caractérisée par une faible cinétique de libération est expliquée par une augmentation de la distance au sein du comprimé.

Ce phénomène de libération est expliqué par la théorie de la percolation. Ainsi une variation du contenu en PA ou un changement de la taille du système induira un changement des profils de libération par modification du nombre de voies de diffusion. Une plus forte teneur en PA entrainera dès lors une augmentation des voies de diffusion par dissolution/diffusion du PA, et donc les cinétiques de libération seront plus rapides.

Certains auteurs utilisent la théorie de percolation pour expliquer les cinétiques de libération, c'est notamment le cas de l'équipe de Leuenberger. Un raté dans la théorie de percolation est observé dans le cas des dispersions solides où la molécule active est emprisonnée au sein d'un polymère insoluble dans l'eau, mais peut être expliqué. A un faible taux de chargement, des groupes de PA sont complètement entourés par le polymère et la non dégradation de ce dernier entraîne une inaccessibilité du PA au milieu de dissolution. Une libération incomplète est alors observée (figure 2.4). Une augmentation du contenu en PA occasionne une connexion plus importante entre les particules de PA, créant un réseau dans lequel le PA peut être libéré (Caraballo, 1996). Cette limite où le groupement de PA est emprisonné et ne peut être libéré est nommé "percolation threshold".

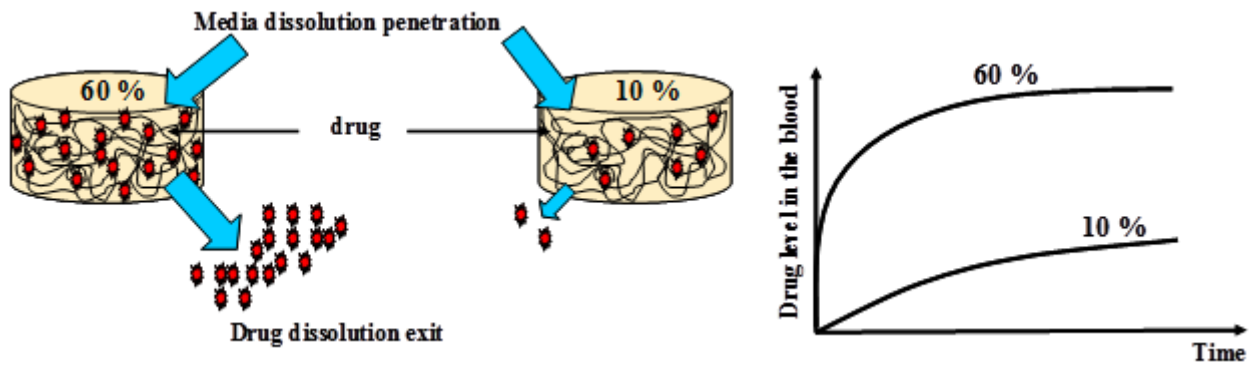


Figure 8: Représentation de comprimés matriciels obtenus par extrusion en phase chauffante/broyage/compression directe et l'impact du taux de chargement initial sur les cinétiques de libération.

Les buts de ce travail ont été de :

- (i) Identifier une théorie concernant un modèle mathématique pour des comprimés de glyceryl dibehante préparés par compression directe ou par une série d'extrusion en phase chauffante/broyage/compression directe,
- (ii) Utiliser cette théorie pour mieux comprendre l'importance des procédés impliqués dans le phénomène de transport de masse,
- (iii) Utiliser cette théorie pour prédire quantitativement l'impact de la forme du comprimé (notamment la composition, la dimension et la méthode de préparation) sur les cinétiques de libération résultantes,
- (iv) Evaluer la validité de ce modèle sur différents lots expérimentaux. La niacine, vitamine B3, est utilisée comme PA modèle.

2.4. Imagerie terahertz pulsée pour la caractérisation de film d'enrobage

(from Haaser et al., 2013)

Les formes multiparticulaires montrent une multitude d'avantages par rapport aux formes uniques, comme un meilleur contrôle du temps de transit gastrique et une faible susceptibilité au phénomène de dose dumping (Bechgaard and Nielsen, 1978). Fréquemment les minigranules sont enrobées pour modifier les cinétiques de libération. Alors, les performances du produit seront directement corrélées avec les attributs de qualité du film d'enrobage, incluant l'épaisseur et l'uniformité (Haddish-Berhane et al., 2006).

La détermination des modèles mathématiques est liée à la connaissance de chacune des phases impliquées dans le profil de libération (Ringqvist et al., 2003). La technique la plus ancienne mais néanmoins toujours d'actualité reste le calcul de la prise en masse par mesure de la quantité de produits pulvérisés dans le cas des formes enrobées. Toutefois cette technique, qui ne permet qu'une approximation de la masse gagnée liée à la perte de produits durant la technique d'enrobage, semble de plus en plus désuète (Ho et al., 2008; Ho et al., 2010).

Dans le cas d'études de libérations à partir de sous unités, il est préférable d'élucider les propriétés des sous unités séparément afin d'étudier les variabilités entre chaque individus. Un grand nombre d'études utilisant des analyses mécaniques, par exemple des tests de libération in-vitro, sont utilisées pour obtenir plus d'informations sur les structures des films d'enrobages appliqués sur les mini granules et leurs effets sur les cinétiques de libération (Siepmann et al., 2007; Siepmann et al., 2008; Muschert et al., 2009). Ces méthodes permettent alors de comprendre plus profondément les mécanismes de libération à partir de formes enrobées, mais ne sont pas forcément utiles pour donner en détails des informations sur la qualité des films attribuée à l'épaisseur, l'uniformité et la morphologie du film. Ces informations peuvent être obtenues par d'autres techniques analytiques comme la Microscopie Electronique à balayage (MEB)(Heinicke and Schwartz, 2007), microscopie à fluorescence (Andersson et al., 2000), microscopie à force atomique (AFM)(Ringqvist et al., 2003), Imagerie Raman (Ringqvist et al., 2003), imagerie rayon X à énergie dispersive

(EDX)(Ensslin et al., 2008), spectroscopie à résonance magnétique nucléaire (NMR)(Ensslin et al., 2008), spectroscopie à résonance paramagnétique nucléaire (EPR)(Ensslin et al., 2009) et la microscopie à laser confocal à balayage (CLSM)(Depypere et al., 2009).

Toutefois, la plupart de ces méthodes de sont restreintes à l'enrobage de surface, ou encore certaines méthodes comme la MEB, la microscopie à fluorescence ou EDX requièrent la coupure de l'échantillon pour déterminer les informations sur l'épaisseur de l'enrobage. Enfin, les techniques de spectroscopie RMN et EPR permettant d'attribuer des informations sur la qualité de l'enrobage ne peuvent donner des résultats directement, mais nécessitent l'aide de modèles chimiométrique.

Des observations de l'échantillon par comparaison avant et après contact avec le milieu de libération permettent de comprendre l'effet du milieu et donc les mécanismes de transport dus à une dissolution des polymères solubles. La présence en surface de cracks à la fin de la libération permet de même d'attester de l'existence d'une rupture de la résistance mécanique du film polymérique. Pour attester de ces mécanismes des études telles que les cellules de diffusion pour calculer la perméabilité des films, le calcul des coefficients de diffusion ou encore le coefficient de partition peuvent être utilisées (Marucci, 2007). Les phénomènes de gonflement quant à eux peuvent être expliqués par une accumulation de solvants impliquant les pompes osmotiques. Une accumulation de milieu au sein de la minigranule peut être mesurée par la prise en eau avant et après libération du PA et de mesures microscopiques.

Récemment, la technologie d'Imagerie Pulsée Terahertz a été introduite en tant que nouvel outil pour déterminer l'épaisseur de l'enrobage appliqué à des comprimés et granules de grandes tailles, mais aussi une nouveauté avec la création d'un indice pour définir l'homogénéité de produits appliqués à la surface (Ho et al., 2007; Ho et al., 2009; Ho et al., 2010) Les radiations terahertz font parties de la région de l'infrarouge lointain du spectre électromagnétique (2 cm^{-1} and 120 cm^{-1}) et il est bien établi que les formulations à base de polymères utilisées pour l'enrobage sont transparentes ou semi-transparentes à la lumière utilisée par le TPI (Zeitler et al., 2007b). L'obtention des images est liée à une pénétration des ondes terahertz à la surface de la minigranule qui permet par une combinaison des images d'obtenir une résolution en 3D. Après pénétration à la surface de l'échantillon, tout changement structural ou chimique est mesuré par un changement dans

l'indice de réfraction, entraînant une modification dans les pics obtenus. Le calcul du temps parcouru par l'onde entre les différentes couches permet alors de calculer l'épaisseur de la couche traversée (Fitzgerald et al., 2005).

Les calculs sont obtenus par dérivation de la distance entre plusieurs pics observés, à savoir que pour des systèmes multicouches, plusieurs pics représenteront les différentes interfaces au sein du système (Fitzgerald et al., 2005). Quand un changement structural ou chimique est mesuré dans le système, une réfraction des signaux est obtenue et la présence de plusieurs signaux dans la longueur d'onde des terahertz est observée. Le premier changement mesuré est donc rattaché à l'interface entre l'enrobage de surface et le milieu environnant. Le signal suivant correspond au changement de nature du système, dans le cas de différent polymère d'enrobage, et le temps de délai entre les deux signaux sert à caractériser l'épaisseur de la couche d'enrobage. Pour ce faire, il suffit d'appliquer la formule suivante:

$$2 d_{(coating)} = \Delta t c/n,$$

où $d_{(coating)}$ est l'épaisseur de la couche d'enrobage, Δt est le temps de délai entre les réflexions Terahertz, c est la vitesse de la lumière et n l'indice de réfraction du polymère d'enrobage (Ho et al., 2007).

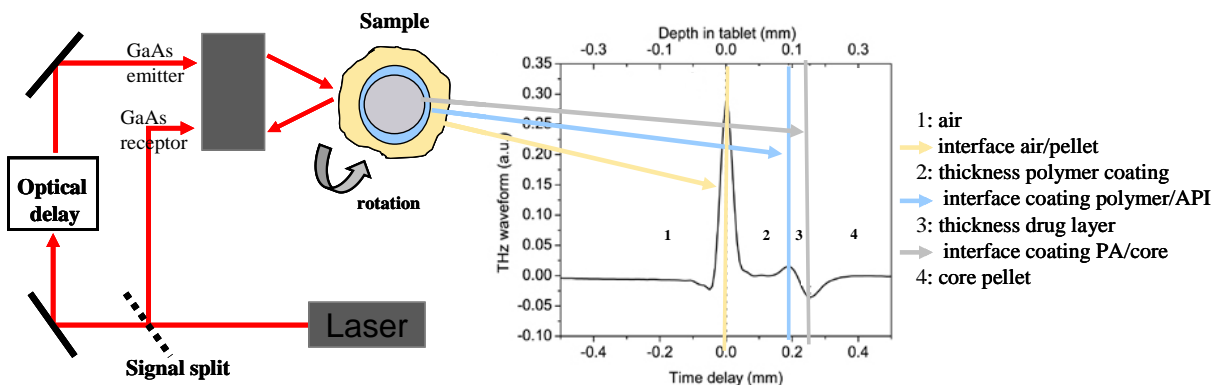


Figure 9: Représentation schématique du système TPI pour examiner des films d'enrobage appliqués à des systèmes oraux. Les signaux terahertz réfléchis montrent un changement d'interface liés à un changement dans l'indice de réfraction (Schéma adapté de Shen, 2011).

Spencer et al, en 2008, corrèlent les données de terahertz, l'épaisseur de l'enrobage et le temps moyen de dissolution (MDT). La couche active et polymérique déposée à la surface de la minigranule est mesurée et construite en modèle 3D. Dans une autre étude, l'utilisation de la technique Terahertz sur des minigranules enrobées de taille de 6 mm de diamètre a été décrite. L'influence de la méthode de préparation des noyaux, la couche de principe actif, le film d'enrobage et la méthode de mûrissement ont été analysées. Les résultats montrent une relation entre la taille et la densité de l'épaisseur de la couche polymérique avec les profils de libération *in-vitro* (Ho et al, 2009b). Ces résultats étant toujours confirmés par les analyses au MEB..

Cette technique est un bon outil pour l'analyse en ligne et le design de la qualité. Plusieurs types de défauts sont recensés pour les formes enrobées; l'uniformité de surface, la forme des noyaux d'origines, la présence de défauts de surface qui influencent les cinétiques de libération. Une étude en 2006 d'Haddish a permis de corrélérer les formes irrégulières des minigranules aux variations des profils de libération.

Un nouveau paramètre le TEFPS (*Terahertz Electric Field Peak Strength*) a été mis au point par l'équipe de Ho en 2008. Ce paramètre est obtenu par extraction du signal obtenu avec le signal de référence émis. Il permet de prédire la relative densité de surface de l'enrobage mais aussi de son homogénéité (Shen and today, 2008 ; Ho *et al.*, 2008).

Dans cette partie, la technologie terahertz a été employée pour la première fois pour analyser l'épaisseur de l'enrobage de film et d'une couche active interne, ainsi que leur uniformité, cela appliqués à des minigranules de taille standard (1 mm de diamètre). De plus, les caractéristiques de l'impact de l'enrobage sur les profils de libération ont été investiguées.

2.5. Ethanol-resistant polymeric film coatings

Dans les systèmes multiparticulaires, la quantité de la molécule active est divisée au sein de plusieurs entités, contrairement aux formes classiques unidoses. Dans le cas d'un défaut d'enrobage, les conséquences sont plus limitées avec seulement une fraction des sous unités impactées contrairement aux unidoses qui sont fortement sensibles au phénomène de « dose dumping ». Les formes multiparticulaires prennent de ce fait une plus grande importance, spécialement les minigranules. L'un des nombreux avantages des formes multiparticulaires est le temps de vidange gastrique qui est moins variable ici et une plus haute reproductibilité dans le tractus gastro intestinal (GIT), ainsi qu'une minimisation des variabilités inter et intra individus sur les profils plasmatiques (Varum, 2010).

Les systèmes à libération contrôlée contiennent généralement une concentration plus importante en PA que pour les formes conventionnelles, cela pour permettre au PA de se délivrer au niveau du corps humain à un taux préprogrammer sur un laps de temps prolongé. Si le mécanisme de contrôle de libération est basé sur les fonctions de la barrière polymérique, qui est insensible à l'eau mais soluble à l'éthanol, alors la consommation concomitante avec des substances alcoolisées pourra entraîner un « dose dumping », à savoir : « une libération de PA rapide et inattendue sur une courte période, de la totalité ou d'une fraction signifiante du contenu en PA issue de formes à libération modifiées » (Meyer and Hussain, 2005; Varum et al., *in press*). Les conséquences peuvent être sévères: (i) des concentrations toxiques de PA peuvent être atteintes, avec éventuellement des effets secondaires fatals, et (ii) un effet thérapeutique qui n'est plus garanti durant la période envisagée. Cela est particulièrement vrai pour des PA hautement actif avec une fenêtre thérapeutique étroite, comme beaucoup de PA à base d'opioïde.

IMPACT DE L'ALIMENTATION ET DE L'ALCOOL SUR LES FONCTIONS BIOLOGIQUES GASTRIQUES

Depuis plusieurs années, la qualité des systèmes et des procédés de fabrication se développe de plus en plus. La digestion orale peut être estimée et prise en considération pour les profils de libération *in-vitro*. Des différences physiologiques dans la sécrétion

salivaire, la vidange gastrique, les valeurs de pH (Van Aken et al., 2007), la mobilité GIT et l'état à jeun ou après repas va être relevant dans les cinétiques de dissolution. D'où le fait qu'en 2006, la FDA recommande pour tous nouveaux produits basés sur des PA de faire des études sur l'impact de l'alimentation sur les cinétiques de libération.

La biodisponibilité du PA à partir de comprimés est très souvent plus lente après l'administration dans un temps proche avec la prise alimentaire. Cela étant lié à l'impact du bol alimentaire sur le transit intestinal (Fadda, 2008 ; Park and al., 2004). En effet, Freire et al., (2011) montre l'importance d'une multitude de facteurs qui influence ce transit. Mais plusieurs études montrent également l'importance de la composition des comprimés. Il est à noter que l'absorption est généralement diminuée, mais parfois on observe un « dose dumping » et dans de rares cas aucun effet (Abrahamsson et al., 2004; Schug et al., 2002). Le taux d'absorption du PA pour les SDM est principalement régulé par la vidange gastrique qui influence le temps de résidence des SDM dans l'estomac, spécialement pour les particules de hautes tailles comme les comprimés. Les comprimés sont retenus de par leur grande taille au niveau de l'estomac où ils seront broyés et hydratés dans un état suffisant pour passer le sphincter pylorique, tandis que les petites particules comme les minigranules peuvent passer directement cette barrière (Meyer et al., 1981). Une augmentation du temps de rétention dans l'estomac entraîne une exposition prolongée au fluide gastrique qui prolonge le temps de dissolution. Une étude à propos de l'absorption dans l'estomac, de la caféine venant de minigranules enrobées, montre une corrélation entre le délai dans la libération et le taux de vidange gastrique. Une augmentation de la rétention du système au niveau de l'estomac pourra entraîner le phénomène de « dose dumping » du PA (Weitschies et al., 2005).

Il est en plus connu l'inhibition que vont pouvoir entraîner l'éthanol ou les solutions alcoolisées sur la vidange gastrique (Franke et al., 2005). Park and al, dans plusieurs études, font la distinction entre l'éthanol et la composition des substances alcoolisées et leurs impacts sur la vidange gastrique. En 2005 ils ont montré que pour différents types de solutions alcoolisées (bière ou vin rouge montrant des différences au niveau du contenu en éthanol et calorique) une divergence sur le temps de vidange gastrique est observée. De là, l'hypothèse d'un autre ingrédient non-alcoolique impliqué ici est posée. Un autre impact lié

à l'alcool est une augmentation de la perméabilité sur la muqueuse intestinale (Lavo et al., 1992).

IMPACT DE L'ETHANOL SUR LES SDM

En effet, le « dose dumping » peut être due au délai de la vidange gastrique mais aussi dans le cas d'un accident ou changement intentionnel dans les libérations du système. Les systèmes multiparticulaires sont généralement basés sur l'enrobage polymérique. Toutefois, la majorité des polymères étant soluble dans l'éthanol, une dissolution prématurée en cas de contact avec de fortes concentrations en éthanol. Une combinaison de l'augmentation de la perméabilité ainsi qu'une augmentation de la dissolution du système sous contact de l'éthanol induit une plus haute valeur de C_{max} et dans le pire des cas un dose dumping. Ce phénomène a été à l'origine de la suspension du Palladone en 2005 par la FDA, qui entraînait des effets secondaires néfastes dû à un « dose dumping » de PA opiacé (FDA Alert, July 2005).

Une idée pour réduire le risque de dose dumping est d'apposer un signal d'avertissement sur le produit pharmaceutique. Toutefois, il est reporté que des personnes sous emprise d'alcool souffrant de douleur chronique (low back pain) ne peuvent réduire leurs utilisations de dérivés opiacés, et donc ne respectent en rien les avertissements concernant la prise concomitante d'alcool et d'opiacés (Booker et al., 2003). De plus, le Système de Surveillance sur les Facteurs de Risques Comportementaux reportent qu'un buveurs sur 3 aux USA est un alcoolique compulsif: Ils consomment 4 ou 5 boissons (femme ou homme respectivement) sur une courte période (Serdula et al., 2004). Alors, les systèmes à libération contrôlée seront en pratique exposés à de fortes concentrations d'éthanol dans l'estomac. En 2009, Lennernaes a publié une revue très intéressante sur les formulations vulnérables à l'éthanol, pointant le risque de défaut des formes galéniques due à la co-consommation d'alcool. Les réservoirs aussi bien que les systèmes matriciels peuvent être affectés. Le produit oral à libération contrôlée « Palladone » en est un bon exemple, qui montre le risque de dose dumping quand les boissons alcoolisées sont administrées en concomitance. Le principe actif est l'hydromorphone HCl dans ce cas, la libération est issue de minigranules enrobées par un film d'éthylcellulose, ammonium methacrylate copolymère type B et alcool stéarique (Fadda et al., 2008). Les minigranules sont

administrées par une capsule à prise journalière. Une étude clinique avec des sujets sains révèle que la co-administration de 240 mL de 40 % d'alcool avec une capsule de Palladone de 12-mg résulte en un pic moyen d'hydromorphone, qui est 6 fois plus important que pour les capsules prises avec de l'eau (HMP1013). Un sujet a même montré une augmentation de la concentration plasmatique maximale 16 fois plus importante. Comme les opiacés sont hautement actifs avec de sévères effets secondaires, les fabricants ont décidé de suspendre la production industrielle. La sensibilité à l'éthanol des capsules de Palladone SR a alors été montré *in vitro* (Walden et al., 2007).

Fadda et al. (2008) ont comparés 3 formes commerciales à libération contrôlée, disponibles sur le marché contenant de l'acide 5-aminosalicylique : Pentasa, Asacol, Salofalk. Ils ont montré que l'addition d'un maximum de 40% d'éthanol au milieu de libération affecte significativement les profils de libération. Il est à noter que les modifications observées sont très variables selon le type de formulation. En 2010, Smith et al. ont reporté des études *in vitro* qui ont été conduites sur 27 produits oraux à libération modifiée. Différents types de PA ont ainsi été étudiées, incluant les opiacés, des canaux calciques bloquant, des antidépresseurs et anti arithmétiques. À noter que 9 des 10 capsules testées et 2 des 17 comprimés ont montré une accélération des cinétiques de libération dans des milieux contenant 40% d'éthanol. Aussi, Traynor et al. (2008) ont étudiés l'impact de l'addition de hautes quantités d'alcool au milieu de libération. Ils ont montrés que le taux de tramadol pour des formulations contrôlées sur 24 heures est significativement augmenté. Dans le cas de formulations à base d'hydroxypropyl methylcellulose (HPMC), Levina et al. (2007) reportent seulement des effets modérés de l'éthanol sur les libérations *in-vitro*. Une étude de Roberts et al. (2007) a montré une augmentation significative des taux d'aspirine libéré à partir de comprimés d'HPMC en présence de hautes concentrations d'éthanol. Cela peut être partiellement expliqué par le faible gonflement de la matrice et une augmentation de la solubilité du PA. Autre constat qu'est celui de Roth et al. (2009) qui ont reporté que la libération de Vérapamil à partir de systèmes matriciels est significativement plus rapide en présence de 40% d'éthanol au milieu de libération. Une autre étude très intéressante sur le potentiel impact de l'éthanol contenu dans le milieu de libération sur les performances des libérations de PA retenues par des films polymériques a été montrée par Larsson et al. (2010). Ils ont préparés des films à base d'éthylcellulose et d'hydroxypropyl cellulose (HPC)

par pulvérisation de solutions organiques polymériques et étudiés la perméabilité à l'eau du système en présence et absence d'éthanol. La perméabilité à l'eau des films avec un faible contenu en HPC augmente avec une augmentation de la concentration d'éthanol, alors que la perméabilité avec un fort contenu en HPC diminue avec une augmentation de la concentration en alcool. L'augmentation de la perméabilité pour les formules à faible contenu en HPC est certainement due à un gonflement plus prononcé de l'éthylcellulose en présence d'éthanol. La diminution de la perméabilité pour les forts contenus en HPC peut être expliquée comme suit: un réseau continu d'HPC est formé, via lequel l'eau peut relativement être transportée. L'addition d'éthanol dans le milieu de libération entraîne un gonflement de l'éthylcellulose plus important, qui peut probablement fermer partiellement les pores d'HPC. Ces différents exemples *in vitro* et *in vivo* démontrent clairement que la consommation de substances alcoolisées peut être dangereuse pour les patients traités avec un principe actif à haute activité compris dans les systèmes oraux à libération contrôlée. De ce fait, il est évident que la diminution de ces risques est nécessaire.

MELANGE DE POLYMERES D'ENROBAGE POUR LA RESISTANCE A L'ETHANOL DE SYSTEMES ORAUX A LIBERATION PROLONGEE

Des études précédentes établies au sein du laboratoire ont portées sur la formulation de films polymériques insensibles à la présence d'éthanol (Rosiaux, 2013). Ce procédé portait sur des films d'enrobages permettant d'éviter les ruptures du système même sous de fortes concentrations d'éthanol. Les dispersions aqueuses d'éthylcellulose, sont régulièrement utilisées pour des libérations contrôlées et soutenues. Le problème étant sa solubilité à l'éthanol entraînant les effets indésirables de « dose dumping » lors de la prise concomitante avec des boissons alcoolisées. L'ajout d'une fraction de gomme guar, polymère hydrophile et résistant à l'éthanol permet alors d'augmenter les cinétiques de libérations qui seront trop soutenue dans des milieux dépourvus d'éthanol, mais aussi de résister aux effets de « dose-dumping » dans des milieux riches en éthanol. Il est alors important de savoir comment ce mélange agit pour avoir cette fonction.

L'éthylcellulose est un matériel d'enrobage hydrophobique généralement utilisé pour le masquage de gout, une protection contre l'humidité et surtout pour permettre de réguler

les cinétiques de libération. C'est un polymère non toxique, non allergisant et aux propriétés non irritantes. Ce polymère peut être appliqué à partir d'une solution organiques mais aussi pour éviter l'utilisation de solvant comme le méthanol, dichlorométhane ou encore l'éthanol, des dispersions aqueuses d'éthylcellulose ont été formulées. L'éthylcellulose étant soluble dans l'éthanol, cela fait de lui un bon candidat pour la formulation de nouveaux SDM résistants à l'éthanol.

Les polysaccharides sont également de bons candidats pour les formes à libération prolongées d'enrobage. Ces excipients sont peu chers et montrent une large variété de structures et de propriétés (Hovgaard and Brondsted, 1996). De plus, ils sont hautement stables, non toxiques, biodégradables et hydrophiles. Cette propriété d'hydrophilie fait que les polysaccharides formeront un gel par gonflement sous exposition au milieu de dissolution et deviendront plus hautement solubles entraînant une rapide libération de PA (Milojevic et al., 1996). Pour contrôler les caractéristiques de solubilité et limiter la libération du PA, des polymères hydrophobiques peuvent être ajoutés. Une famille de polysaccharides naturels comprend des gommes guar avec de bonnes propriétés pour l'enrobage. Il peut être utilisé comme "pore former", pour des libérations à visées coliques (Krishnaiah et al, 1998; Prabakaran, 2011), et dans notre cas vu son insolubilité dans l'éthanol, la gomme guar semble une bonne candidate pour obtenir des enrobages résistants à l'éthanol.

Dans le cadre de cette thèse, les minigranules ont été enrobées par une dispersion de polymère. L'utilisation de l'enrobeur de type Wurster où la dispersion à enrober est pulvérisée par une buse placée au centre de la plaque du bas permet un meilleur enrobage que les autres systèmes. Le flux de pulvérisation est alors dans le même sens que l'air d'alimentation. L'utilisation du cylindre du Wurster ainsi que la surface plane présentant différentes perforations permet un mouvement d'accélération des granules via le flux d'air dans le tube. Les granules passent alors dans le cône de pulvérisation pour être enrobées par la solution/suspension. Du fait de la position de la buse au sein du flux d'air, la solution d'enrobage n'a qu'une courte distance à effectuer avant d'arriver à la surface des granules (Wheatley et al., 1997). Plusieurs théories ont été établies concernant la formation de fins films polymériques. Dans l'optique de rester centré sur notre étude, seulement le cas des films issus de dispersions aqueuses seront mentionnés ici. L'enrobage ayant lieu à une température donnée l'évaporation de l'eau permet le rapprochement des particules. Une

forte pression capillaire provoque la déformation des particules et la fusion en un film continu. La capillarité et la mobilité des chaînes polymériques sont à l'origine de la coalescence complète des particules.

Les profils de libération et la formation des films peuvent être impactés par plusieurs paramètres: le noyau des granules ainsi que le film polymérique, Le contenu solide de la dispersion (Wesseling et al., 1999 ; Pearnchob et al., 2003), Le débit de pulvérisation (Larsen, 2003), Le diamètre de la buse et la pression de l'air comprimé (Ronsse et al., 2008) et La température du procédé (Bodmeier, 1997).

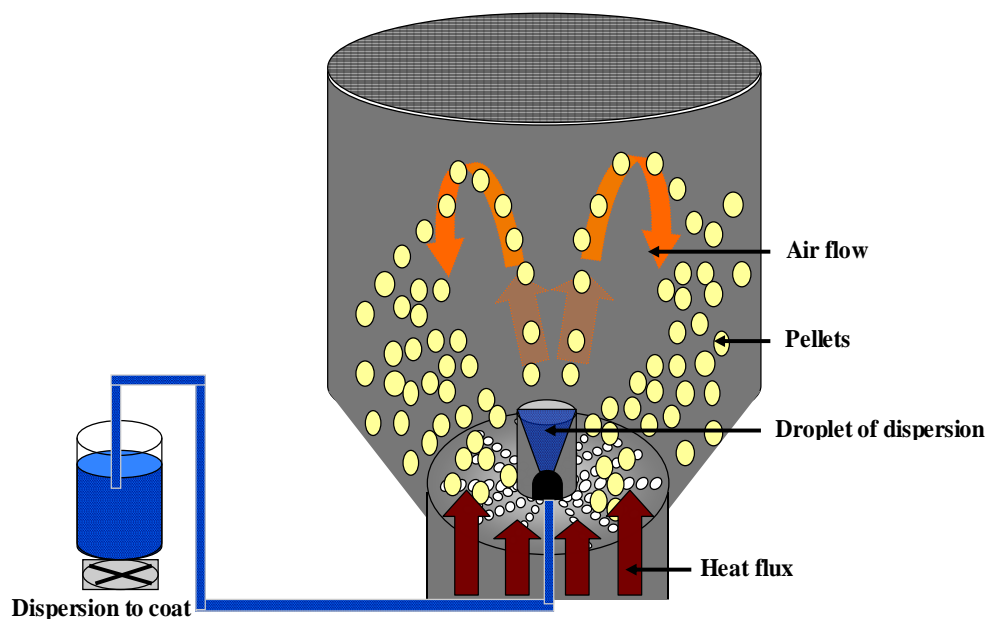


Figure 10: Représentation schématique d'un enrobeur de type Wurster.

Le mélange de polymères a des applications spécifiques sous plusieurs conditions:

- (i) pH résistance (Siepmann et al., 2008b, 2008d) ;
- (ii) ciblage du colon (Karrout et al., 2009) ;
- (iii) augmenter la stabilité au stockage (Siepmann et al., 2008c).

Le but de cette thèse a alors été d'élucider les mécanismes de transport de masse contrôlant la libération à partir de minigranules enrobées avec le nouveau film polymérique résistant à l'éthanol (Lecomte et al., 2005a,b; Siepmann et al., 2005, 2006). Le fait d'avoir fractionnée la dose de PA en plusieurs sous-unités est un grand avantage pour ce type d'application : en cas de dommage accidentel du film seulement la minigranule concernée se retrouvera en état de « dose dumping », la totalité des profils de libération provenant de l'ensemble des minigranules devraient alors être moins impactée. Basée sur des caractérisations expérimentales comprehensive du système avant et après exposition à différent milieu de libération, ainsi que sur des caractérisations de la forme (par exemple des mesures de prise en eau ou des changements dans la résistance après mise en contact avec le milieu), la détermination d'un système spécifique peut être faites avec le coefficient de diffusion apparent du principe actif au travers du film d'enrobage (Siepmann and Peppas, 2001; Siepmann and Goepferich, 2001; Borgquist et al., 2004; Marucci et al., 2008).

Des équations mathématiques appropriées seront développées pour permettre le développement d'un modèle mathématique pour faciliter l'optimisation des produits à base de PA : Des simulations *In silico* devraient être capables de prédire les effets des paramètres de formulation sur les cinétiques de libération.

3. References

Abd El-Halim S., Amin M., El-Gazayerly O., Abd El-Gawad N. Comparative study on the different techniques for the preparation of sustained-release hydrophobic matrices of a highly water-soluble drug. *Drug Discoveries & Therapeutics*. **2010**; 4:484-492.

Abrahamsson B., Albery T., Eriksson A., Gustafsson I., Sjöberg M. Food effects on tablet disintegration. *European Journal of Pharmaceutical Sciences*. **2004**; 22:165–172.

Andersson M., Holmquist B., Lindquist J., Nilsson O., Wahlund K.G. Analysis of film coating thickness and surface area of pharmaceutical pellets using fluorescence microscopy and image analysis. *Journal of Pharmaceutical and Biomedical Analysis*. **2000**; 22:325-339.

Baker, R. (Ed.), *Controlled release of biologically active agents*, John Wiley & Sons, New York, **1987**.

Barthelemy P., Laforêt J., Farah N., Joachim J. Compritol® 888 ATO: an innovative hot-melt coating agent for prolonged-release drug formulations. *European Journal of Pharmaceutics and Biopharmaceutics*. **1999**; 47:87–90.

Bechgaard H., Nielsen G.H. Controlled-release multiple-units and single-unit doses - Literature review. *Drug Development and Industrial Pharmacy*. **1978**; 4:53-67.

Bodmeier R., Guo X., Paeratakul O., Process and formulation factors affecting the drug release from pellets coated with the ethylcellulose-pseudolatex Aquacoat. Aqueous polymeric coatings for pharmaceutical dosage forms. J.W. McGinity, Editor. **1997**; Marcel Dekker: New York.

Booker E.A., Haig A.J., Geisser M.E., Yamakawa, K. Alcohol use self report in chronic back pain--relationships to psychosocial factors, function performance, and medication use. *Disabil. Rehabil.* **2003**; 25: 1271-1277.

Borgquist P., Zackrisson G., Nilsson B. Axelsson, A. Simulation and parametric study of a film-coated controlled-release pharmaceutical. *Journal Controlled Release*. **2002**; 80: 229-245,.

Borgquist P., Nevsten P., Nilsson B., Wallenberg L.R., Axelsson, A. Simulation of the release from a multiparticulate system validated by single pellet and dose release experiments. *Journal of Controlled Release*. **2004**; 97: 453-465.

Borgquist P., Körner A., Piculell L., Larsson A., Axelsson A. A model for the drug release from a polymer matrix tablet—effects of swelling and dissolution. *Journal of Controlled Release*. **2006**; 113:216-225.

Brazel C.S., Peppas N.A. Modeling of drug release from Swellable polymers, *European Journal of Pharmaceutics and Biopharmaceutics*. **2000**; 49:47-58.

Cahyadi C., Karande A.D., Chan L.W., Heng P.W.S. Comparative study of non-destructive methods to quantify thickness of tablet coatings. *International Journal of Pharmaceutics*. **2010**;398:39–49.

Caraballo I., Millán M., Rabasco A.M., Leuenberger H. Zero-order release periods in inert matrices. Influence of the distance to the percolation threshold. *Pharmaceutica Acta Helveticae*. **1996**; 71:335–339.

Cogdill R.P., Forcht R.N., Shen Y., Taday P.F., Creekmore J.R.,Anderson C.A., Drennen J.K. Comparison of terahertz pulse imaging and near-infrared spectroscopy for rapid, non-destructive analysis of tablet coating thickness and uniformity. *Journal Pharmaceutics. Innovation*. **2007**; 2:29–36.

Cole G., in: Cole G., Hogan J., Aulton M.E. (Eds.). *Pharmaceutical Coating Technology*. Taylor & Francis Ltd., Philadelphia. **1995**:1-5.

Crank J., **1975**. *The Mathematics of Diffusion*, Clarendon Press, Oxford.

De Beer T., Burggraeve A., Fonteyne M., Saerens L., Remon J.P., Vervaet C. Near infrared and Raman spectroscopy for the in-process monitoring of pharmaceutical production processes. *International Journal of Pharmaceutics*. **2011**; 417:32–47.

Depypere F., Van Oostveldt P., Pieters, J.G., Dewettinck, K. Quantification of microparticle coating quality by confocal laser scanning microscopy (CLSM). *European Journal of Pharmaceutics and Biopharmaceutics*. **2009**; 73:179-186.

Ensslin S., Moll K.P., Paulus K., Mader K. New insight into modified release pellets - Internal structure and drug release mechanism. *Journal Controlled Release*. **2008**; 128:149-156.

Ensslin, S., Moll, K.P., Metz, H., Otz, M., Mader, K. Modulating pH-independent release from coated pellets: effect of coating composition on solubilization processes and drug release. *European Journal of Pharmaceutics and Biopharmaceutics*. **2009**; 72:111-118.

Fadda H.M., Mohamed M.A.M., Basit,A.W. Impairment of the *in vitro* release behaviour of oral modified release preparations in the presence of alcohol. *International Journal of Pharmaceutics*. **2008**; 360: 171-176.

Faham A., Prinderre P., Farah N., Eichler K.D., Kalantzis G., Joachim J. Hot-Melt Coating Technology. I. Influence of Compritol® 888 Ato and Granule Size on Theophylline Release. *Die Pharmazie*. **2000**; 26:167-176.

Fan, L.T., Singh, S.K. (Eds.), *Controlled release. A quantitative treatment*, Springer-Verlag, Berlin, **1989**.

http://www.fda.gov/ohrms/dockets/ac/05/briefing/2005-4187B1_01_08-Alcohol-Induced.pdf

Fitzgerald A.J., Cole B.E., Taday P.F. Nondestructive analysis of tablet coating thicknesses using terahertz pulsed imaging. *Journal of Pharmaceutical Sciences*. **2005**; 94:177-183.

Franke A, Luebke N, Links M, Harder H, Teyssen S, Singer MV. Effect of ethanol and some alcoholic beverages on gastric emptying of solid meals in men. *Gastroenterology*. 2003;124:A575–A576.

Frenning G. Modelling drug release from inert matrix systems: From moving-boundary to continuous-field descriptions. *International Journal of Pharmaceutics*. **2011**; 418:88-99.

Gao Y., Muzzio F.J., Ierapetritou M.G. A review of the Residence Time Distribution (RTD) applications in solid unit operations. *Powder Technology*. **2012**; 228:416–423.

Grassi M., Colombo I. Mathematical modelling of drug permeation through a swollen membrane, *Journal of Controlled Release*. **1999**; 59:343-359.

Gendre C., Genty M., Boiret M. et al. Development of a Process Analytical Technology (PAT) for in-line monitoring of film thickness and mass of coating materials during a pan coating operation. *European Journal of Pharmaceutical Sciences*. **2011**; 43:244–250.

Gendre C., Genty M., Silva J.C. et al. Comprehensive study of dynamic curing effect on tablet coating structure. *European Journal of Pharmaceutics and Biopharmaceutics*. **2012**; 81:657–665.

Grassi M., Voinovich D., Franceschinis E., Perissutti B., Filipovic-Grcic J. Theoretical and experimental study on theophylline release from stearic acid cylindrical delivery systems. *Journal of Controlled Release*. **2003**; 92:275-289.

Gueres S., Siepmann F., Siepmann J., Kleinebudde P. Drug release from extruded solid lipid matrices: Theoretical predictions and independent experiments. *European Journal of Pharmaceutics and Biopharmaceutics*. **2012**; 80:122-129.

Guse C., Koennings S., Kreye F., Siepmann F., Goepferich A., Siepmann J. Drug release from lipid-based implants: Elucidation of the underlying mass transport mechanisms. *International Journal of Pharmaceutics*. **2006**; 314:137-144.

Haddish-Berhane N., Jeong S.H., Haghghi K., Park K. Modeling film-coat non-uniformity in polymer coated pellets: A stochastic approach. *International Journal of Pharmaceutics*. **2006**; 323:64–71.

Hariharan D., Peppas N.A. Modelling of water transport and solute release in physiologically sensitive gels, *Journal of Controlled Release*. **1993**; 23:123-135.

Herrmann S., Winter G., Mohl S., Siepmann F., Siepmann J. Mechanisms controlling protein release from lipidic implants: Effects of PEG addition. *Journal of Controlled Release*. **2007a**; 118:161-168.

Herrmann S., Mohl S., Siepmann F., Siepmann J., Winter G. New insight into the role of polyethylene glycol acting as protein release modifier in lipidic implants. *Pharmaceutical Research*. **2007b**; 24:1527-1537.

HMP1013. A randomised, open-label, single dose, 4 way crossover study of the effects of varying doses of ethanol on the pharmacokinetic characteristics of 12 mg hydromorphone hydrochloride extended release capsules (Palladone™) in two groups (fed and fasted) of healthy volunteers. Data in house, Purdue Pharma (US).

Haddish-Berhane N., Jeong S.H., Haghghi K., Park, K. Modeling film-coat non-uniformity in polymer coated pellets: a stochastic approach. *International Journal of Pharmaceutics*. **2006**; 323:64-71.

Heinicke G., Schwartz J.B. Ammonio polymethacrylate-coated diltiazem: Drug release from single pellets, media dependence, and swelling behavior. *Pharmaceutical Development and Technology*. **2007**; 12:285-296.

Ho L., Müller R., Römer M., Gordon K.C., Heinämäki J., Kleinebudde P., Pepper M., Rades T., Shen Y.C., Strachan C.J., Taday P.F., Zeitler J.A. Analysis of sustained-release tablet film coats using terahertz pulsed imaging. *Journal Controlled Release*. **2007**;119:253-261.

Ho L., Müller R., Gordon K.C., Kleinebudde P., Pepper M., Rades T., Shen Y.C., Taday P.F., Zeitler J.A. Applications of terahertz pulsed imaging to sustained-release tablet film coating quality assessment and dissolution performance. *Journal Controlled Release*. **2008**; 127:79-87.

Ho L., Müller R., Gordon K.C., et al. Terahertz pulsed imaging as an analytical tool for sustained-release tablet film coating. *European Journal of Pharmaceutics and Biopharmaceutics*. **2009a**; 71:117–123.

Ho L., Cuppok Y., Muschert S., Gordon K.C., Pepper M., Shen Y., Siepmann F., Siepmann J., Taday P.F., Rades T., Effects of film coating thickness and drug layer uniformity on in vitro drug release from sustained-release coated pellets: a case study using terahertz pulsed imaging. *International Journal of Pharmaceutics*. **2009b**; 382:151-159.

Ho L., Müller R., Krüger C., Gordon K.C., Kleinebudde P., Pepper M., Rades T., Shen Y.C., Taday P.F., Zeitler J.A. Investigating Dissolution Performance Critical Areas on Coated Tablets: A Case Study Using Terahertz Pulsed Imaging. *Journal of Pharmaceutical Sciences*. **2010**; 99:392-402.

Hovgaard L., Bronsted H. Current applications of polysaccharides in colon targeting. *Critical review in Therapeutic Drug Carrier System*. **1996**; 13:185-223.

Jannin V., Bérard V., N'Diaye A., Andrès C., Pourcelot Y. Comparative study of the lubricant performance of Compritol® 888 ATO either used by blending or by hot melt coating. *International Journal of Pharmaceutics*. **2003**; 262:39–45.

Karrouit Y., Neut C., Wils D, et al. Novel polymeric film coatings for colon targeting: Drug release from coated pellets. *European Journal of Pharmaceutical Sciences*. **2009**; 37:427–433.

Kaunisto E., Marucci M., Borgquist P., Axelsson A. Mechanistic modelling of drug release from polymer-coated and swelling and dissolving polymer matrix systems. *International Journal of Pharmaceutics*. **2011**; 418:54-77.

Kreye F., Siepmann F., Siepmann J. Lipid implants as drug delivery systems. *Expert Opinion Drug Delivery*. 2008; 3:291-307.

Kreye F., Siepmann F., Siepmann J. Drug release mechanisms of compressed lipid implants. *International Journal of Pharmaceutics*. **2011a**; 404:27-35.

Kreye F., Siepmann F., Zimmer A., Willart J.F., Descamps M., Siepmann J. Controlled release implants based on cast lipid blends, *Eur. J. Pharm. Sci.* **2011b**; 43:78-83.

Kreye F., Siepmann F., Willart J.F., Descamps M., Siepmann J. Drug release mechanisms of cast lipid implants. *European Journal of Pharmaceutics and Biopharmaceutics*. **2011c**; 78:394-400.

Kreye F., Siepmann F., Zimmer A., Willart J.F., Descamps M., Siepmann J. Cast lipid implants for controlled drug delivery: Importance of the tempering conditions. *J. Pharm. Sci.* **2011d**; 100:3471-3481.

Krishnaiah Y.S., Satyanarayana S., Rama Prasad Y., Narasimha Rao S. Evaluation of guar gum as a compression coat for drug targeting to colon. *International Journal of Pharmaceutics*. **1998**; 171:137–146.

Kucera S., Felton L., McGinity J. Physical Aging in Pharmaceutical Polymers and the Effect on Solid Oral Dosage Form Stability. *International Journal of Pharmaceutics.*, **in press**.

Larsson M., Hjaertstam J., Berndtsson J., Stading M., Larsson A. Effect of ethanol on the water permeability of controlled release films composed of ethyl cellulose and hydroxypropyl cellulose. *European Journal of Pharmaceutics and Biopharmaceutics*. **2010**; 76: 428-432.

Lavö B, Colombel JF, Knutsson L, Hällgren R. Acute exposure of small intestine to ethanol induces mucosal leakage and prostaglandin E2 synthesis. *Gastroenterology*. 1992;102:468-73.1.

Lecomte F., Siepmann J., Walther M., MacRae R.J., Bodmeier R. Blends of enteric and GIT-insoluble polymers used for film coating: Physicochemical characterization and drug release patterns. *Journal of Controlled Release*. **2003**; 89: 457-471.

Lecomte F., Siepmann J., Walther M., MacRae R.J., Bodmeier, R. Polymer blends used for the coating of multiparticulates: Comparison of aqueous and organic coating techniques. *Pharmaceutical Research*. **2004a**; 21: 882-890.

Lecomte F., Siepmann J., Walther M., MacRae R.J., Bodmeier, R. Polymer blends used for the aqueous coating of solid dosage forms: Importance of the type of plasticizer. *Journal Controlled Release*. **2004b**; 99:1-13.

Lecomte F., Siepmann J., Walther M., MacRae R.J. Bodmeier R. pH-sensitive polymer blends used as coating materials to control drug release from spherical beads: Elucidation of the underlying mass transport mechanisms. *Pharmaceutical Research*. **2005a**; 22:1129-1141.

Lecomte F., Siepmann J., Walther M., MacRae R.J., Bodmeier R. pH-sensitive polymer blends used as coating materials to control drug release from spherical beads: Importance of the type of core. *Biomacromolecules* **2005b**; 6: 2074-2083.

Lee M.J., Seo D.Y., Lee H.E., et al. In line NIR quantification of film thickness on pharmaceutical pellets during a fluid bed coating process. *International Journal of Pharmaceutics*. **2011**;403:66–72.

Lennernaes H. Ethanol-drug absorption interaction: Potential for a significant effect on the plasma pharmacokinetics of ethanol vulnerable formulations. *Mol. Pharm.* **2009**; 6: 1429-1440.

Levina M., Vuong H., Rajabi-Siahboomi A.R. The influence of hydro-alcoholic media on hypromellose matrix systems. *Drug Development and Industrial Pharmacy*. **2007**; 33:1125–1134.

Mallapragada S.K., Peppas N.A. Crystal dissolution-controlled release systems: I. Physical characteristics and modeling analysis. *Journal of Controlled Release*. **1997**; 45:87-94.

Maroni A., Zema L., Loreti G., Palugan L., Gazzaniga A. Film coatings for oral pulsatile release. *International Journal of Pharmaceutics.*, **in press**.

Marucci M., Ragnarsson G., Nyman U., Axelsson A. Mechanistic model for drug release during the lag phase from pellets coated with a semi-permeable membrane. *Journal Controlled Release*. **2008**; 127: 31–40.

Marucci M., Ragnarsson G., Nilsson B., Axelsson A. Osmotic pumping release from ethyl–hydroxypropyl–cellulose-coated pellets: A new mechanistic model. *Journal Controlled Release*. **2010**; 142: 53-60.

Marucci M., Ragnarsson G., von Corswant C., Welinder A., Jarke A., Iselau F., Axelsson A. Polymer leaching from film coating: Effects on the coating transport properties. *International Journal of Pharmaceutics*. **2011**; 411: 43-48.

Maschke A., Lucke A., Vogelhuber W., Fischbach C., Appel B., Blunk T., Göpferich A. Lipids: An Alternative Material for Protein and Peptide Release. *Carrier Based drug Delivery*. **2004**; 13:176-196.

Maurer L., Leuenberger H. Terahertz pulsed imaging and near infrared imaging to monitor the coating process of pharmaceutical tablets. *International Journal of Pharmaceutics*. **2009**;370:8–16.

McConnell EL, Fadda HM, Basit AW. Gut instincts: Explorations in intestinal physiology and drug delivery. *International Journal of Pharmaceutics*. 2008;364:213–226.

Meyer R.J., Hussain A.S. Awareness Topic: Mitigating the risks of ethanol induced dose dumping from oral sustained/controlled release dosage forms. FDA’s ACPS Meeting, October **2005**.

Milojevic S., Newton J.M., Cummings J.H., et al. Amylose as a coating for drug delivery to the colon: Preparation and in vitro evaluation using glucose pellets. *Journal of Controlled Release*. **1996**; 38:85–94.

Müller J, Brock D, Knop K, Axel Zeitler J, Kleinebudde P. Prediction of dissolution time and coating thickness of sustained release formulations using Raman spectroscopy and terahertz pulsed imaging. *European Journal of Pharmaceutics and Biopharmaceutics*. **2012**;80:690–697.

Muschert S., Siepmann F., Leclercq B., Carlin B., Siepmann J. Drug release mechanisms from ethylcellulose:PVA-PEG graft copolymer coated pellets. *European Journal of Pharmaceutics and Biopharmaceutics*. **2009a**; 72: 130-137.

Muschert S., Siepmann F., Cuppok Y., Leclercq B., Carlin B., Siepmann, J. Improved long term stability of aqueous ethylcellulose film coatings: Importance of the type of drug and starter core. *International Journal of Pharmaceutics*. **2009**; 368:138-145.

Muschert S., Siepmann F., Leclercq B., Carlin B., Siepmann J. Prediction of drug release from ethylcellulose coated pellets. *Journal of Controlled Release*. **2009b**; 135: 71-79.

Moes J.J., Ruijken M.M., Gout E., Frijlink H.W., Ugwoke M.I. Application of process analytical technology in tablet process development using NIR spectroscopy: Blend uniformity, content uniformity and coating thickness measurements. *International Journal of Pharmaceutics*. **2008**;357:108–118.

Möltgen C.V., Herdling T., Reich G. A novel multivariate approach using science-based calibration for direct coating thickness determination in real-time NIR process monitoring. *European Journal of Pharmaceutics and Biopharmaceutics*. (2013).

Obaidat A.A., Obaidat R.M. Controlled release of tramadol hydrochloride from matrices prepared using glyceryl behenate. *European Journal of Pharmaceutics and Biopharmaceutics*. **2001**; 52:231–235.

Pearnchob N., Bodmeier R. Dry polymer powder coating and comparison with conventional liquid-based coatings for Eudragit® RS, ethylcellulose and shellac. *European Journal of PPharmaceutics and Biopharmaceutics*. **2003**; 56:363–369.

Peppas N.A. Historical perspective on advanced drug delivery: How engineering design and mathematical modeling helped the field mature. *Adv Drug del rev*. 2013; 65:5-9.

Prabaharan M. Prospective of guar gum and its derivatives as controlled drug delivery systems. *International Journal of Biological Macromolecules*. **2011**; 49:117–124.

Reitz E., Podhaisky H., Ely D., Thommes M. Residence time modeling of hot melt extrusion processes. *European Journal of Pharmaceutics and Biopharmaceutics*. **2013**. In Press, Corrected Proof.

Ringqvist A., Taylor L.S., Ekelund K., Ragnarsson G., Engstrom S., Axelsson, A. Atomic force microscopy analysis and confocal Raman microimaging of coated pellets. *International Journal of Pharmaceutics*. **2003**;267:35-47.

Roberts M., Cespi M., Ford J.L., Dyas A.M., Downing J., Martini L.G., Crowley P.J. Influence of ethanol on aspirin release from hypromellose matrices. *International Journal of Pharmaceutics*. **2007**; 332: 31-37.

Römer M., Heinämäki J., Strachan C., Sandler N., Yliruusi J. Prediction of tablet film-coating thickness using a rotating plate coating system and NIR spectroscopy. *AAPS PharmSciTech*. **2008**; 9:1047–1053

Ronsse F., Pieters J.G., Dewettinck K. Modelling side-effect spray drying in top-spray fluidised bed coating processes. *Journal of Food Engineering*. **2008**; 86:529–541.

Rosiaux Y., Muschert S., Chokshi R., Leclercq B., Siepmann F., Siepmann J. Ethanol-resistant polymeric film coatings for controlled drug delivery. *Journal of Controlled Release*. **2013**; 169:1–9.

Roth W., Setnik B., Zietsch M., Burst A., Breitenbach J., Sellers E., Brennan D. Ethanol effects on drug release from Verapamil Meltrex, an innovative melt extruded formulation. *International Journal of Pharmaceutics*. **2009**; 368:72-75.

Sackett C.K., Narasimhan B. Mathematical modeling of polymer erosion: Consequences for drug delivery. *International Journal of Pharmaceutics*. **2011**; 418:104-114.

Saerens L., Dierickx L., Lenain B., Vervaet C., Remon J.P., Beer T.D.. Raman spectroscopy for the in-line polymer–drug quantification and solid state characterization during a pharmaceutical hot-melt extrusion process. *European Journal of Pharmaceutics and Biopharmaceutics*. **2011**;77:158–163.

Saerens L., Dierickx L., Quinten T., et al. In-line NIR spectroscopy for the understanding of polymer–drug interaction during pharmaceutical hot-melt extrusion. *European Journal of Pharmaceutics and Biopharmaceutics*. **2012**;81:230–237.

Schug B.S., Brendel E., Wolf D., Wonnemann M., Wargenau M., Blume H.H. Formulation-dependent food effects demonstrated for nifedipine modified-release preparations marketed in the European Union. *European Journal of Pharmaceutical Sciences*. **2002**; 15:279–285.

Serdula M.K., Brewer R.D., Gillespie C., Denny C.H., Mokdad A. Trends in alcohol use and binge drinking, 1985-1999 results of a multi-state survey. *Am. J. Prev. Med.* **2004**; 26:294-298.

Shen Y.C., Taday P.F., Pepper M. Elimination of scattering effects in spectral measurement of granulated materials using terahertz pulsed spectroscopy. *Applied Physics Letters*. **2008**;92.

Siepmann J., Goepferich A. Mathematical modeling of bioerodible, polymeric drug delivery systems. *Advanced Drug Delivery Reviews*. **2001**; 48:229-247.

Siepmann J., Peppas N.A. Modeling of drug release from delivery systems based on hydroxypropyl methylcellulose (HPMC). *Advanced Drug Delivery Reviews*. **2001**; 48:139-157.

Siepmann F., Siepmann J., Walther M., MacRae R.J., Bodmeier R. Blends of aqueous polymer dispersions used for pellet coating: Importance of the particle size. *Journal Controlled Release*. **2005**; 105: 226-239.

Siepmann J., Siepmann F., Florence A.T. Local controlled drug delivery to the brain: Mathematical modeling of the underlying mass transport mechanisms. *International Journal of Pharmaceutics*. **2006**; 314:101-119.

Siepmann F., Siepmann J., Walther M., MacRae R.J., Bodmeier R. Polymer blends for controlled release coatings. *Journal Controlled Release*. **2007a**; 125:1-15.

Siepmann F., Hoffmann A., Leclercq B., Carlin B., Siepmann J. How to adjust desired drug release patterns from ethylcellulose-coated dosage forms. *Journal Controlled Release*. **2007b**; 119:182-189.

Siepmann F., Herrmann S., Winter G., Siepmann J. A novel mathematical model quantifying drug release from lipid implants. *Journal of Controlled Release*. **2008**; 128:233-240.

Siepmann F., Muschert S., Leclercq B., Carlin, B., Siepmann J. How to improve the storage stability of aqueous polymeric film coatings. *Journal Controlled Release*. **2008**; 126:26-33.

Siepmann J., Siepmann F. Mathematical modeling of drug delivery. *International Journal of Pharmaceutics*. **2008**; 364: 328-343.

Siepmann F., Eckart K., Maschke A., Kolter K., Siepmann J. Modeling drug release from PVAc/PVP matrix tablets. *Journal of Controlled Release*. **2010**; 141:216-222.

Siepmann J., Siepmann F. Mathematical modeling of drug release from lipid dosage forms. *International Journal of Pharmaceutics*. **2011**; 418:42-53.

Siepmann, J; Siepmann, F. Modeling of diffusion controlled drug delivery. *Journal of Controlled Release*. **2012a**;161, 351-362.

Siepmann J., Siepmann F. Modeling of diffusion controlled drug delivery. *Journal Controlled Release*. **2012b**; 61:351-362.

Siepmann J., Peppas N.A. Modeling of drug release from delivery systems based on hydroxypropyl methylcellulose (HPMC), *Adv. Drug Del. Rev.* **2012b**; 64:163-174.

Siepmann J. In-silico simulations of advanced drug delivery systems: What will the future offer? *International Journal of Pharmaceutics*. **2013a**; 454:512-516.

Siepmann J., Siepmann F. Mathematical modeling of drug dissolution, *International Journal of Pharmaceutics*. **2013b**; 453:12-24.

Siepmann J., Siepmann F. Mathematical modeling of drug dissolution. *International Journal of Pharmaceutics*, **in press**.

Singer M.V., Leffmann C., Eysselein V.E., Calden H., Goebell H. Action of ethanol and some alcoholic beverages on gastric acid secretion and release of gastrin in humans. *Gastroenterology*. 1987; 93:1247-1254.

Smith G.D. Numerical Solution of Partial Differential Equations: Finite Difference Methods, Clarendon Press, Oxford, **1985**.

Smith A.P., Moore T.W., Westenberger B.J., Doub W.H. *In vitro* dissolution of oral modified-release tablets and capsules in ethanolic media. *International Journal of Pharmaceutics*. **2010**; 398:93-96.

Tanquary, A.C., Lacey, R.E. (Eds.), *Controlled release of biologically active agents*, Plenum Press, New York, **1974**.

Traynor M.J., Brown M.B., Pannala A., Beck P., Maetin G.P. Influence of alcohol on the release of tramadol from 24-h controlled-release formulations during *in vitro* dissolution experiments. *Drug Development and Industrial Pharmacy*. **2008**; 34:885-889.

Van Aken G.A., Vingerhoeds M.H., de Hoog EHA. Food colloids under oral conditions. *Current Opinion in Colloid & Interface Science*. **2007**; 12:251–262.

Varum FJO, Merchant HA, Basit AW. Oral modified-release formulations in motion: The relationship between gastrointestinal transit and drug absorption. *International Journal of Pharmaceutics*. **2010**;395:26–36.

1.

Varum F.J.O., Hatton G.B., Basit A. Food, physiology and drug delivery. *International Journal of Pharmaceutics.*, **in press**.

Vercruyse J., Córdoba Díaz D., Peeters E., et al. Continuous twin screw granulation: Influence of process variables on granule and tablet quality. *European Journal of Pharmaceutics and Biopharmaceutics*. **2012**; 82:205–211.

Verhoeven E., De Beer T.R.M., Van den Mooter G., Remon J.P., Vervaet C. Influence of formulation and process parameters on the release characteristics of ethylcellulose sustained-release mini-matrices produced by hot-melt extrusion. *European Journal of Pharmaceutics and Biopharmaceutics*. **2008**; 69:312–319.

Vogelhuber W., Magni E., Gazzaniga A., Göpferich A. Monolithic glyceryl trimyristate matrices for parenteral drug release applications. *European Journal of Pharmaceutics and Biopharmaceutics*. **2003**; 55:133–138.

Walden M., Nicholls F.A., Smith K.J. The effect of ethanol on the release of opioids from oral prolonged-release preparations. *Drug Development and Industrial Pharmacy*. **2007**; 33:1101-1111.

Weitschies W., Wedemeyer R-S, Kosch O., et al. Impact of the intragastric location of extended release tablets on food interactions. *Journal of Controlled Release*. **2005**; 108:375–385.

Wesseling M., Bodmeier R. Drug release from beads coated with an aqueous colloidal ethylcellulose dispersion, Aquacoat®, or an organic ethylcellulose solution. *European Journal of Pharmaceutics and Biopharmaceutics*. **1999**; 47:33–38.

Wheatley T.A., Steuernagel C.R., Latex emulsion for controlled drug delivery. Aqueous polymeric coatings for pharmaceutical dosage forms. J.W. McGinity, Editor. 1997; Marcel Dekker: New York.

Yin C., Li X. Anomalous diffusion of drug release from a slab matrix: Fractional diffusion models, *International Journal of Pharmaceutics*. **2011**; 418:78-87.

Zaky A., Elbakry A., Ehmer A., Breunig M., Goepferich A. The mechanism of protein release from triglyceride microspheres. *Journal of Controlled Release*. **2010**; 147:202-210.

Zeitler J.A., Shen Y., Baker C., Taday P.F., Pepper M., Rades T. Analysis of coating structures and interfaces in solid oral dosage forms by three dimensional terahertz pulsed imaging. *Journal of Pharmaceutical Sciences*. **2007a**; 96:330-340.

Zeitler J.A., Taday P.F., Newnham D.A., Pepper M., Gordon K.C., Rades, T. Terahertz pulsed spectroscopy and imaging in the pharmaceutical setting-a review. *Journal of Pharmacy and Pharmacology*. **2007b**; 59, 209-223.

Zhong S., Shen Y.C., Ho L., et al. Non-destructive quantification of pharmaceutical tablet coatings using terahertz pulsed imaging and optical coherence tomography. *Optics and Lasers in Engineering*. **2011**;49:361–365.

4. *In-silico* simulation of niacin release from lipid tablets:

Theoretical predictions and independent experiments

4.1. Materials and methods

4.1.1. Materials

Niacin (Sigma-Aldrich, Steinheim, Germany); glyceryl dibehenate (Compritol[®] 888; Gattefosse, Saint Priest, France); lactose (monohydrate, Flowlac 100; Meggle, Wasserburg, Germany); magnesium stearate (Fagron, Waregem, Belgium).

4.1.2. Tablet preparation

Direct Compression: Niacin, lactose and glyceryl dibehenate powders were sieved (0.71 mm) and blended: 10 min manually using a pestle and mortar, followed by 15 min with a turbula mixer at 64 rpm (Bachoven, Basle, Switzerland). Magnesium stearate was added and the powders were further blended in the turbula mixer for 2 min. The percentages of lactose and magnesium stearate were kept constant in all formulations: 12.4 and 0.5 % (w:w), respectively. The niacin:lipid ratios were varied as indicated. The blends were compressed on a single punch tableting machine (EK 0, Korsch, Berlin, Germany), equipped with flat-faced punches. The tablet diameter and height were varied as indicated. The tablet weight (500 mg) and hardness (60 N, Dr. Schleuniger Pharmatron, Solothurn, Switzerland) were kept constant.

Hot-melt extrusion/grinding/compression: Niacin and glyceryl dibehenate powders were sieved (0.71 mm) and blended: 10 min manually using a pestle and mortar, followed by 15 min with a turbula mixer at 64 rpm (Bachoven). The blend ratio was varied as indicated. The mixtures were hot-melt extruded using a Leistritz “Nano 16” apparatus (Leistritz, Nuremberg, Germany), equipped with a co-rotating twin screw (diameter = 16 mm, 5 heating zones, kneading elements in zones 2 and 3, diameter of the die orifice = 1 mm). The screw speed and feeding rate were kept constant at 50 rpm and 5 mL/min, respectively. The feeding zone (zone 1) was kept at room temperature. The temperature of zones 2 to 5 was: 60, 60, 60, and 65°C, respectively (the melting point of the lipid being equal to 70 °C). The extrudates were air-cooled and manually ground using a pestle and mortar for 10 min.

The obtained powder was sieved (0.71 mm) and blended with sieved lactose (0.71 mm). The mixture was blended manually using a pestle and mortar for 10 min, followed by 15 min in a turbula mixer (64 rpm, Bachoven). Magnesium stearate was added, and mixing was continued for 2 min. The percentages of lactose and magnesium stearate were kept constant in all formulations: 12.4 and 0.5 % (w:w), respectively. The obtained powder blend was compressed using a single punch tableting machine (EK 0), as described above.

4.1.3. Tablet characterization

In vitro niacin release from the lipid tablets was measured using the USP 35 paddle apparatus (900 mL, 0.1 N HCl; 37 °C; 100 rpm; $n=6$) (Sotax, Basle, Switzerland). At predetermined time points, 3 mL samples were withdrawn (not replaced), filtered (0.45 μm) and analyzed UV-spectrophotometrically ($\lambda=261.0$ nm; Shimadzu UV-1650PC, Shimadzu France, Champs-sur-Marne, France). The tablet dimensions (diameter and height) and potential dynamic changes thereof upon exposure to the release medium were measured using a thickness gauge (samples were treated as for the *in vitro* vitamin release measurements). The tablet morphology was observed with an optical image analysis system (Nikon SMZ-U; Nikon, Tokyo, Japan), equipped with a Zeiss camera (AxioCam ICc 1, Zeiss, Jena, Germany) before and after exposure to the release medium (under the same conditions as described above).

4.1.4. Equilibrium solubility measurements

The equilibrium solubility of niacin powder (as received) was determined in agitated flasks in 0.1 M HCl. Excess amounts of niacin were exposed to 100 mL medium at 37°C under horizontal shaking (80 rpm, GFL 3033; Gesellschaft fuer Labortechnik, Burgwedel, Germany). Every 24 h, samples were withdrawn, filtered and analyzed by UV-spectroscopy for their vitamin content (as described above) until equilibrium was reached. Each experiment was conducted in triplicate.

4.2. Results and discussion

4.2.1. Model development

A mechanistically realistic mathematical model should be based on a thorough physico-chemical characterization of the respective dosage form before and after exposure to the release medium. Crucial features to be studied include potential changes in the system's geometry and size (e.g. due to device disintegration and/or dissolution), limited drug/vitamin solubility effects (within the dosage form and/or the surrounding bulk fluid) as well as time- and/or position-dependent changes in the conditions for mass transport (e.g. due to excipient swelling). Figure 11 shows macroscopic pictures of glyceryl dibehenate-based tablets loaded with 30 % niacin before and after 24 h exposure to 0.1 N HCl at 37 °C. The tablets on the left hand side were prepared by direct compression, the tablets on the right hand side by hot-melt extrusion/grinding/compression. These pictures are representative for all other investigated formulations (data not shown). Clearly, the tablets remained intact during the observation period and did not swell. Upon exposure to the release medium the systems became more and more porous.

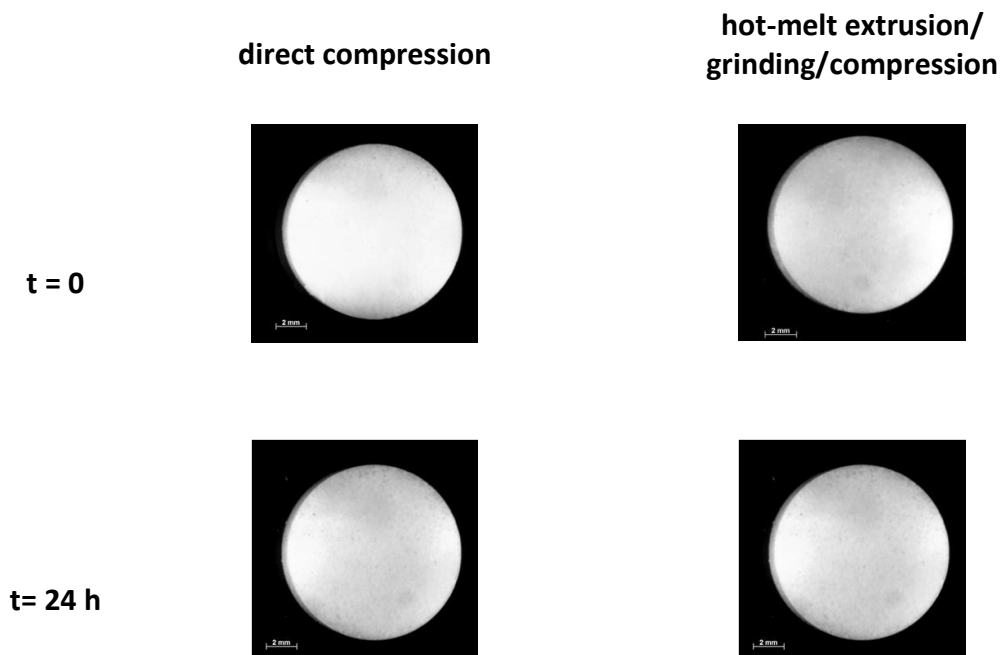


Figure 11: Macroscopic pictures of niacin-loaded, glyceryl dibehenate-based tablets prepared by direct compression or hot-melt extrusion/grinding/compression before and after 24 h exposure to the release medium (30 % initial vitamin loading).

The fact that the size and geometry of the tablets did not change during vitamin release is very important information, allowing for a significant simplification of the mathematical description of mass transport: The lengths of the pathways to be overcome remain unaltered during vitamin release, and various symmetries can be considered. The symbols in Figures 12 and 13 illustrate the experimentally determined niacin release kinetics from glyceryl dibehenate-based tablets prepared by direct compression (figure 12) or hot-melt extrusion/grinding/compression (figure 13). The diagrams on the left hand side show the *relative* vitamin release rates, the diagrams on the right hand side the respective *absolute* niacin release rates. Clearly, the slope of the release curves continuously decreases with time (vitamin release slows down), irrespective of the type of preparation method. Comparing figures 12 and 13, it becomes evident that niacin release from tablets prepared by hot-melt extrusion/grinding/compression was much slower than from tablets of identical composition, but prepared by direct compression, irrespective of the initial vitamin loading (note the different scaling of the x-axes). For both types of preparation methods the relative

and absolute release rate increased with increasing initial niacin content (figures 12 and 13). The solubility of niacin in the release medium at 37 °C was determined to be equal to 32.3 mg/mL, perfect sink conditions were maintained throughout all release experiments

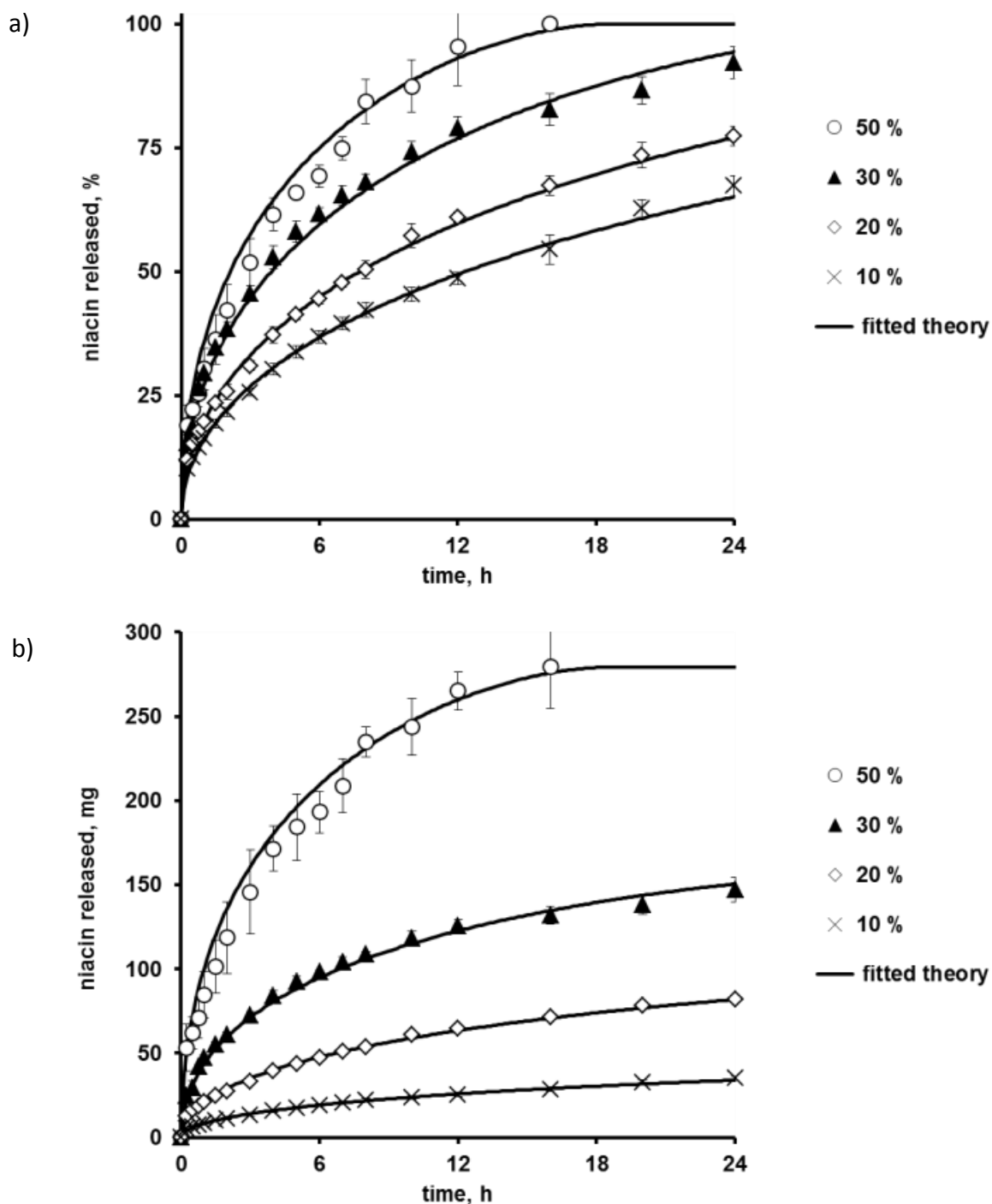


Figure 12: Effects of the initial niacin loading (indicated in the diagrams) on vitamin release from glyceryl dibehenate-based tablets prepared by direct compression on the resulting: (a) relative vitamin release rates, and (b) absolute vitamin release rates. The symbols represent the experimental results, the curves the fitted theory (Eqs. 1-6).

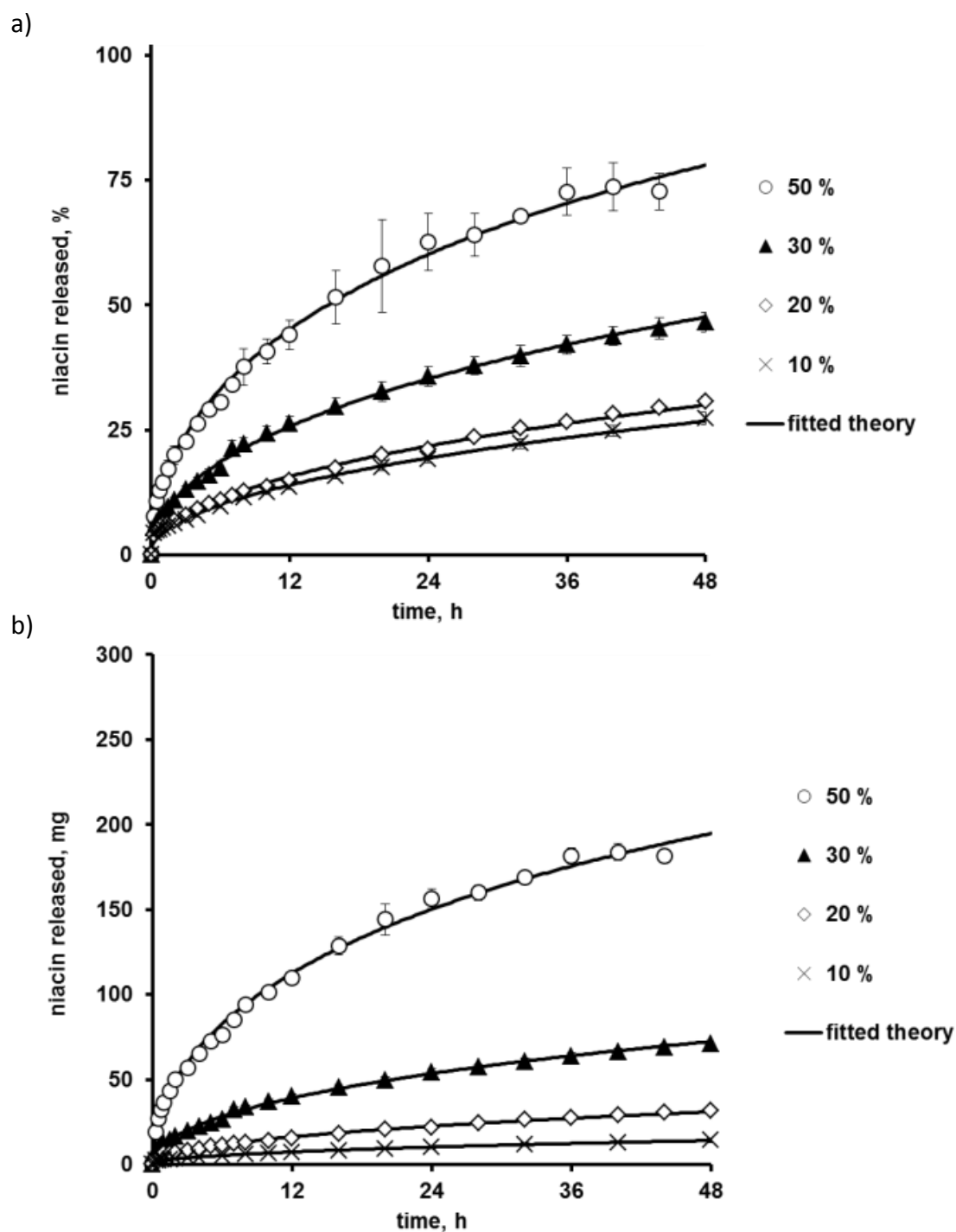


Figure 13: Effects of the initial niacin loading (indicated in the diagrams) on vitamin release from glyceryl dibehenate-based tablets prepared by hot-melt extrusion/grinding/compression on the resulting: (a) relative vitamin release rates, and (b) absolute vitamin release rates. The symbols represent the experimental results, the curves the fitted theory (Eqs. 1-6).

Based on these experimental results, the following model assumptions were defined:

- The tablets are cylindrical in shape, mass transport takes place in axial and radial direction.
- The tablet dimensions remain unaltered during vitamin release (stationary boundary conditions).
- Niacin dissolved in the bulk fluid does not hinder the release of vitamin remaining in the lipid tablets.
- The diffusion of niacin within the lipid tablets (through water-filled pores) is the dominant mass transport step, and this process is affected by the limited solubility of the vitamin in the low amounts of water present within the tablets. The rationale for this hypothesis is as follows: Upon exposure to the bulk fluid, three phenomena occur in a sequence: (1) The release medium penetrates into the system. (2) The vitamin particles dissolve in this liquid. (3) Dissolved niacin diffuses out of the lipid matrices, through water-filled channels and pores (since the lipid is in a crystalline state, vitamin diffusion through glyceryl dibehenate plates is highly unlikely). The process of vitamin dissolution (Siepmann, 2013) can be expected to be rapid compared to the two diffusional mass transport steps. In this study, the diffusion of the vitamin through the pores within the lipid matrix is assumed to be slower than water penetration into the system. Since the three processes occur in a sequence, and since niacin diffusion is considered to be the slowest mass transport step, the overall mass transport rate can be approximated by the diffusion rate of dissolved niacin through the porous lipid matrix (Siepmann, 2012). Importantly, the amount of water penetrating into the lipid matrix is very limited and the investigated niacin loadings are relatively high. Thus, not all of the vitamin can be expected to be immediately available for diffusion: Dissolved and non-dissolved niacin are considered to co-exist in the tablets.
- The mass transfer resistance due to the existence of a liquid, unstirred boundary layer surrounding the tablets is considered negligible compared to the mass transport resistance within the tablets.
- The vitamin and lipid powders are initially homogeneously distributed throughout

the tablets.

Based on these assumptions, a solution of Fick's second law of diffusion can be derived, considering the given initial and boundary conditions (this means the conditions given before exposure to the release medium and the conditions given at the edges of the tablets during niacin release). Since limited vitamin solubility within the matrix is considered (co-existence of dissolved and non-dissolved niacin), the model could not be solved analytically, but was solved numerically. For details, the reader is referred to the literature (Crank, 1975 ; Smith, 1985), and only the key equations are briefly described in the following.

Fick's law considering axial and radial diffusion in cylinders can be written as (Crank, 1975):

$$\frac{\partial c}{\partial t} = \frac{1}{r} \cdot \left\{ \frac{\partial}{\partial r} \left(r \cdot D \cdot \frac{\partial c}{\partial r} \right) + \frac{\partial}{\partial \theta} \left(\frac{D}{r} \cdot \frac{\partial c}{\partial \theta} \right) + \frac{\partial}{\partial z} \left(r \cdot D \cdot \frac{\partial c}{\partial z} \right) \right\} \quad (1)$$

where c is the concentration of *dissolved* niacin, which is dependent on time and position within the cylinder; t represents time; r , z denote the radial and axial coordinates and θ the angle perpendicular to the r - z -plane; D is the apparent diffusion coefficient of the vitamin within the tablet. The center of the cylinder is located at $r = 0$, $z = 0$ and $\theta = 0$. The tablet radius is R , the tablet half-height is Z . To reduce the calculation time, several system symmetries were considered, namely the symmetry at the $z = 0$ plane and the rotational symmetry of the tablets around the z -axis. Thus, it is sufficient to calculate the mass transport in only one quarter of a vertical cross-section of the tablets going through the center: Upon rotation around the z -axis, the mass transport in half of the cylinder can be calculated and due to the $z = 0$ symmetry plane, vitamin diffusion in the entire tablet can be calculated.

The initial condition expresses the fact that the niacin concentration within the tablets is homogeneous throughout the system and equal to its initial concentration, c_{ini} :

$$t=0 \quad c = c_{ini} \quad 0 \leq r \leq R \quad 0 \leq z \leq Z \quad (2)$$

The perfect sink boundary conditions read as follows:

$$t > 0 \quad c = 0 \quad 0 \leq r \leq R \quad z = Z \quad (3)$$

$$t > 0 \quad c = 0 \quad 0 \leq z \leq Z \quad r = R \quad (4)$$

Due to the symmetries at $z = 0$ and $r = 0$, there are no concentration gradients at $z = 0$ for $0 \leq r \leq R$, and at $r = 0$ for $0 \leq z \leq Z$:

$$t > 0 \quad \frac{\partial c}{\partial z} = 0 \quad 0 \leq r \leq R \quad z = 0 \quad (5)$$

$$t > 0 \quad \frac{\partial c}{\partial r} = 0 \quad 0 \leq z \leq Z \quad r = 0 \quad (6)$$

In order to calculate the niacin concentration at each time point and every position in the tablets, the above described 2-dimensional rectangle was divided into small pixels, introducing I and J space intervals, Δr and Δz , and generating a grid of $(I+1) \times (J+1)$ grid points. The release time was divided into g time intervals Δt (for most of the simulations $I = J = 50$ and $g = 500,000$ were chosen). At $t = 0$ (before exposure to the release medium), the niacin concentration was known from the initial vitamin loading and the system's dimensions. Using Eqs. 1-6, the concentration profiles of niacin at a new time step ($t = 0 + \Delta t$) was calculated, based on the concentration profiles at the previous time step ($t = 0$): The concentration at a certain *inner* grid point ($ix\Delta r, jx\Delta z$) at a new time step ($t = t_0 + \Delta t$) was calculated from its concentrations at the same grid point ($ix\Delta r, jx\Delta z$) and the four direct neighbors [$(i-1)x\Delta r, jx\Delta z$; $ix\Delta r, (j-1)x\Delta z$; $ix\Delta r, (j+1)x\Delta z$; $(i+1)x\Delta r, jx\Delta z$] at the previous time step ($t = 0$). The respective concentrations at *outer* grid points ($i = 0 \vee i = I \vee j = 0 \vee j = J$) at a new time step ($t = 0 + \Delta t$) were calculated using the boundary conditions (Eqs. 3-6). As the initial niacin concentration was known, the concentration profiles at $t = 0 + \Delta t, t = 0 + 2\Delta t,$

$t = 0 + 3\Delta t, \dots, t = 0 + g\Delta t$ could be calculated sequentially. Importantly, at each time point the vitamin concentration at a given position is compared to the niacin solubility: If the latter is exceeded, the excess amount of vitamin is considered to be non-dissolved and not available for diffusion. If the vitamin concentration is equal to or below niacin solubility, all of the niacin is considered to be available for diffusion at this position and this time point. The determined apparent diffusion coefficients are time- and position-averaged parameters. For

the implementation of the mathematical model the programming language C++ was used (Code::Blocks 10.05).

4.2.2. Model fittings to experimental results

The curves in figures. 12 and 13 show the fittings of the above described mathematical model to the experimentally determined niacin release kinetics from the different types of lipid tablets. It must be pointed out that the term “*fitting*” means that at least one model parameter was adjusted to minimize the resulting differences between theory and experiment. In the present study, only 1 parameter was initially unknown and adjusted by model fitting: the apparent diffusion coefficient of the vitamin in the lipid matrix. As is can be seen, good agreement between theory and experiment was obtained in all cases, irrespective of the initial niacin loading and type of preparation method (curves and symbols in figures. 12 and 13). This can serve as an indication for the fact that the hypothesized model assumptions might indeed be valid. However, caution must be paid: The observed good agreement is not a real proof for the validity of the model, since these are model *fittings*.

Based on these calculations, the apparent diffusion coefficient of niacin in the glyceryl dibehenate matrices could be determined. The black bars in figure 14 show the values obtained with tablets prepared by direct compression, the white bars the respective values for tablets prepared via hot-melt extrusion/grinding/compression. Two tendencies are clearly visible: (i) With increasing initial vitamin content, the apparent niacin mobility within matrices significantly increases, irrespective of the type of preparation technique. (ii) Niacin mobility in tablets prepared by direct compression is much higher than in tablets of identical composition, but prepared via hot-melt extrusion/grinding/compression. These tendencies are consistent with the vitamin release kinetics illustrated in figures. 12 and 13. The substantial increase in vitamin mobility with increasing initial niacin loading can be attributed to the fact that the lipid matrices become more and more porous upon vitamin exhaust with increasing initial vitamin content: As discussed above, the outer dimensions of the lipid tablets remained unaltered during niacin release. Thus, positions initially occupied by vitamin particles become occupied by water once niacin is released. The higher the initial

vitamin content, the higher the resulting porosity. Consequently, the vitamin mobility in the lipid matrices increases with increasing initial niacin loading. Note that the D values are time- and position-averaged parameters. Importantly, the following quantitative empirical relationships could be established between the apparent niacin diffusivity in the lipid tablets and the initial vitamin content:

In the case of tablets prepared by direct compression:

$$D(\% \text{ niacin}) = 0.2224 \cdot (\% \text{ niacin})^{2.016} \cdot 10^{-6} \text{ cm}^2/\text{s} \quad (7)$$

In the case of tablets prepared via hot-melt extrusion/grinding/compression:

$$D(\% \text{ niacin}) = [0.0665 \cdot (\% \text{ niacin})^2 - 0.196 \cdot (\% \text{ niacin}) + 0.1494] \cdot 10^{-6} \text{ cm}^2/\text{s} \quad (8)$$

These equations allow calculating the vitamin diffusivities for arbitrary initial niacin loadings for both types of preparation techniques.

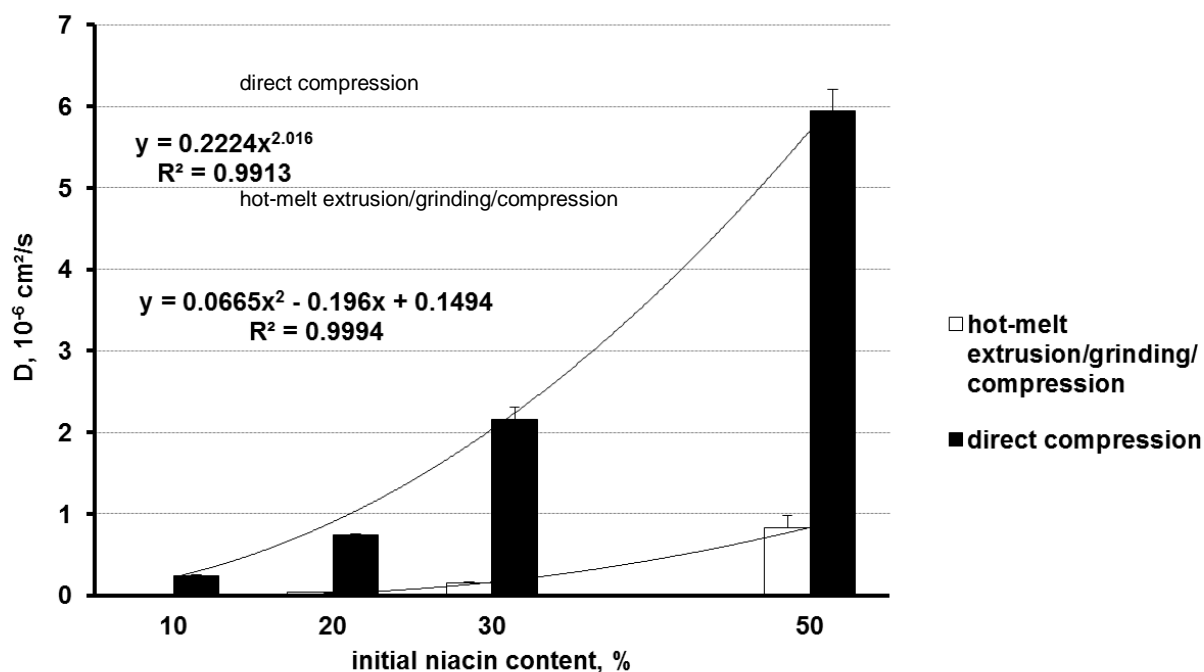


Figure 14: Dependence of the apparent niacin diffusion coefficient in glyceryl dibehenate-based tablets on the type of preparation technique and initial vitamin content.

The fact that vitamin mobility is much higher in lipid tablets prepared by direct compression than in tablets prepared via hot-melt extrusion/grinding/compression can be explained as follows: During hot-melt extrusion a much more intense embedding of the niacin particles with lipid mass can be expected, since the temperature is elevated, high shear forces are acting and the vitamin and lipid are intensively mixed. Consequently, the geometry and size of the pores through which the niacin has to diffuse (once dissolved) is likely to be substantially different (e.g., increased tortuosity and length of the pathways to be overcome).

4.2 3. Deeper insight into vitamin release mechanisms

Based on these observations and calculations, the relative importance of “niacin diffusion through the porous lipid matrix” versus “limited niacin solubility within the tablets” can be estimated. Knowing the apparent diffusion coefficient of the vitamin in the tablets, Eqs. 1-6 can be used to calculate the resulting niacin concentration distance profiles within the systems at any time point. The scheme in figure 15 illustrates the view points used for the presentation of such concentration distance profiles shown in figure 16. A cross-section is made through the center of a cylindrical tablet, perpendicular to the circular surfaces. The diagrams in figure 16 show the 3-dimensional total niacin concentration distance profiles calculated after 6, 12 and 24 h exposure to the release medium. The view point for these diagrams is illustrated by the solid eye in figure 5. Tablets prepared by direct compression are shown on the left hand side, tablets prepared via hot-melt extrusion/grinding/compression are illustrated on the right hand side. The total vitamin concentration is plotted in mg/mL on the vertical axes, the radial and axial positions are plotted on the other two axes, as indicated in figure 15. The total vitamin concentration at a specific position is the sum of the concentration of *dissolved* niacin (being available for diffusion) and *non-dissolved* niacin (not being available for diffusion). At the top of each diagram, a 2-dimensional projection of the 3-dimensional concentration distance profiles is given. The view point for these projections is illustrated by the dotted eye in figure 15.

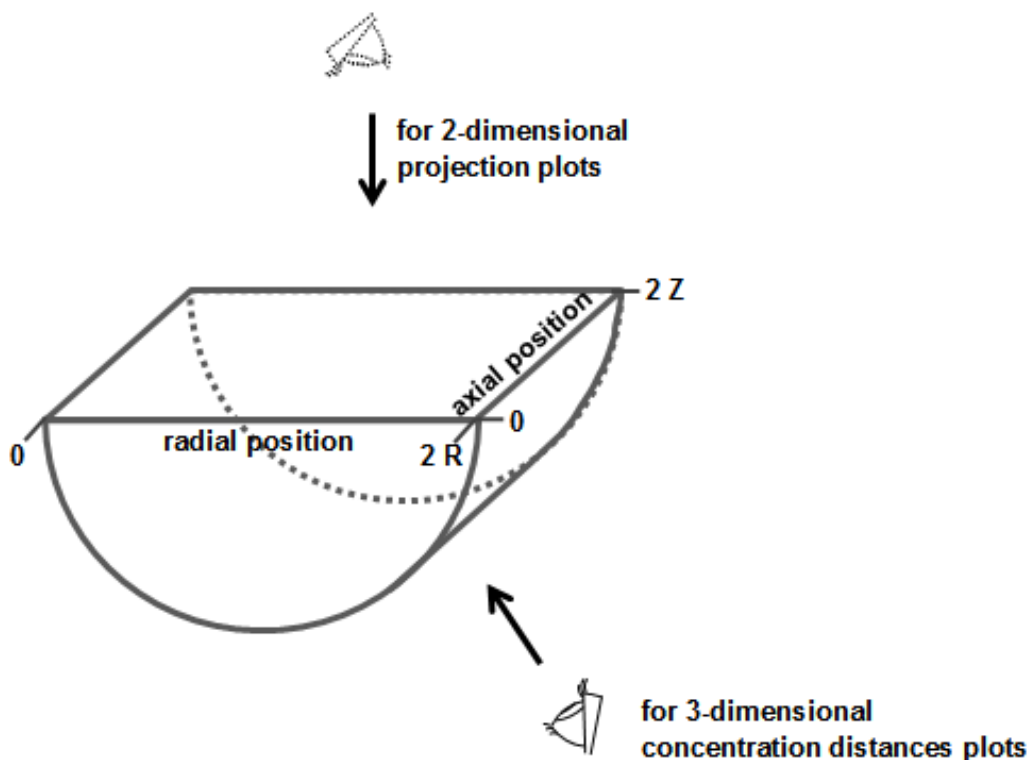


Figure 15: Schematic representation of the viewpoints used in Figure 6 to illustrate the time-dependent changes in the niacin concentration distance profiles within the glyceryl dibehenate-based tablets upon exposure to the release medium. The solid eye shows the viewpoint for the 3-dimensional concentration distance profiles, the dotted eye the viewpoint for the 2-dimensional projections shown at the top of each diagram in figure 16.

Clearly, extremely steep concentration gradients exist within the tablets, irrespective of the selected time point and type of preparation technique. Zooms on the lower concentration ranges of these diagrams are shown in figure 17. The solubility of niacin in the release medium was determined to be 32.3 mg/mL.

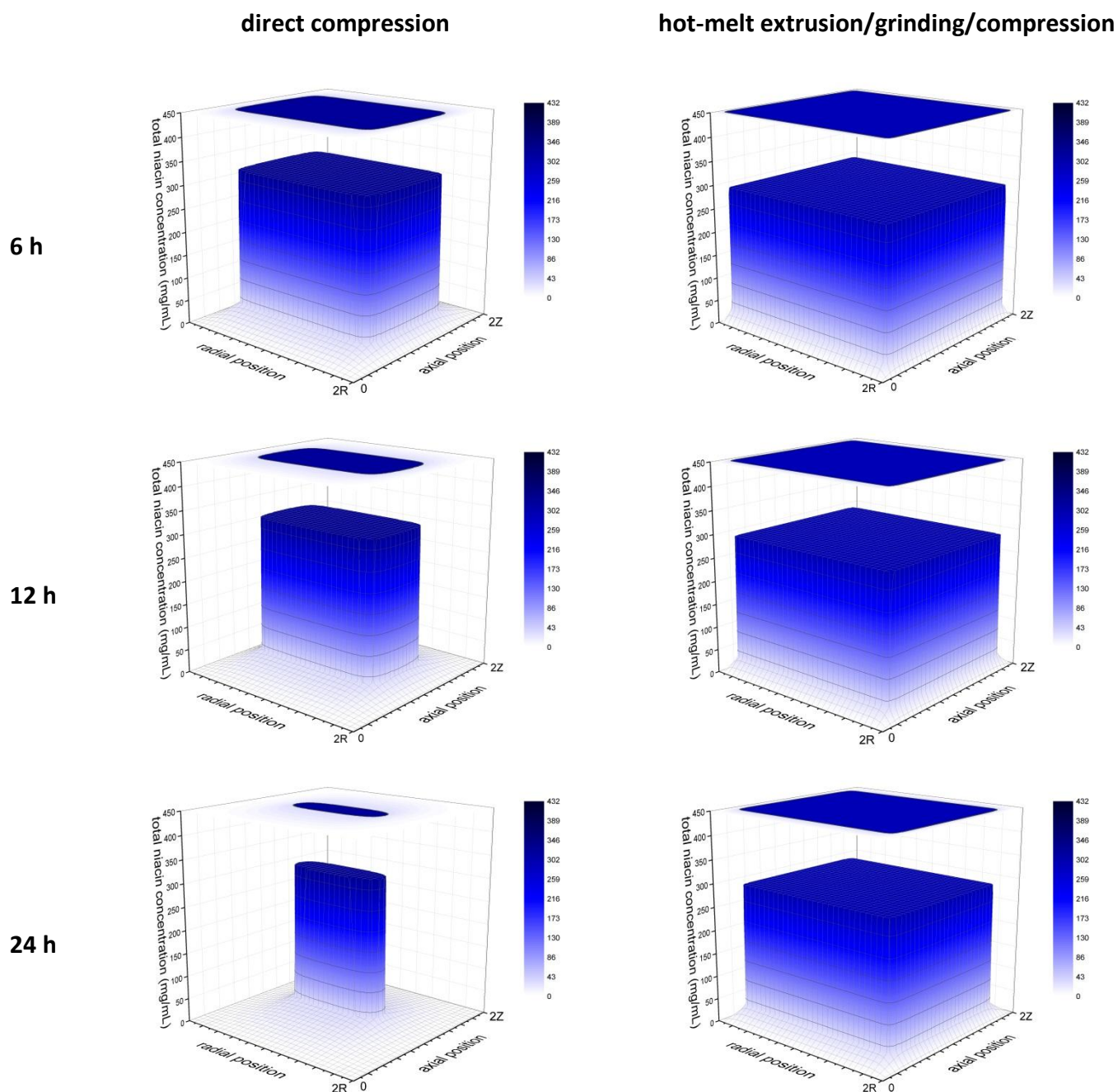


Figure 16: Dynamic changes in the total niacin concentration distance profiles within glyceryl dibehenate-based tablets prepared by direct compression (left hand side) or hot-melt extrusion/grinding/compression (right hand side) upon exposure to the release medium. The initial vitamin content was 30 %. The view points are schematically illustrated in figure 15. At the top of each diagram, a 2-dimensional projection of the respective concentration profile is shown. The color scales are given on the right hand side. The exposure times to the release medium are indicated on the left hand side. Zooms on lower vitamin concentrations are shown in figure 17.

Looking at figures 16 and 17 it becomes obvious that the limited amount of water available in the lipid tablets can only dissolve a small part of the vitamin present in the system. Most of the niacin is not dissolved and not available for diffusion. The front separating the zone of the tablet containing both, dissolved and non-dissolved vitamin, from the zone of the tablet containing only dissolved niacin can be called "dissolution front". This front steadily moves inwards, towards the center of the tablets. Importantly, net diffusional mass transport of niacin in the tablets only takes place between this dissolution front and the edges of the tablets. Within the saturated center, the concentration of dissolved niacin is always equal to the solubility of the vitamin. Thus, there is no concentration gradient of dissolved vitamin and, hence, no net mass flux. The concentration gradients of dissolved niacin illustrated in figure 17 clearly show that with time the driving forces for niacin release decrease: The lengths of the diffusion pathways monotonically increase, while the concentration difference remains unaltered (saturated solution versus perfect sink). This is consistent with the observed monotonically decreasing niacin release rates shown in figures 12 and 13. Comparing the left and right hand sides of figures. 16 and 17, it becomes obvious that the dissolution front movement is much more rapid in tablets prepared by direct compression than in tablets prepared via hot-melt extrusion/grinding/compression. This is due to the much higher niacin mobility in these systems, as discussed above (see also figure. 14). Interestingly, close to the center of the tablets a large excess of niacin still exists at late time points, when vitamin release is almost complete. In contrast, most of the niacin is released from the surface-near regions of the system. Note that figures 16 and 17 only show a few examples of concentration distance profiles in tablets with an initial vitamin loading of 30 %, but the tendencies are similar in all other systems. With increasing initial vitamin loading, the rate at which the dissolution front moves towards the center of the tablet increases, irrespective of the type of preparation technique.

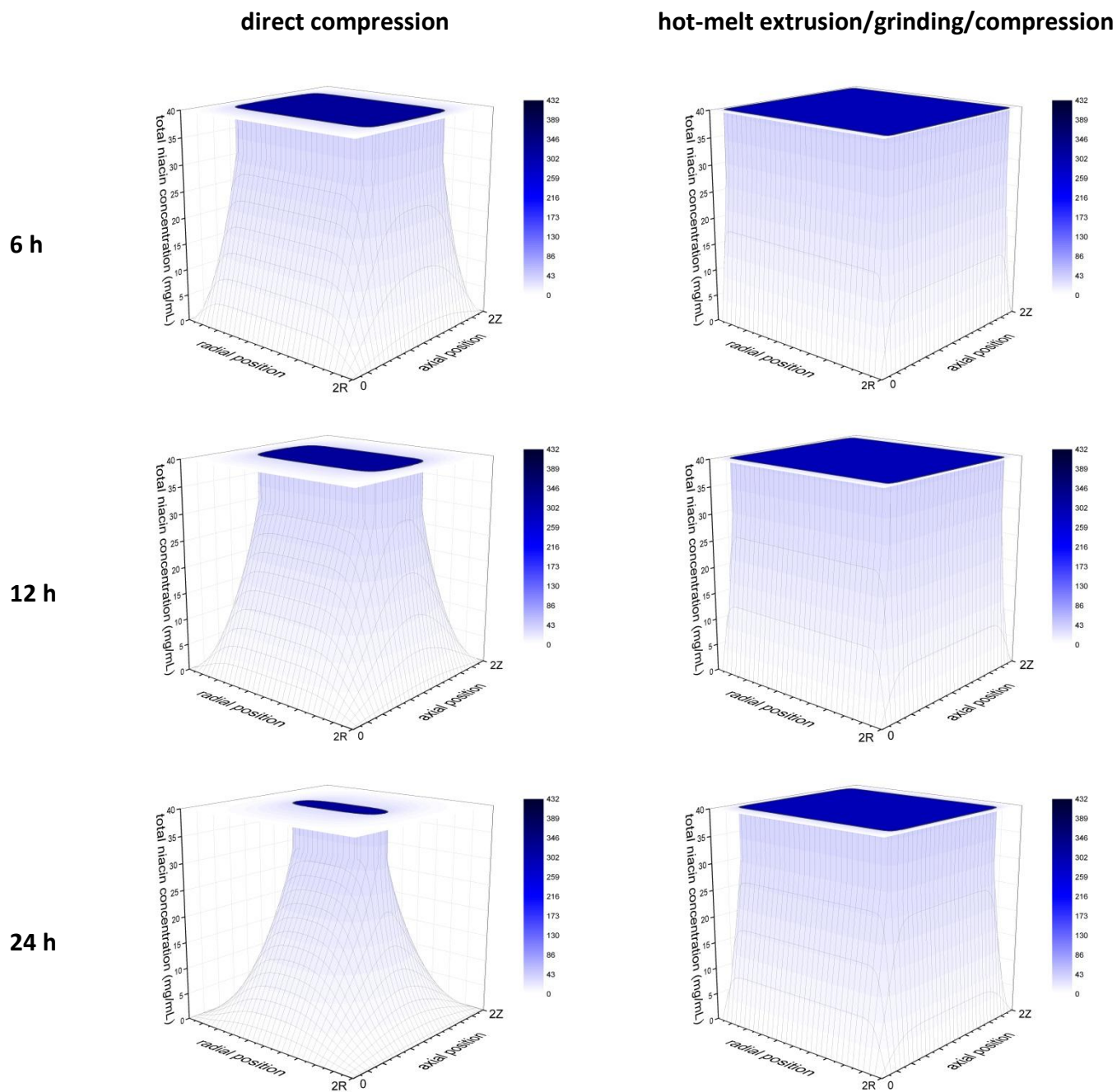


Figure 17: Zooms on the lower niacin concentrations of the diagrams shown in figure 16.

A major key message from these *in-silico* simulations is that limited vitamin solubility is of utmost importance in the investigated systems (in addition to niacin diffusion through water-filled pores in the lipid matrix), and net vitamin transport does not occur *all over* the tablets at the same time, but is restricted to specific, well-defined regions during major parts of the release period.

4.2.4. *Model predictions and independent experiments*

As pointed out above, obtaining good agreement between a mathematical theory and experimental results when *fitting* the model to the experimental data, is not a real proof for the validity of the theory. A much more reliable evaluation of the validity of a mathematical theory is based on the comparison of quantitative model predictions (e.g., concerning the impact of a formulation or processing parameter on the resulting system properties) with *independent* experiments (experiments, which were only performed after the *in-silico* simulations were made). In these cases, deviations between theory and experiment are not minimized by a fitting procedure. Also, in practice this type of *in-silico* simulations (quantitative predictions of the impact of formulation and processing parameters) is highly valuable, allowing for the substitution of time-intensive and costly series of experimental studies by rapid computer calculations.

The dashed curves in figure 8 show the theoretically predicted release kinetics of niacin from glyceryl dibehenate-based tablets initially containing 40 % vitamin, with a diameter of 11.3 mm and a height of 4.6 mm. Eqs. 1-6 were used for these calculations, the D values were estimated from Eqs. 7-8. As expected, the theory predicts a much higher vitamin release rate from tablets prepared by direct compression compared to systems prepared via hot-melt extrusion/grinding/compression (for the reasons discussed above) as well as monotonically decreasing release rates, irrespective of the type of preparation technique. To evaluate the validity of these predictions, the respective tablets were prepared in a second step and niacin release was measured experimentally (only after the predictions were made). These independent experimental results are shown as symbols in figure 18. Importantly, good agreement was obtained between the theoretical predictions and the independent experiments, irrespective of the type of preparation technique. This indicates that: (i) the theory is likely to be valid for the investigated lipid tablets, and (ii) *in-silico* simulations can help saving time and reducing costs during product development.

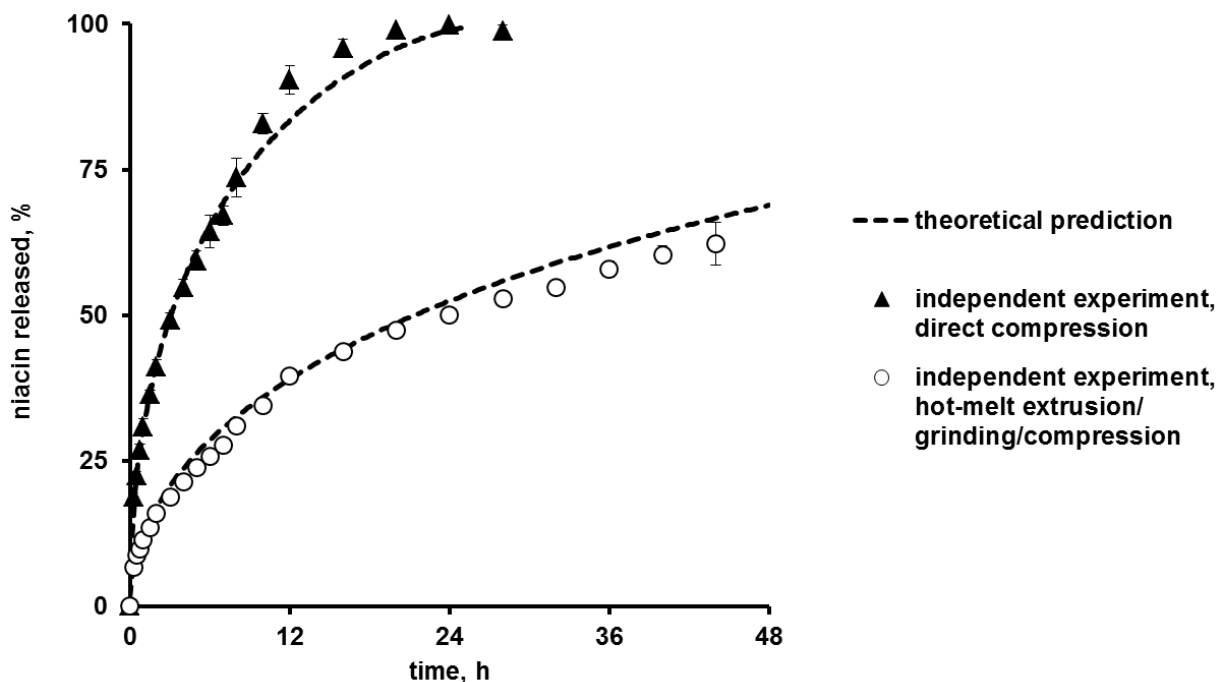


Figure 18: Theoretical predictions (dashed curves) and independent experiments (symbols): Impact of the tablet composition and type of manufacturing procedure on niacin release from glyceryl dibehenate-based tablets with an initial loading of 40 %, prepared by direct compression (upper curve) or hot-melt extrusion/grinding/compression (lower curve). The tablet diameter was 11.3 mm, the tablet height 4.6 mm.

Further examples for quantitative model predictions and independent experiments are illustrated in figure 19: The dashed curves in the diagram on the left hand side represent the theoretically predicted impact of varying the tablet *height* of glyceryl dibehenate-based tablets from 2.5 to 7.5 mm, while keeping the tablet diameter constant at 11.3 mm. The initial vitamin loading was 35 %, the tablets were prepared by direct compression in these cases. Again, Eqs. 1-6 were used for the *in-silico* simulations and Eq. 7 for the estimation of the apparent niacin diffusion coefficient. Clearly, the model predicts decreasing relative vitamin release rates with increasing tablet height, which can be attributed to the decreasing surface area:volume ratio. Importantly, independent experiments confirmed all model predictions (symbols versus dashed curves in figure 19a). The theoretically predicted impact of varying the *diameter* of glyceryl dibehenate-based tablets from 7.0 to 16.0 mm is illustrated by the dashed curves in figure 19b. In these cases, the tablet height was constant (5 mm), the initial vitamin loading was 25 %, and the systems were prepared by direct

compression. Again, a decrease in the relative release rate is predicted with increasing system dimension (due to the decreasing surface area:volume ratio). Interestingly, the expected decrease in the relative release rate is less pronounced compared to the decrease upon varying the tablet height shown in figure 19a. This can be explained by the less pronounced reduction in the surface area:volume ratio of the tablets in these cases: by 33 % in figure 19b compared to 46 % in figure 19a. Importantly, these predicted moderate effects on the resulting vitamin release rates were again confirmed by independent experiments (symbols in figure 19b). This further confirms the validity of the presented mathematical model and demonstrates its practical usefulness.

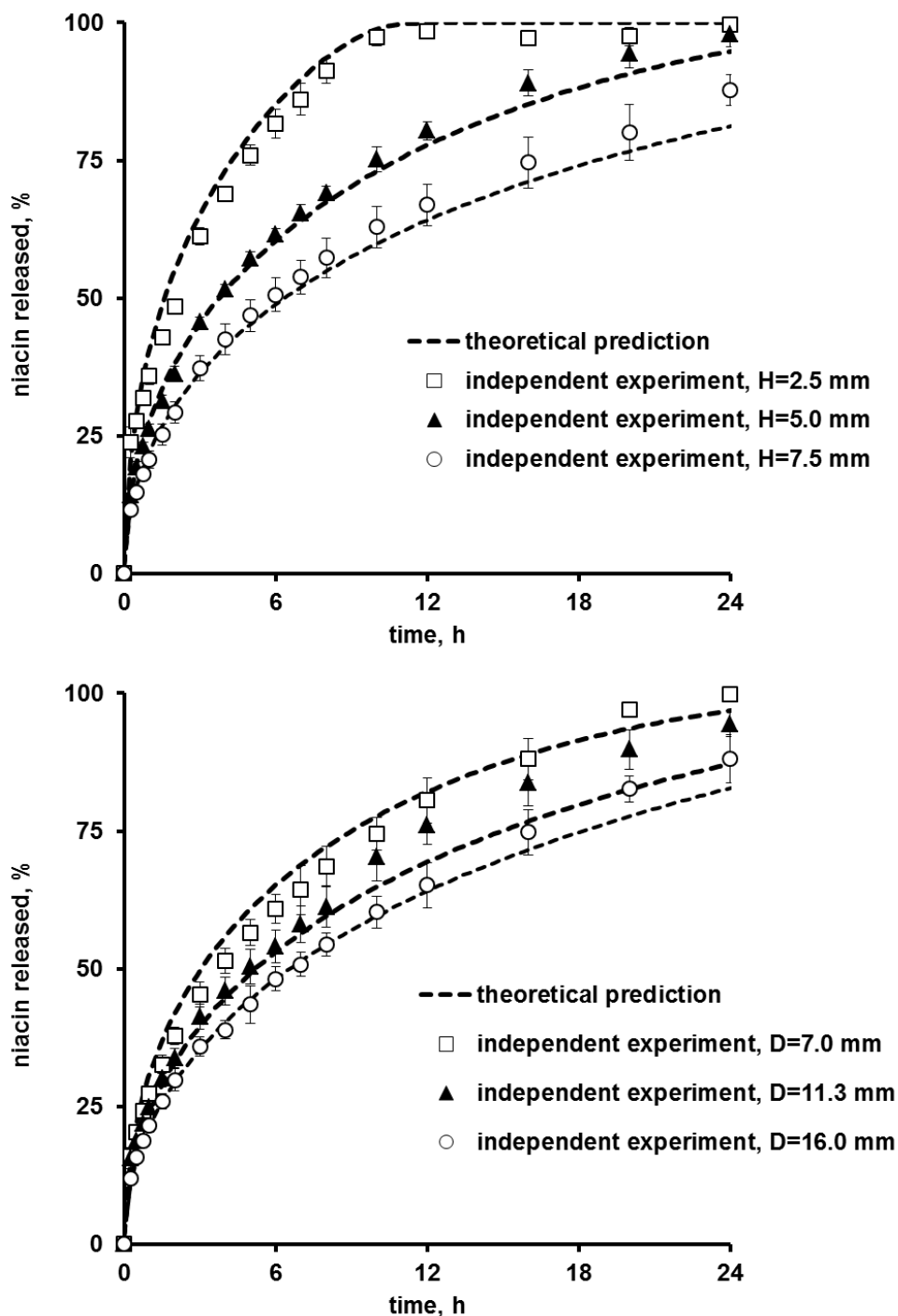


Figure 19: Theoretical predictions (dashed curves) and independent experiments (symbols): Impact of the tablet dimensions and composition on niacin release from glyceryl dibehenate-based tablets with: a) a height of 2.5, 5.0 or 7.5 mm, a constant diameter of 11.3 mm, and an initial vitamin loading of 35 %, and b) a diameter of 7.0, 11.3 or 16.0 mm, a constant height of 5 mm and an initial vitamin loading of 25 %. The tablets were prepared by direct compression.

4.3. Conclusion

In-silico simulations of lipid tablets using mechanistically realistic mathematical theories can help understanding how vitamin release is controlled and can very much facilitate the optimization of this type of advanced delivery systems.

**5. Investigation of the viscosity grade of guar gum
in polymer blends to overcome ethanol
sensitivity of ethylcellulose-based coated pellets.**

5.1. Materials and Methods

5.1.1. Materials

Theophylline matrix pellets (70 % drug content, diameter: 0.71-1.25 mm; FMC BioPolymer, Philadelphia, PA, USA); Ethylcellulose Aqueous Dispersion NF (Aquacoat[®] ECD 30; FMC BioPolymer); dibutyl sebacate (DBS; Morflex, Greensboro, NC, USA); ethanol (Fisher Bioblock Scientific, Illkirch, France); *low viscosity* guar gum (*low η* guar gum, apparent viscosity of a 1 % aqueous guar gum solution = 52 cPs; TIC Pretested Gum Guar TICOLV FCC Powder; TIC Gums); *medium viscosity* guar gum (*medium η* guar gum, apparent viscosity of a 1 % aqueous guar gum solution = 320 cPs; Polygum 240/80; Polygal Trading, Maerstetten, Switzerland). The apparent viscosities were measured using an AR2000Ex rheometer (TA Instruments, New Castle, DE, USA) at a shear rate of 50 s⁻¹ in a 1 % aqueous guar gum solution measured rotationally at 20 °C after 1 min equilibration using a 6 cm acrylic cone (1°), wherein the shear was ramped up linearly from 1 to 50 s⁻¹ in 25 steps over 29 s.

5.1.2. Preparation and characterization of thin polymeric films

Aquacoat[®] ECD 30 was plasticized for 1 d with 25 % DBS (w/w; based on the ethylcellulose mass). Guar gum (*medium η* guar gum in all cases) was dissolved in purified water at 65 °C (0.7 % or 1 % w/w, as indicated; 100 % reference value = total coating formulation; 2 h stirring) and cooled down to room temperature. The two liquids were blended and stirred for 30 min prior to use. Films were prepared by casting Aquacoat[®] ECD 30:guar gum blends onto Teflon plates and subsequent controlled drying for 24 h at 60 °C.

The dry mass loss kinetics and changes in the (water + ethanol) content of the films upon exposure to the release media were determined as follows: Pieces of 5 cm × 5 cm were placed into 100 mL plastic flasks filled with 100 mL pre-heated 0.1 M HCl:ethanol 60:40 (V/V), followed by horizontal shaking (37 °C, 80 rpm; GFL 3033, Gesellschaft fuer Labortechnik, Burgwedel, Germany). At pre-determined time points, samples were withdrawn, accurately weighed [wet mass (*t*)] and dried to constant mass at 60°C [dry mass

(*t*)]. The (water + ethanol) content (%) and dry film mass (%) at time *t* were calculated as follows:

$$(\text{water} + \text{ethanol}) \text{ content } (\%) (t) = \frac{\text{wet mass}(t) - \text{dry mass}(t)}{\text{wet mass}(t)} \cdot 100 \% \quad (1)$$

$$\text{dry film mass } (\%) (t) = \frac{\text{dry mass}(t)}{\text{dry mass}(t=0)} \cdot 100 \% \quad (2)$$

5.1.3. Pellet coating

Theophylline matrix cores were coated with Aquacoat[®] ECD 30:guar gum 85:15 blends (if not otherwise stated). Aquacoat[®] ECD 30 was plasticized for 1 d with 25 % DBS (w/w; based on the ethylcellulose mass). Guar gum (*medium* η guar gum, if not otherwise stated) was dissolved in purified water at 65 °C (0.7 % or 1 % w/w, as indicated; 100 % reference value = total coating formulation; 2 h stirring) and cooled down to room temperature. The two liquids were blended and stirred for 30 min prior to use. The coating dispersions were sprayed onto theophylline pellets using a fluidized bed coater (Strea 1, Wurster insert; Niro; Aeromatic-Fielder, Bubendorf, Switzerland). The process parameters were as follows: inlet temperature = 38 °C, product temperature = 38 ± 2 °C, spray rate = 2g/min, atomization pressure = 1.2 bar, nozzle diameter = 1.2 mm. After coating the pellets were further fluidized for 10 min and subsequently cured for 24 h at 60 °C in an oven.

5.1.4. Drug release measurements

Theophylline release from coated pellets was measured in 0.1 M HCl or 0.1 M HCl: ethanol 60:40 (V/V), followed by phosphate buffer pH 7.4 (USP 36) using the USP 36 paddle apparatus (Sotax, Basel, Switzerland) (900 mL, complete medium change after 2 h; 37 °C, 100 rpm, *n* = 3). At pre-determined time points, 3 mL samples were withdrawn and analyzed UV-spectrophotometrically [λ = 270.4 nm in 0.1 N HCl, λ = 272.2 nm in 0.1 M HCl:ethanol 60:40 (V/V) and phosphate buffer pH 7.4] (UV 1650 PC, Shimadzu, Champs-sur-Marne,

France). If indicated, the pellets were stored in open glass vials (no packing material) at ambient conditions (25 °C and 60 % relative humidity) or stress conditions (40 °C and 75 % relative humidity) prior to the measurements.

5.1.5. SEM studies

The morphology of coated pellets was characterized using a cold field emission high resolution scanning electron microscope (S-4700 Field Emission Gun; Hitachi, Hitachi High-Technologies Europe, Krefeld, Germany). Samples were covered under vacuum with a carbon layer.

5.2. Results and discussion

5.2.1. Ethylcellulose/guar gum ratio

Recently, ethanol-resistant drug release was reported from ethylcellulose:guar gum coated pellets (Rosiaux et al., *submitted*). The blend ratio in that study was 85:15 or 90:10, respectively. The presence of the ethanol insoluble guar gum effectively avoided undesired ethylcellulose dissolution in ethanol-rich media. However, so far it is unclear what the minimal guar gum content in the film coating is, required to provide ethanol-resistant drug release kinetics.

In the present study, ethylcellulose was blended in various ratios with *medium* μ guar gum and used to coat theophylline matrix cores. Figure 20 shows the resulting drug release kinetics in: 0.1 M HCl for 2 h, followed by phosphate buffer pH 7.4 for 6 h (open symbols); or 0.1 M HCl:ethanol 60:40 for 2 h, followed by phosphate buffer pH 7.4 for 6 h (filled symbols). Firstly, the drug release rate generally increased with increasing guar gum content, irrespective of the type of release medium. This can be attributed to the higher permeability of guar gum compared to ethylcellulose for the drug. Secondly, it can be seen that at low to very low guar gum contents, the difference in drug release observed between ethanol-containing and ethanol-free release media increased. This is likely due to the fact that too small amounts of guar gum are not able to effectively prevent undesired ethylcellulose dissolution in ethanol-rich media. Consequently, the permeability of the film coating increases and drug release is accelerated in the presence of 40 % ethanol. The term “ethanol-resistant” drug release might be defined as follows: A solid dosage form is called “ethanol-resistant” (or “not sensitive to ethanol”), if the *in vitro* drug release data in 0.1 M HCl is compared with and without 40% ethanol for 2 h at 37 °C and the difference throughout the 2 h period in release profiles between the ethanol-free media and ethanol-containing media is: i) < 15 %, when less than 20% of the drug is released in the ethanol-free media, or ii) < 30 % when more than 20 % of the drug is released in ethanol-free media. According to this definition all pellets coated with ethylcellulose:guar gum blends containing more than 5 % guar gum exhibited ethanol-resistant drug release (Figure 20).

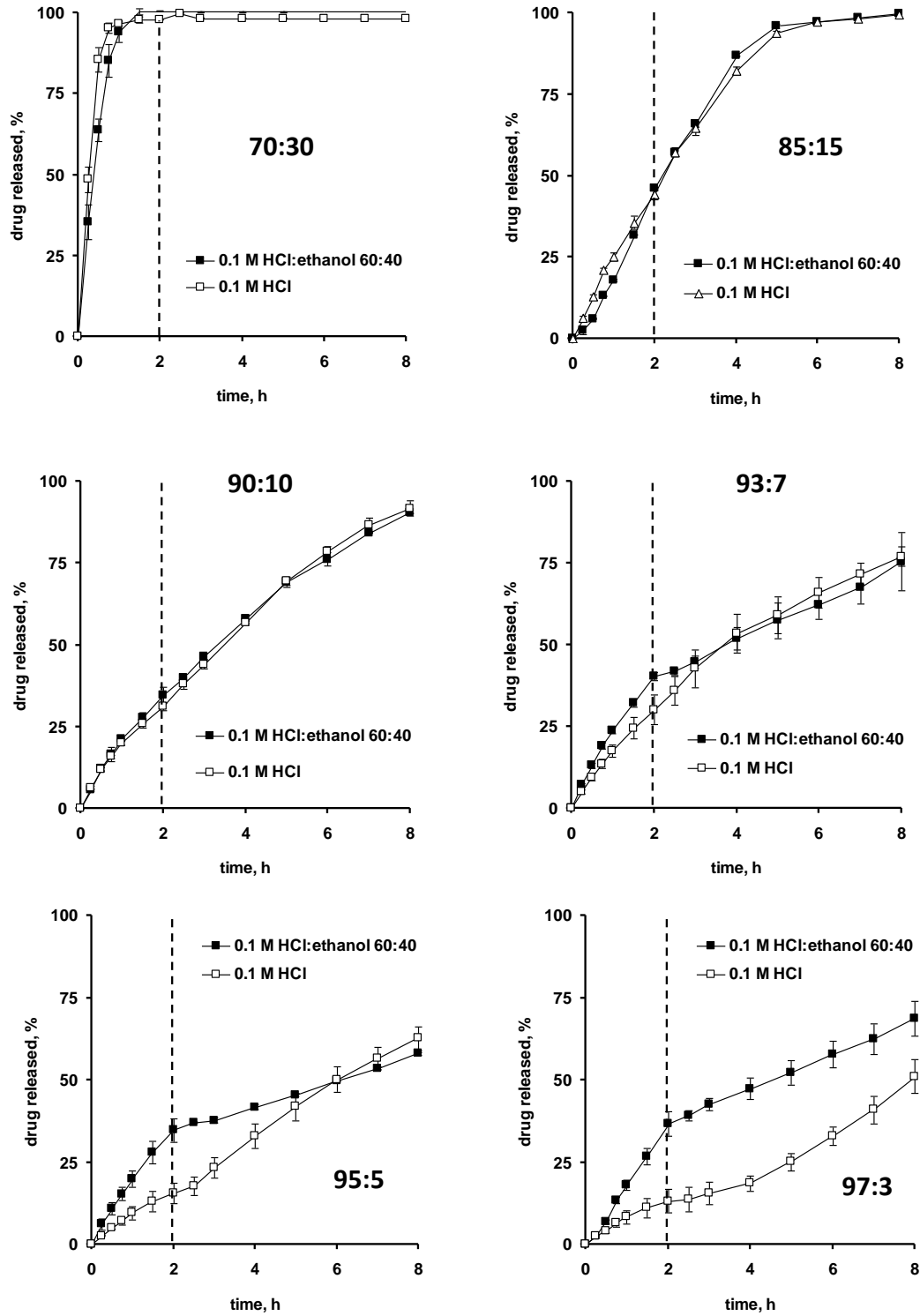


Figure 20. Impact of the ethylcellulose:guar gum blend ratio (indicated in the diagrams) on the sensitivity of drug release to ethanol: Theophylline release from pellets (15 % coating level) in 0.1 M HCl for 2 h, followed by phosphate buffer pH 7.4 (open symbols) or 0.1 M HCl:ethanol 60:40 for 2 h, followed by phosphate buffer pH 7.4 (filled symbols).

These findings are consistent with the experimentally measured dry mass loss kinetics of thin, free films of identical composition as the investigated film coatings. Film pieces of 5 cm x 5 cm were exposed to 0.1 M HCl:ethanol 60:40 (in horizontally shaken flasks at 80 rpm and 37 °C). As it can be seen in Figure 21a, the dry mass loss of the systems containing 7, 10 or 15 % guar gum (referred to the total polymer mass) was very similar: The presence of the guar gum effectively prevented undesired ethylcellulose dissolution even at high ethanol contents of the release medium in all cases. Figure 21b shows the experimentally measured (water + ethanol) contents of these films upon exposure to 0.1 M HCl:ethanol 60:40 at 37 °C. As it can be seen, there is a slight increase in the liquid uptake of the films with increasing guar gum content. This is probably due to a higher uptake of water by the more hydrophilic guar gum compared to ethylcellulose. The observed tendency is also consistent with the above discussed effects on drug release (irrespective of the presence/absence of ethanol in the bulk fluid): The release rate generally increased with increasing guar gum content, which can at least partially be attributed to increased water uptake, resulting in higher macromolecular mobility and, thus, higher permeability of the film for the drug.

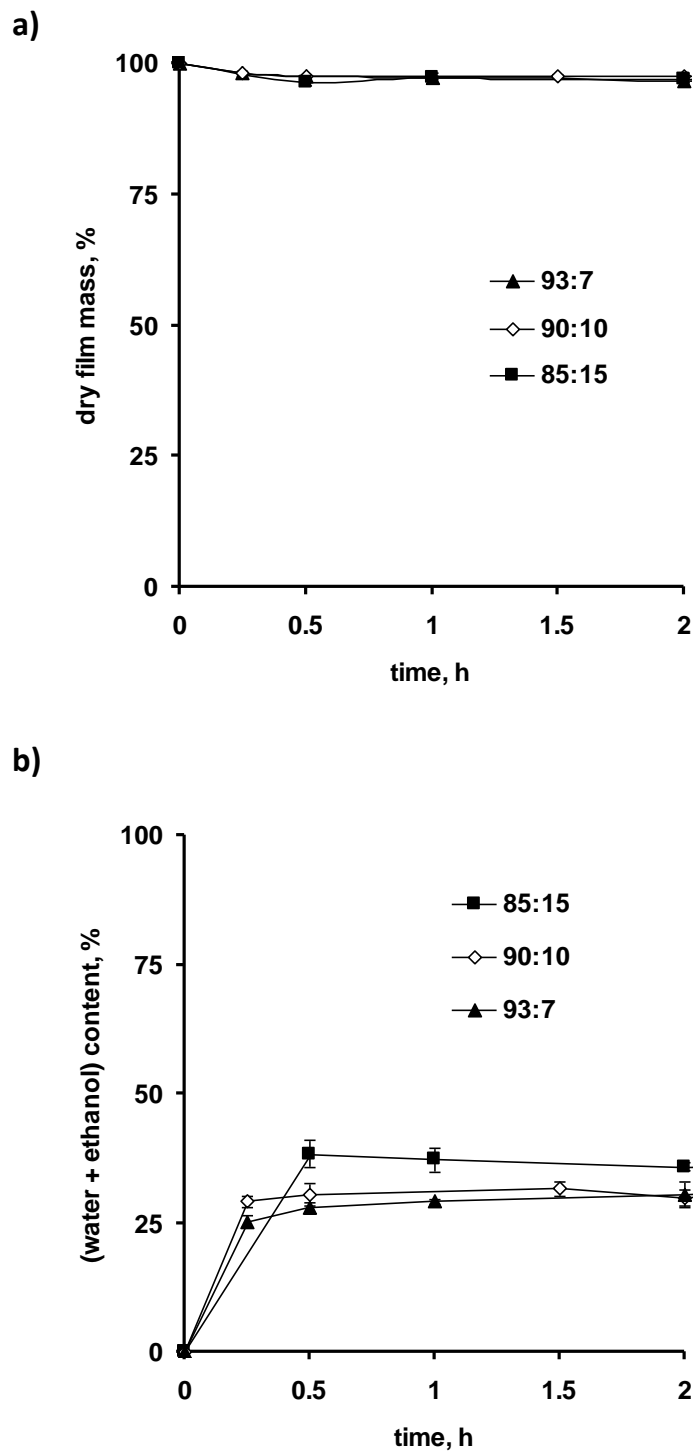


Figure 21: Effects of the ethylcellulose:guar gum blend ratio (indicated in the diagrams) on changes in the: a) dry mass, and b) (water + ethanol) content of thin, free films upon exposure to 0.1 M HCl:ethanol 60:40.

Figure 22 shows the morphology of theophylline matrix pellets, coated with 93:7, 90:10, or 85:15 ethylcellulose: *medium* η guar gum blends (as indicated). The coating level was 20%. As it can be seen, in all cases a continuous polymer coating was achieved, without any sign for cracks. Thus, the coating composition did not significantly affect the outer morphology of the coated pellets in the investigated range.

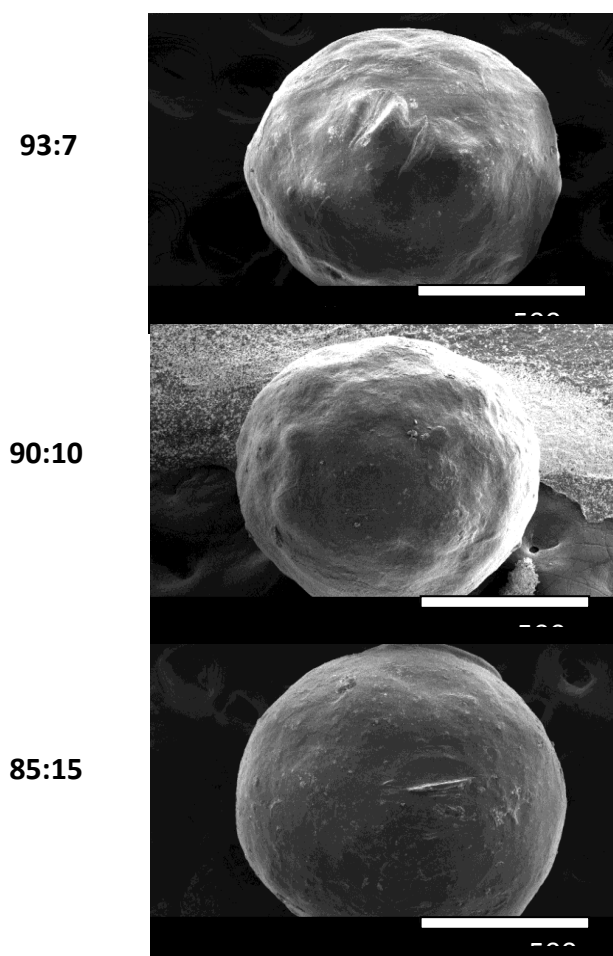


Figure 22: SEM pictures of theophylline pellets coated with different ethylcellulose:guar gum blends (the ratio is indicated on the left hand side) (before exposure to the release medium, 20 % coating level).

5.2.2. Guar gum concentration in the total dispersion

From a processing point of view the viscosity of the coating formulation, which is sprayed onto the pellets, is of major importance. If the viscosity is too high, extensive sticking can result. In contrast, too low viscosities can lead to very long processing times. However, so far it is unknown whether the concentration of the guar gum in the coating formulation (affecting the latter's viscosity) does or does not impact the ethanol-resistance of the coated pellets. In this study, two guar gum concentrations in the final coating dispersion were compared: 0.7 and 1 % (w/w). The 100 % reference value is the total mass of the coating dispersion. Figure 23 shows the experimentally measured drug release kinetics from theophylline loaded pellets coated with 90:10 (top row), or 85:15 (bottom row) ethylcellulose:guar gum blends. In the left column, the guar gum concentration was 0.7 %, in the right column 1 %. The open symbols indicate drug release in 0.1 M HCl for 2 h, followed by phosphate buffer pH 7.4 for 6 h. The filled symbols show drug release in 0.1 M HCl:ethanol 60:40 for 2 h, followed by phosphate buffer pH 7.4 for 6 h. Clearly, ethanol-resistance was not affected by the investigated guar gum concentration in the coating formulation: In all cases, the difference in theophylline release in media free of ethanol versus media containing 40 % ethanol was only minor. Comparing the release profiles on the left and on the right hand side, it can be seen that an increase in the guar gum concentration in the coating dispersion (from 0.7 to 1 %) led to a slight decrease in the resulting theophylline release rate, irrespective of the ethylcellulose:guar gum blend ratio. This slight effect might be attributable to a somewhat denser polymer network structure, resulting from a higher polymer concentration in the liquid deposited on the pellets' surface during coating.

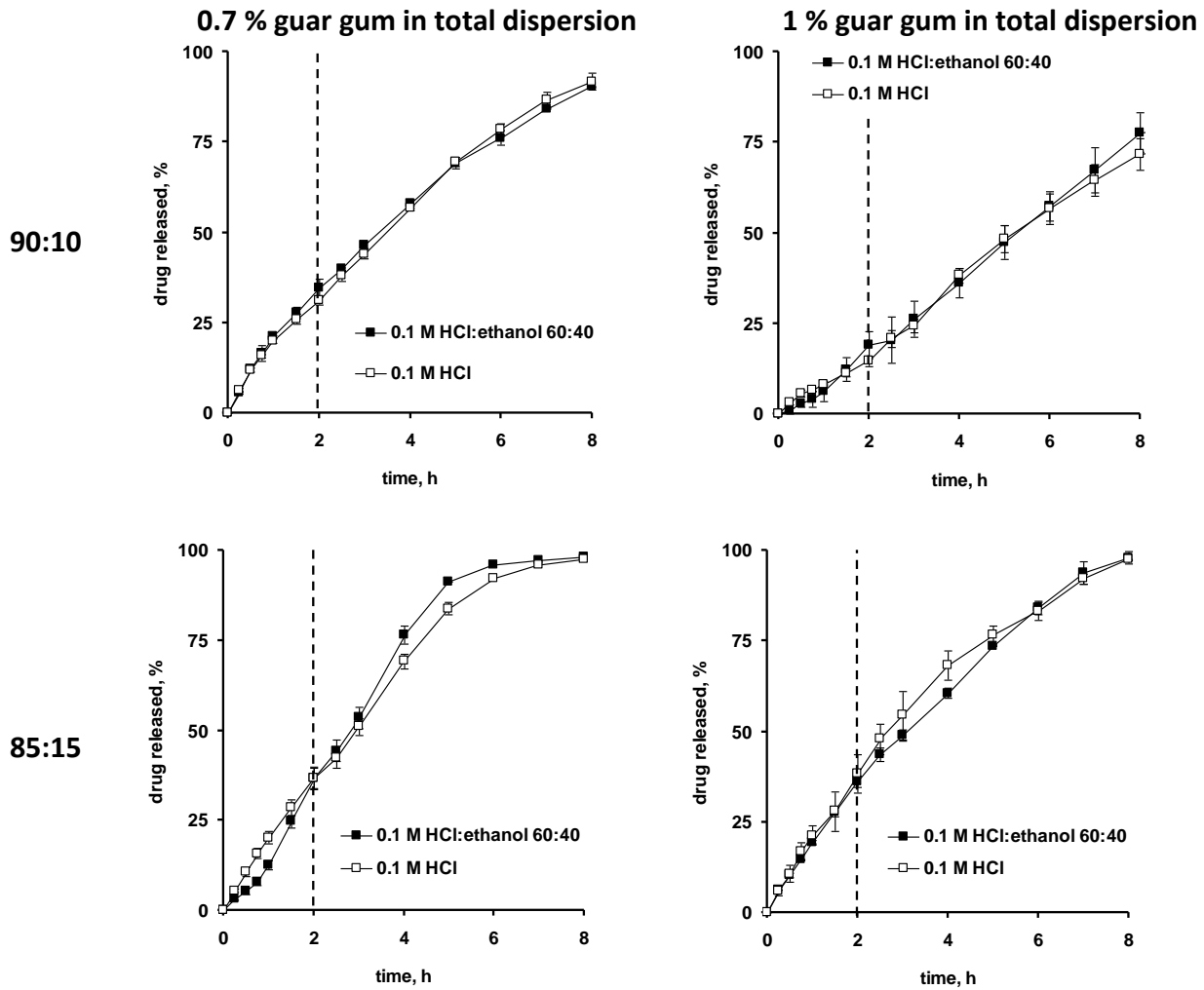


Figure 23: Impact of the guar gum dilution (left versus right column) on theophylline release from pellets coated with different ethylcellulose:guar gum blends (as indicated on the left hand side) (coating level: 20 % w/w). Release medium: 0.1 M HCl for 2 h, followed by phosphate buffer pH 7.4 (open symbols); or 0.1 M HCl:ethanol 60:40 for 2 h, followed by phosphate buffer pH 7.4 (filled symbols).

Figure 24a shows how the viscosity of the coating dispersion increases with increasing guar gum concentration in the formulation (the 100 % reference value is the total mass of the dispersion), irrespective of the ethylcellulose:guar gum blend ratio (indicated by the white, gray and black bars). This increase in viscosity can be attributed to more pronounced macromolecular interactions at higher guar gum concentrations. Note that at a given guar gum concentration in the total coating dispersion, the viscosity of the formulation generally decreases when increasing the relative guar gum content with respect to the ethylcellulose content (blend ratios 93:7 vs. 90:10 vs. 85:15). This is due to the higher water contents of the systems: With increasing relative guar gum content with respect to ethylcellulose, the water content of the total dispersion increases. For the viscosity of the coating dispersion not the guar gum percentage referred to the total dispersion mass is most decisive, but the guar gum percentage referred to the “guar gum + water” mass. Figure 24b shows the relationship between these two percentages. As it can be seen, for a given guar gum percentage referred to the total dispersion mass, the guar gum percentage referred to the “guar gum + water” mass decreases with increasing guar gum content with respect to the ethylcellulose content (white vs. gray vs. black bars). Thus, the amount of water present per amount of guar gum increases when varying the ethylcellulose:guar gum blend ratio from 93:7 to 90:10 to 85:15, resulting in decreasing coating dispersion viscosity (Figure 24a).

5.2.3. Coating level

Figure 25 shows the effects of varying the coating level of theophylline matrix pellets coated with 85:15 or 90:10 ethylcellulose:medium η guar gum blends on drug release in 0.1 M HCl for 2 h, followed by phosphate buffer pH 7.4 for 6 h (open symbols); or 0.1 M HCl:ethanol 60:40 for 2 h, followed by phosphate buffer pH 7.4 for 6 h (filled symbols). Firstly, ethanol-resistant drug release was observed at all coating levels and polymer blend ratios. Secondly, drug release generally decreased with increasing coating level (as expected), due to the increasing length of the diffusion pathways to be overcome. Thus, desired drug release profiles can effectively be fine-tuned by varying the coating level.

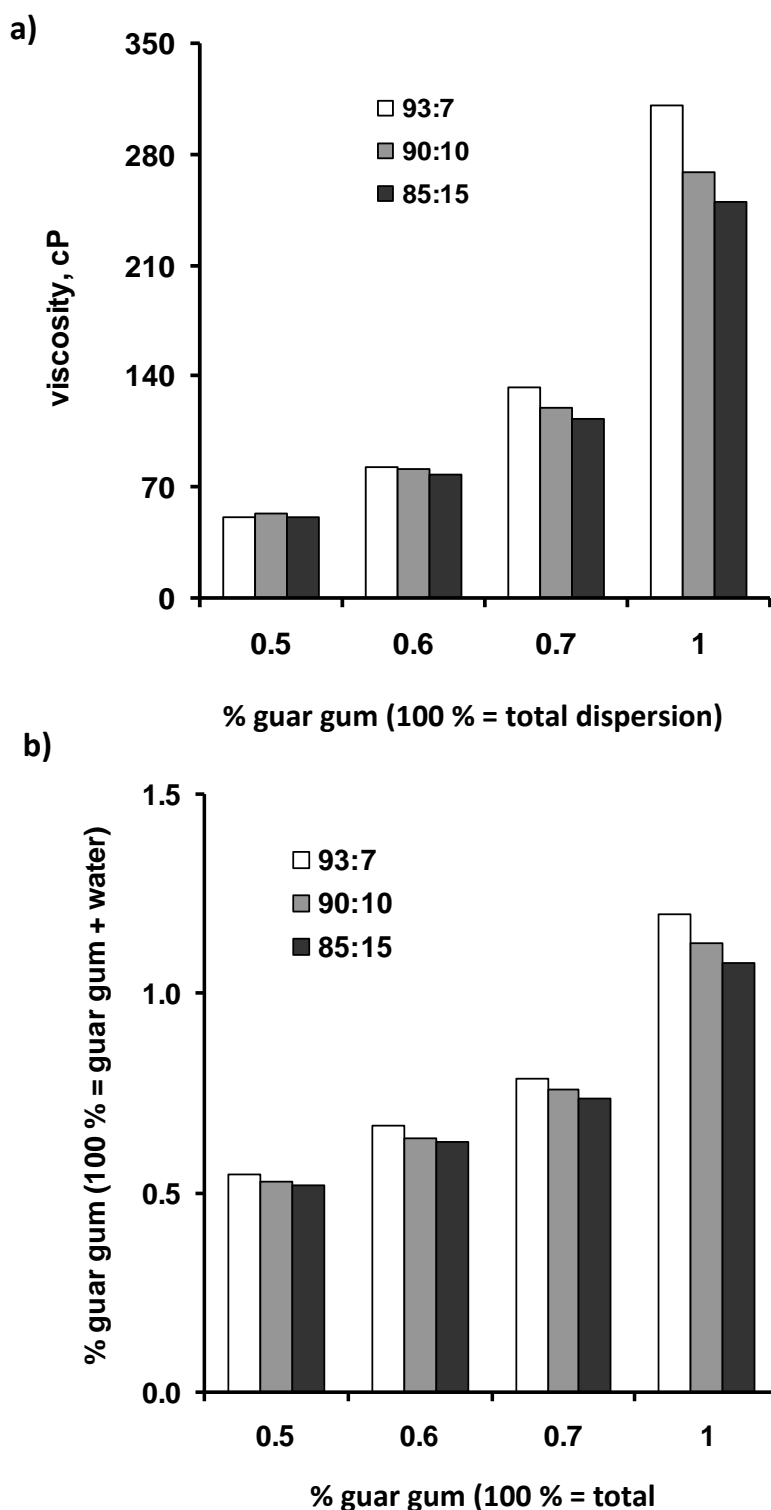


Figure 24. Effects of the degree of dilution of the coating dispersion: a) dependence of the viscosity of the applied formulation on the guar gum concentration in the dispersion (100 % being the total dispersion mass), and b) relationship between the “percentage of guar gum, 100 % being the guar gum + water mass” and the “percentage of guar gum, 100 % being the total dispersion mass”. The white, gray and black bars refer to ethylcellulose:guar gum 93:7, 90:10 and 85:15 (w:w) blends, respectively.

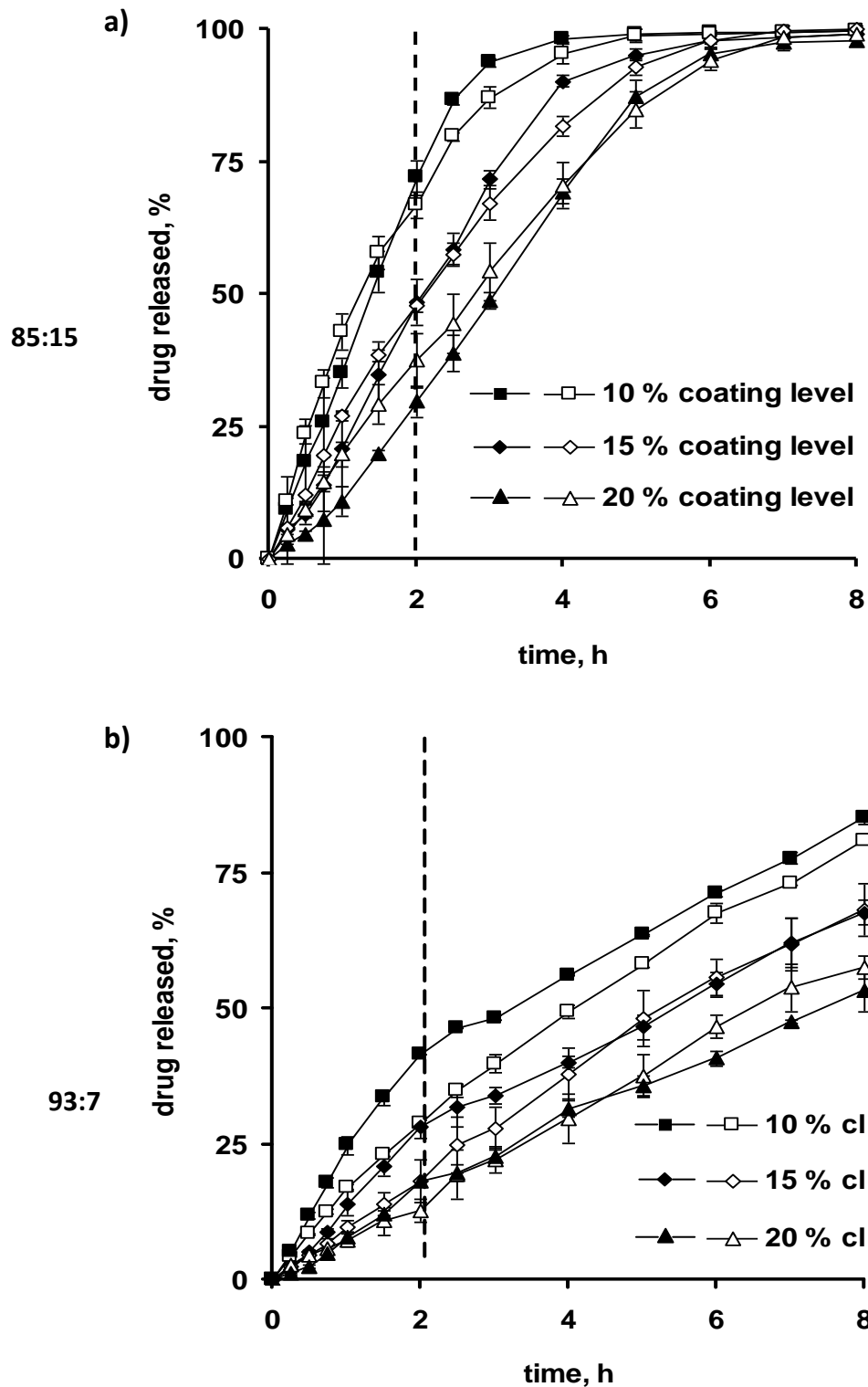


Figure 25: Impact of the coating level (indicated in the diagrams) on theophylline release from pellets coated with: a) ethylcellulose:guar gum 85:15 (0.7% guar gum in total dispersion), or b) 93:7 (1% guar gum in total dispersion) in 0.1 M HCl for 2 h, followed by phosphate buffer pH 7.4 (open symbols); or 0.1 M HCl:ethanol 60:40 for 2 h, followed by phosphate buffer pH 7.4 (filled symbols).

5.2.4. Storage stability

When using polymeric film coatings for controlled drug delivery, stability during long term storage is of utmost importance (Siepmann et al., 2008). Potential changes in the inner film structure might lead to changes in the resulting drug release rates (Kucera et al., *in press*). Importantly, the investigated film coatings based on ethylcellulose:guar gum blends were all long term stable without packaging material (“open storage” in glass vials), irrespective of the investigated polymer blend ratio, coating level, percentage of guar gum in the final coating dispersion and storage conditions: ambient (25 °C and 60 % relative humidity) or stress conditions (40 °C and 75 % relative humidity). Figures 26a and 26b just show a few examples: The solid curves in figure 26a illustrate theophylline release from 93:7 pellets coated with ethylcellulose:guar gum blends (20 % coating level) before storage, the dotted curves show the respective release profiles after 6 months open storage under ambient conditions. The release medium was 0.1 M HCl for the first 2 h, followed by phosphate buffer pH 7.4 for the subsequent 6 h in this case. The solid curves in figure 26b illustrate drug release from pellets coated with 15 or 20 % 85:15 ethylcellulose:guar gum blends before storage. The dotted curves show drug release from these systems after 6 months open storage under stress conditions. Here, the release medium was 0.1 M HCl:ethanol 60:40 for the first 2 h, followed by phosphate buffer pH 7.4 for the subsequent 6 h. As it can be seen, in all cases the solid and dotted curves are very similar, indicating good storage stability.

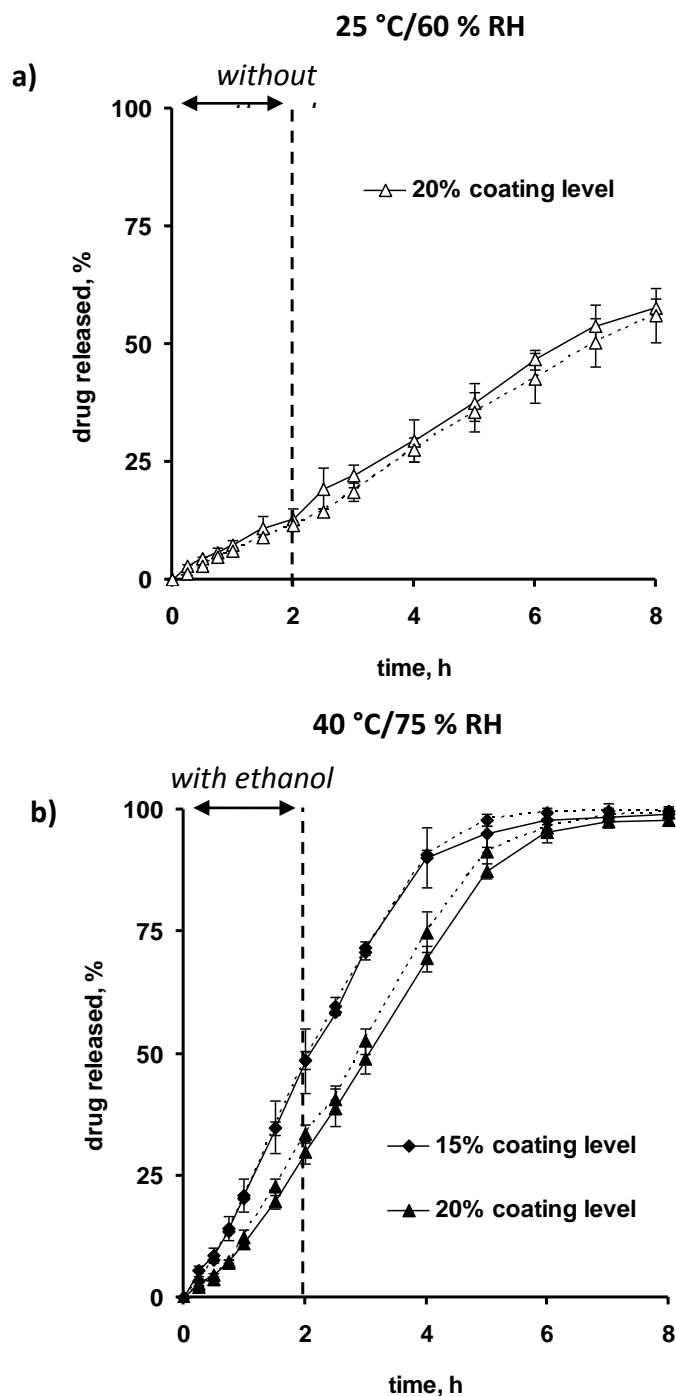


Figure 26: Storage stability (examples) of theophylline pellets coated with ethylcellulose:guar gum blends: a) 93:7 blend ratio, 20 % coating level, before and after 6 months open storage under ambient conditions (25 °C and 60 % relative humidity), 2 h exposure to 0.1 M HCl, followed by phosphate buffer pH 7.4; b) 85:15 blend ratio, 15 or 20 % coating level (as indicated), before and after 6 months open storage under stress conditions (40 °C and 75 % relative humidity), 2 h exposure to 0.1 M HCl:ethanol 60:40, followed by phosphate buffer pH 7.4. The solid curves show drug release before storage, the dotted curves after storage.

5.2.5. Guar gum viscosity

It has recently been reported that blends of ethylcellulose with *very low* or *low* η guar gum were not able to provide ethanol-resistant drug release from coated pellets (Rosiaux et al., *submitted*). In contrast, blends of ethylcellulose with *medium* or *high* η guar gum provided ethanol-resistant theophylline from coated pellets. So far, it was unknown what the critical guar gum viscosity might be, required to provide ethanol-resistant drug release. This is why various blends of *low* and *medium* η guar gum were prepared, exhibiting viscosities ranging from 149 to 304 cP (Table 1). Ethylcellulose was mixed with these guar gum blends at a ratio of 90:10. Theophylline matrix cores were coated with these blends (coating level = 20 %) and drug release was measured in 0.1 M HCl for 2 h, followed by phosphate buffer pH 7.4 for 6 h (open symbols); or 0.1 M HCl:ethanol 60:40 for 2 h, followed by phosphate buffer pH 7.4 for 6 h (filled symbols). Figure 27 shows some examples of the observed drug release profiles, Table 1 indicates the amounts of drug released after 2 h exposure to 0.1 M HCl and the differences to the amounts of drug released after 2 h exposure to 0.1 M HCl:ethanol 60:40. As it can be seen, the previously reported trend has been confirmed: With decreasing guar gum viscosity the ethanol sensitivity increases. According to the above mentioned definition, the guar gum viscosity must be equal to or greater than 151.0 cP. The decrease in ethanol-resistance with decreasing guar gum viscosity can probably at least partially be attributed the lower polymer molecular weight of the guar gum, resulting in an decreased degree of polymer-polymer chain entanglement: The shorter the guar gum chains, the less intensively they are entangled with each other and with the ethylcellulose molecules present in the film coating. Thus, the dissolution of the ethylcellulose in ethanol-containing media is less effectively hindered.

Table 1: Impact of the guar gum viscosity on the ethanol-resistance of theophylline pellets coated with 20 % 90:10 ethylcellulose:guar gum blends.

Blend ratio (w:w) <i>medium η: low η</i> guar gum	Viscosity \pm SD of the guar gum blend (n = 3) (cP)	Drug release in 0.1 N HCl after 2 h (%)	Difference in drug release after 2 h exposure to "0.1 N HCl" versus "0.1 M HCl:ethanol 60:40" (%)	Ethanol sensitivity
90:10	303.5 \pm 2.1	52.9	5.9	No
80:20	276.0 \pm 1.4	63.6	5.3	No
75:25	260.0 \pm 7.2	49.8	8.8	No
70:30	234.0 \pm 7.6	48.1	4.9	No
50:50	173.6 \pm 3.1	40.1	14.3	No
47.5:52.5	160.0 \pm 6.2	51.4	15.1	No
45:55	151.0 \pm 2.7	45.6	25.1	No
40:60	148.5 \pm 2.1	37.9	42.1	Yes

SD = standard deviation

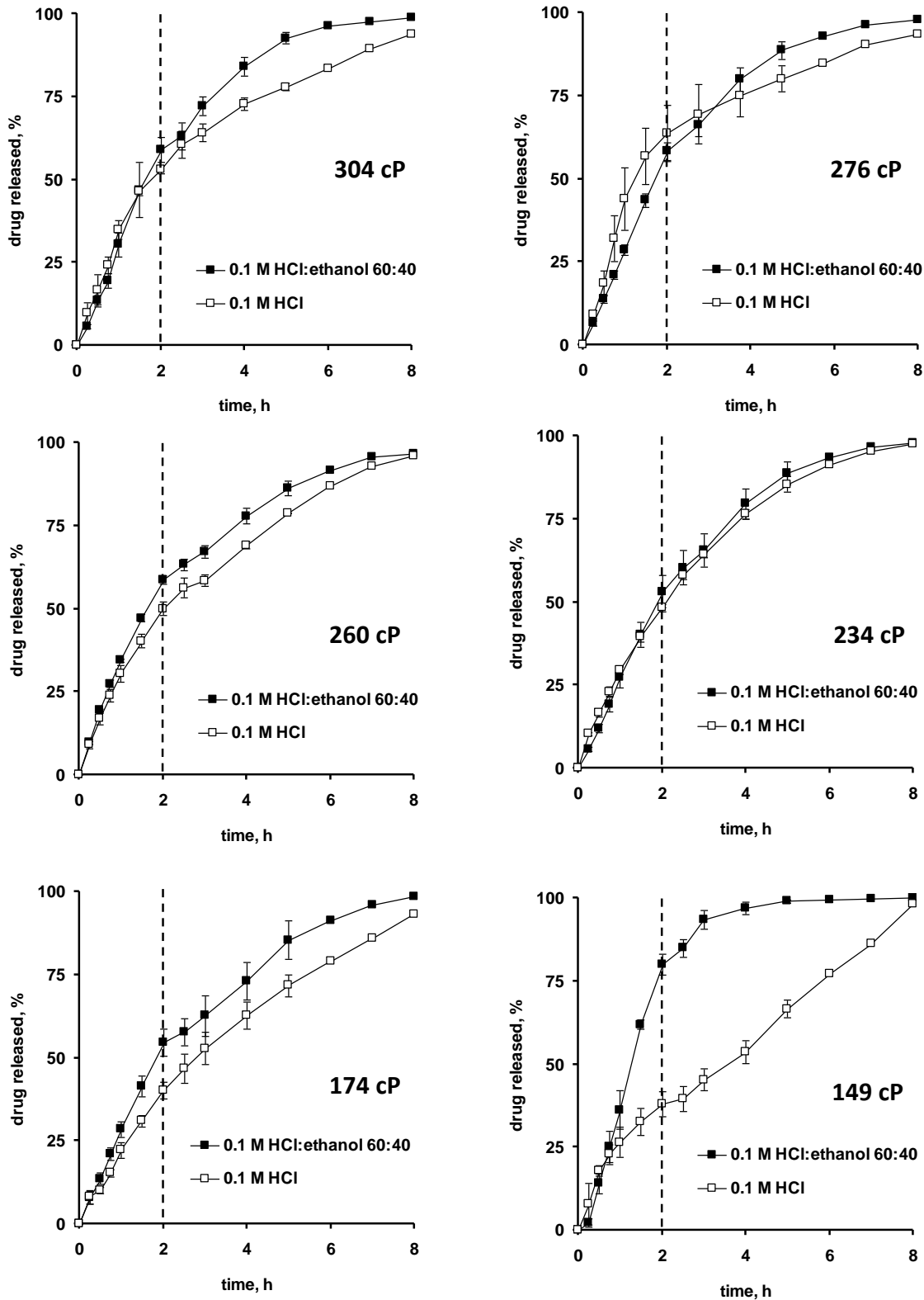


Figure 27: Importance of the apparent viscosity of the guar gum (indicated in the diagrams) used in 90:10 ethylcellulose:guar gum blends on theophylline release from coated pellets in 0.1 M HCl for 2 h, followed by phosphate buffer pH 7.4 (open symbols); or 0.1 M HCl:ethanol 60:40 for 2 h, followed by phosphate buffer pH 7.4 (filled symbols) (20 % coating level).

5.3. Conclusion

Appropriate blends of ethylcellulose and guar gum offer a great potential as controlled release film coatings providing ethanol-resistant drug release. The risk of dose dumping due to co-consumption of alcoholic beverages is likely to be effectively minimized. Important formulation parameters to be considered during product development include: (i) the amount of guar gum incorporated in the film coating (a minimum quantity being required to avoid undesired ethylcellulose dissolution in ethanol-rich media), and (ii) the guar gum viscosity (a minimum viscosity being required to assure sufficient polymer-polymer chain entanglement).

6. Elucidation of the underlying drug release mechanism of ethanol resistant coated pellets

6.1. Materials and methods

6.1.1. Materials

Theophylline matrix pellets (70 % drug content, diameter: 0.71-1.25 mm; FMC BioPolymer, Philadelphia, PA, USA); Ethylcellulose Aqueous Dispersion NF (Aquacoat[®] ECD 30; FMC BioPolymer); dibutyl sebacate (DBS; Morflex, Greensboro, NC, USA); ethanol (Fisher Bioblock Scientific, Illkirch, France); *medium viscosity* guar gum (*medium η* guar gum, apparent viscosity of a 1 % aqueous guar gum solution = 320 cPs; Polygum 240/80; Polygal Trading, Maerstetten, Switzerland); *high viscosity* guar gum (*high η* guar gum, apparent viscosity of a 1 % aqueous guar gum solution in the range of 575-625 cPs; Guar HV 225; Alland & Robert, Port-Mort France, France). The apparent viscosities were measured using an AR2000Ex rheometer (TA Instruments, New Castle, DE, USA) at a shear rate of 50 s⁻¹ in a 1 % aqueous guar gum solution measured rotationally at 20 °C after 1 min equilibration using a 6 cm acrylic cone (1°), wherein the shear was ramped up linearly from 1 to 50 s⁻¹ in 25 steps over 29 s.

6.1.2. Preparation and characterization of thin polymeric films

Aquacoat[®] ECD 30 was plasticized for 1 d with 25 % DBS (w/w; based on the ethylcellulose mass). Guar gum was dissolved in purified water at 65 °C (2 h stirring) and cooled down to room temperature. The two liquids were blended and stirred for 30 min prior to use. Films were prepared by spraying (0.8 mm spray nozzle) or casting (as indicated) Aquacoat[®] ECD 30:guar gum blends onto Teflon plates and subsequent controlled drying for 24 h at 60 °C.

The (water + ethanol) uptake and dry mass loss kinetics of the films were determined as follows: Pieces of 5 cm × 5 cm were placed into 100 mL plastic flasks filled with 100 mL pre-heated release medium, followed by horizontal shaking (37 °C, 80 rpm; GFL 3033, Gesellschaft fuer Labortechnik, Burgwedel, Germany). At pre-determined time points, samples were withdrawn, accurately weighed [wet mass (*t*)] and dried to constant mass at

60°C [dry mass (t)]. The (water + ethanol) content (%) and dry film mass (%) at time t were calculated as follows:

$$(water + ethanol) \text{ content } (\%)(t) = \frac{wet \ mass(t) - dry \ mass(t)}{wet \ mass(t)} \cdot 100 \% \quad (1)$$

$$dry \ film \ mass (\%)(t) = \frac{dry \ mass(t)}{dry \ mass(t=0)} \cdot 100 \% \quad (2)$$

The mechanical properties of the films (puncture strength, percent elongation and energy at break) in the dry and wet state were measured using the puncture test and a texture analyzer (TAXT.Plus, Swantech, Villeneuve la Garenne, France). Film specimens were mounted on a film holder ($n = 6$). The puncture probe (spherical end: 5 mm diameter) was fixed on the load cell (5 kg) and driven downward with a cross-head speed of 0.1 mm/s to the center of the film holder's hole (diameter: 10 mm). Load versus displacement curves were recorded until rupture of the film and used to determine the mechanical properties as follows:

$$puncture \ strength = \frac{F}{A} \quad (3)$$

where F is the load required to puncture the film; A represents the cross-sectional area of the edge of the film located in the path.

$$\% \text{ elongation at break} = \frac{\sqrt{R^2 + d^2} - R}{R} \cdot 100 \% \quad (4)$$

Here, R denotes the radius of the film exposed in the cylindrical hole of the holder and d the displacement to puncture.

$$\text{energy at break per unit volume} = \frac{AUC}{V} \quad (5)$$

where *AUC* is the area under the load versus displacement curve and *V* the volume of the film located in the die cavity of the film holder (the energy at break is normalized to the film's volume).

6.1.3. Pellet coating

Theophylline matrix cores were coated with Aquacoat[®] ECD 30:guar gum 85:15 or 93:7 blends (as indicated). Aquacoat[®] ECD 30 was plasticized for 1 d with 25 % DBS (w/w; based on the ethylcellulose mass). Guar gum (*medium* η guar gum, if not otherwise stated) was dissolved in purified water at 65 °C (0.7 % or 1 % w/w, as indicated; 100 % reference value = total coating formulation; 2 h stirring) and cooled down to room temperature. The two liquids were blended and stirred for 30 min prior to use. The coating dispersions were sprayed onto theophylline pellets using a fluidized bed coater (Strea 1, Wurster insert; Niro; Aeromatic-Fielder, Bubendorf, Switzerland). The process parameters were as follows: inlet temperature = 38 °C, product temperature = 38 ± 2 °C, spray rate = 2g/min, atomization pressure = 1.2 bar, nozzle diameter = 1.2 mm. After coating the pellets were further fluidized for 10 min and subsequently cured for 24 h at 60 °C in an oven.

6.1.4. Drug release measurements

From ensembles of pellets: Theophylline release from coated pellets was measured in 0.1 M HCl or 0.1 M HCl: ethanol blends (optionally containing different amounts of NaCl), followed by phosphate buffer pH 7.4 (USP 36) using the USP 36 paddle apparatus (Sotax, Basel, Switzerland) (900 mL, complete medium change after 2 h; 37 °C, 100 rpm, n = 3). At pre-determined time points, 3 mL samples were withdrawn and analyzed UV-spectrophotometrically [λ = 270.4 nm in 0.1 N HCl, λ = 272.2 nm in 0.1 M HCl:ethanol blends and phosphate buffer pH 7.4] (UV 1650 PC, Shimadzu, Champs-sur-Marne, France).

From single pellets: Single pellets were placed into 100 mL plastic flasks filled with 100 mL pre-heated release medium (0.1 M HCl or 0.1 M HCl:ethanol 60:40 for 2 h, followed by phosphate buffer pH 7.4). The flasks were horizontally shaken at 37 °C and 80 rpm (GFL 3033). At pre-determined time points, 3 mL samples were withdrawn and analyzed UV-spectrophotometrically as described above.

6.1.5. SEM studies

The morphology of coated pellets was characterized using a cold field emission high resolution scanning electron microscope (S-4700 Field Emission Gun; Hitachi, Hitachi High-Technologies Europe, Krefeld, Germany). Samples were covered under vacuum with a carbon layer.

6.1.6 Diffusion cell studies

Drug permeation through thin, drug-free (sprayed) polymeric films was measured using vertical diffusion cells, which were placed in a horizontal shaker (37 °C, 80 rpm; GFL 3033). The film surface exposed to the medium was 1.8 cm². The donor compartment (at the top) was filled with an excess of dry theophylline powder (as received). The acceptor compartment (at the bottom) was filled with 350 mL 0.1 M HCl or 0.1 M HCl:ethanol 60:40. At pre-determined time intervals, 3 mL samples were withdrawn and analysed UV-spectrophotometrically as described above. All experiments were conducted in triplicate.

6.1.7. Determination of the drug solubility and of the partition coefficient of the drug

Excess amounts of theophylline powder were exposed to 0.1 M HCl or 0.1 M HCl:ethanol 60:40 in flasks in a horizontal shaker (80 rpm, 37 °C). At pre-determined time intervals, samples were withdrawn, filtered and analysed for their drug content (as described above) until equilibrium was reached.

Thin, initially drug free (cast) films were exposed to saturated theophylline solutions in 0.1 M HCl or 0.1 M HCl:ethanol 60:40 containing excess amounts of non-dissolved drug in flasks in

a horizontal shaker (80 rpm, 37 °C). At pre-determined time intervals, film samples were withdrawn, surface water removed and the films analyzed for their drug content. The latter was determined UV-spectrophotometrically upon film dissolution in ethanol. The partition coefficient was calculated from the plateau concentrations reached in the films and in the aqueous phase at equilibrium. All experiments were conducted in triplicate.

6.2. Results and discussion

6.2.1. Drug release from single pellets

When elucidating the drug release mechanisms from multiple dosage forms, it is decisive to first of all clarify whether the single units release in a similar way, or whether the release profile observed with ensembles of multiple units is the sum of very different individual release profiles (Borgquist et al., 2002). For instance, in an extreme case, zero order release kinetics observed from an ensemble of hundreds of pellets might result from pulsatile drug release from individual pellets with highly variable lag-times, evenly distributed within the observation period.

Figure 28 shows the experimentally measured theophylline release kinetics from single pellets coated with 20 % (w/w) 85:15 ethylcellulose:guar gum. The release medium was either 0.1 M HCl for 2 h (Figure 28a), or 0.1 M HCl:ethanol 60:40 for 2 h (Figure 28b), in both cases followed by 6 h phosphate buffer pH 7.4. The thick curves indicate the respective mean values, the error bars standard deviations. Clearly, the investigated single pellets behave similarly, irrespective of the type of release medium: About zero order release kinetics are observed in all cases. Thus, the underlying drug release mechanism seems to be the same for all single units. Slight variations in the film coatings' thickness are likely to be responsible for the observed differences in the individual release profiles (Ho et al., 2009). Importantly, the presence or absence of 40 % ethanol in the release medium does not seem to affect the underlying drug release mechanism.

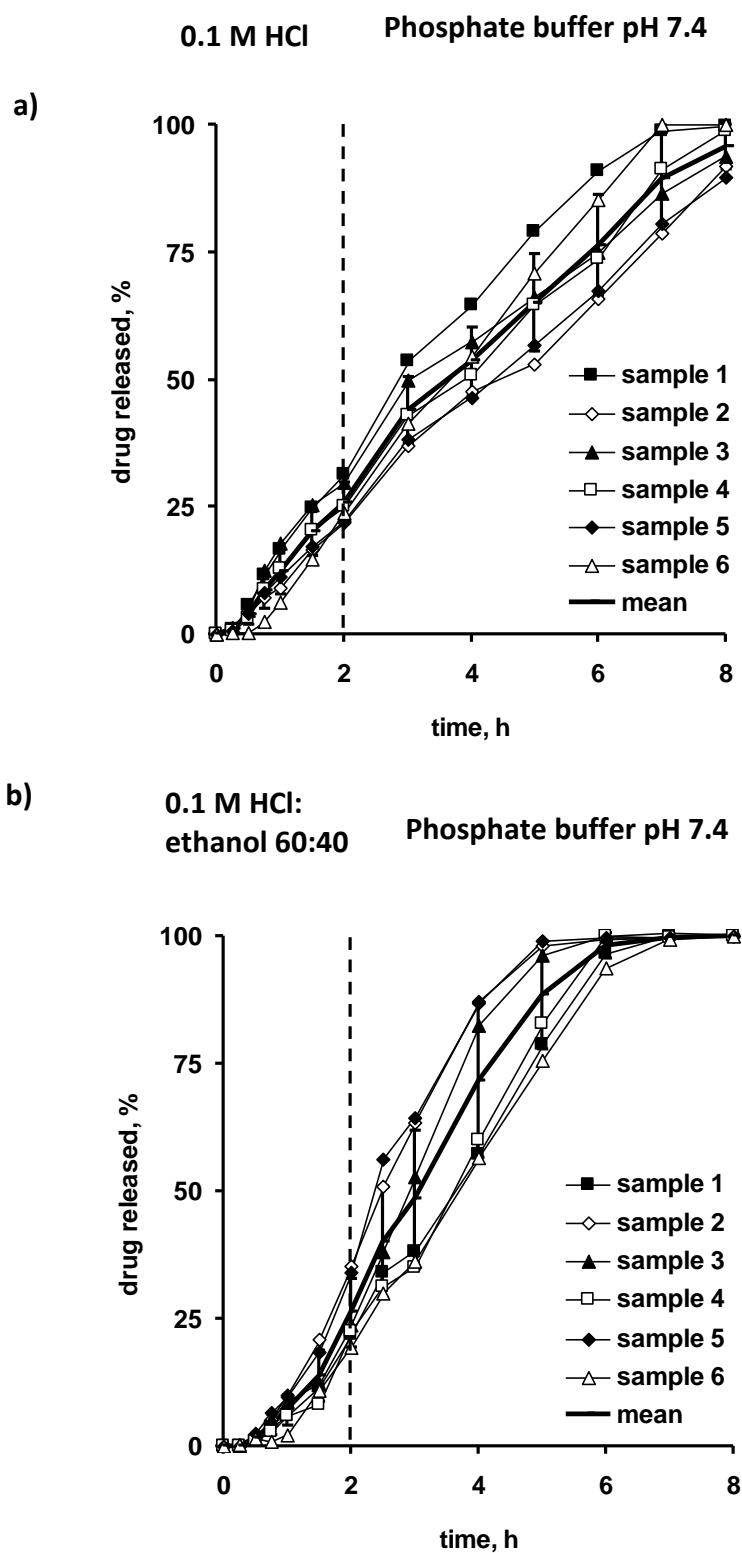


Figure 28: Theophylline release from single pellets coated with 20% (w/w) ethylcellulose:guar gum 85:15 in: **a)** 0.1 M HCl for 2 h, followed by phosphate buffer pH 7.4, or **b)** 0.1 M HCl:ethanol 60:40 for 2 h, followed by phosphate buffer pH 7.4. The thick curves show the respective mean values, error bars indicate standard deviations. The guar gum concentration in the total dispersion was 0.7% w/w.

6.2.2. Impact of the osmolality of the release medium

Drug release from coated pharmaceutical dosage forms might be strongly affected by the osmolality of the release medium. In certain cases, osmotic effects play a major role for the control of drug release (Marucci et al., 2010). Importantly, considerable variations in the osmolality of the release medium did not significantly affect the resulting drug release kinetics, irrespective of the ethylcellulose:guar gum blend ratio and the presence/absence of ethanol in the release medium. Figures 29a and 29b show for example the experimentally measured release of theophylline from pellets coated with 20 % w/w ethylcellulose:guar gum 85:15 in 0.1 M HCl or 0.1 M HCl:ethanol 60:40 containing 0-5 % NaCl for 2 h, followed in both cases by phosphate buffer pH 7.4 for 6 h. As it can be seen, there was only a limited impact on drug release, despite the important variations in the osmolality of the surrounding bulk fluid. Also, no general tendency was visible for any release medium. Based on these results, osmotic pumping as the dominant release mechanism can be excluded.

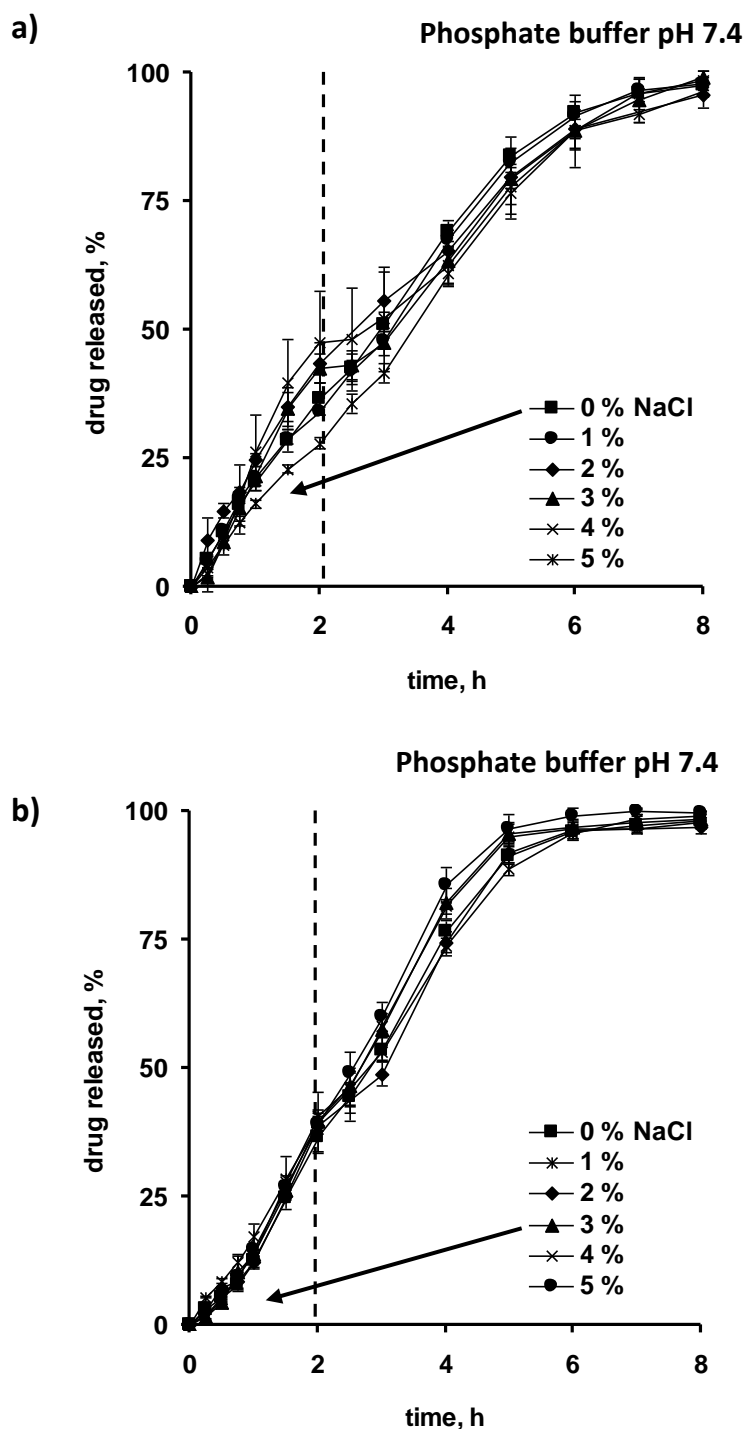


Figure 28: Impact of the osmolality of the bulk fluid: Theophylline release from pellets coated with 20 % w/w ethylcellulose:guar gum 85:15 in: **a)** 0.1 M HCl containing different amounts of NaCl for 2 h, followed by phosphate buffer pH 7.4, or **b)** 0.1 M HCl:ethanol 60:40 containing different amounts of NaCl for 2 h, followed by phosphate buffer pH 7.4. The guar gum concentration in the total dispersion was 0.7% w/w.

6.2.3. Morphology and mechanical properties of the film coatings

Drug release from polymer coated solid dosage forms can either occur through the continuous macromolecular network, or through water-filled cracks or pores (potentially created upon water penetration into the system) (Maroni et al., *in press*). Often, drug transport through water-filled pores/cracks is much faster than drug transport through a continuous polymeric network (Siepmann and Siepmann, 2008). In order to know whether water-filled channels exist in the investigated ethylcellulose:guar gum coated pellets, the latter's morphology was studied using Scanning Electron Microscopy before and after exposure to 0.1 M HCl, or 0.1 M HCl:ethanol 60:40 for 2 h. As it can be seen in figure 30, no signs for the presence of pores or cracks are visible before exposure to the release media. The same was true after 2 h contact with the bulk fluids, irrespective of the presence of high ethanol concentrations (data not shown). However, it must be pointed out that the pellets had to be dried after exposure to the release media and that eventually created pores/cracks were closed during this drying step. In addition, nanosized pores would not necessarily be visible (e.g., being hidden by the carbon layer or due to the limited resolution). Thus, caution has to be paid when drawing conclusions from this type of measurements. Nevertheless, the absence of visible macro-sized pores under the given conditions is interesting information. In other studies on different polymeric film coatings such pores could clearly be seen in SEM pictures (Lecomte, 2005a).

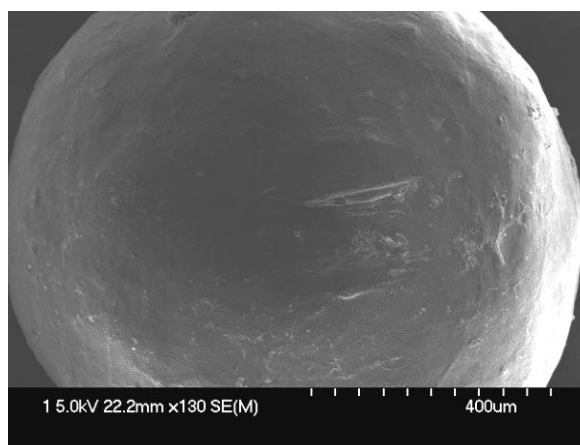


Figure 30: SEM picture of the surface of a pellet coated with 20 % w/w ethylcellulose:guar gum 85:15 before exposure to the release medium. The guar gum concentration in the total dispersion was 0.7% w/w.

Obviously, the mechanical strength of the polymeric film coatings is decisive for the potential creation of cracks upon water penetration into the systems. Water entering the pellet core builds up a hydrostatic pressure, which acts against the macromolecular membrane. If the latter is fragile, the onset of crack formation is likely after a system-specific lag-time. In contrast, if the film coating is mechanically sufficiently stable and withstands the generated hydrostatic pressure, drug release might be controlled by diffusion through the intact polymeric network. Table 2 shows the puncture strength, percent elongation at break and energy required to break thin, free films based on different ethylcellulose:guar gum blends in the dry state (before exposure to the release medium). As it can be seen, the mechanical stability of the films increases with increasing guar gum content. This might at least partially be attributable to the fact that an aqueous *dispersion* of ethylcellulose nanoparticles and an aqueous *solution* of guar gum were used for film preparation (as this is the case for the film coatings). Thus, the underlying film formation mechanism is fundamentally different for the two polymers and a much higher degree of polymer-polymer chain entanglement can be expected in the case of guar gum (Lecomte et al., 2004a).

Table 2: Impact of the ethylcellulose:guar gum blend ratio (w/w) on the mechanical properties (puncture strength, percent elongation at break and energy at break) of free (cast) films in the dry state (mean values +/- standard deviation).

	Ethylcellulose:guar gum blend ratio				
	100:0	93:7	90:10	85:15	0:100
Puncture strength, MPa	0.2 ± 0.0	0.4 ± 0.0	0.5 ± 0.0	0.6 ± 0.1	11.6 ± 4.4
Elongation at break, %	1.1 ± 0.2	1.2 ± 0.2	1.1 ± 0.1	0.8 ± 0.1	0.6 ± 0.3
Energy at break, J/m³	11 ± 0.2	18 ± 0.2	19 ± 2	22 ± 5	323 ± 185

However, it is well known that the composition of polymeric film coatings can significantly change upon exposure to the release medium (Lecomte et al., 2004b). For instance, water penetrating into the film coating can act as a plasticizer for a polymeric component. On the other hand, water soluble film compounds (e.g. guar gum) might at least partially leach out of the film into the bulk fluid. Also, plasticizers (such as dibutyl sebacate) might be partially lost into the surrounding medium. Due to these changes in the film coatings' composition, the latter's mechanical properties might be significantly altered. Figure 31 shows the dynamic changes in the puncture strength, percent elongation at break and energy required to break thin films based on 93:7, 90:10, or 85:15 ethylcellulose blends upon exposure to 0.1 M HCl:ethanol 60:40 (37 °C) for different time periods. As it can be seen, the mechanical stability of the systems: (i) decreases with time in the case of 85:15 ethylcellulose:guar gum blends, (ii) generally only slightly, and more or less arbitrarily varies with time in the case of the other two blends. In most cases, the energy required to break the films increases with increasing guar gum content, irrespective of the exposure time to the release medium (for the reasons discussed above).

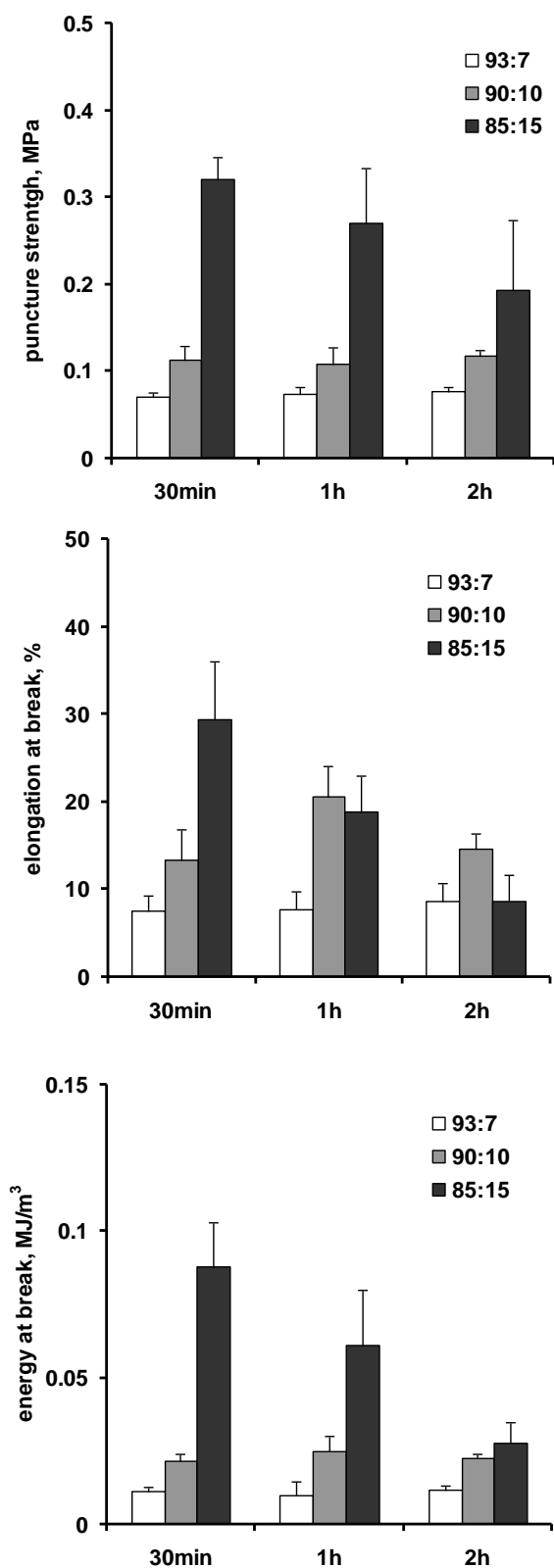


Figure 31: Dynamic changes in the mechanical properties of free ethylcellulose:guar gum (cast) films upon exposure to 0.1 M HCl:ethanol 60:40 (37 °C) for different time periods. The ethylcellulose:guar gum blend ratio (w/w) is indicated in the diagrams. The guar gum concentration in the total dispersion was 0.7% w/w.

6.2.4. Drug mobility within the film coatings

Based on the experimentally observed drug release kinetics from single pellets, the very limited impact of changes in the osmolality of the release medium on drug release, the morphology of the pellets' surface before and after exposure to the release medium as well as the mechanical properties of the polymeric films and dynamic changes thereof upon contact with the bulk fluids, one hypothesis can be that drug release is eventually controlled primarily by diffusion through the intact polymeric film coatings. In order to evaluate the validity of this assumption, the apparent diffusion coefficient of theophylline in thin, free films of identical composition as the investigated film coatings was experimentally determined using vertical diffusion cells. Knowing these values and using an appropriate solution of Fick's law of diffusion the resulting drug release kinetics from coated pellets can be quantitatively predicted (based on the hypothesis of purely diffusion controlled drug release). Comparison of these theoretical predictions with independent experimental results should allow evaluating the validity of the hypothesized release mechanism.

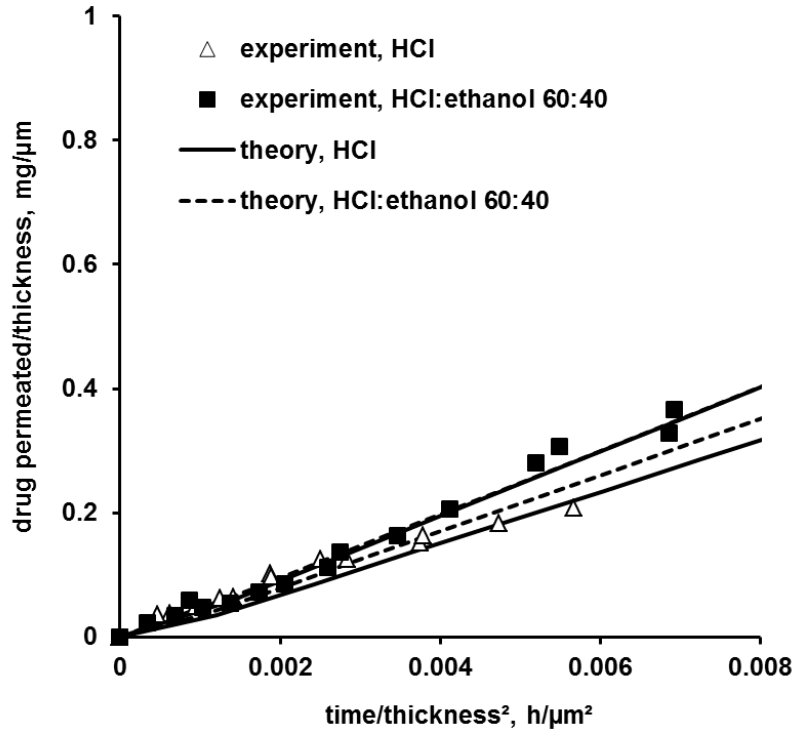


Figure 32: Drug permeation through sprayed thin ethylcellulose:guar gum 85:15 films (measured in vertical diffusion cells): The donor compartment initially contained dry theophylline powder, the acceptor compartment pre-heated (37 °C) 0.1 M HCl or 0.1 M HCl:ethanol 60:40 (as indicated). The symbols represent the experimentally measured results, the curves the fitted theory (Eq. 6). Note that “time/thickness²” is plotted on the x-axis, and “drug permeated/thickness” is plotted on the y-axis to account for slight variations in the film thickness from sample to sample. This type of normalization is possible, according to Eq. 6. The guar gum concentration in the total dispersion was 0.7% w/w.

The symbols in Figure 32 show the experimentally measured permeation of theophylline through thin ethylcellulose:guar gum 85:15 films. Vertical diffusion cells were used, the films were initially drug-free, the donor compartment contained theophylline powder, the acceptor compartment either 0.1 M HCl (filled symbols), or 0.1 M HCl:ethanol 60:40 (open symbols) (37 °C), in order to mimic the conditions in initially dry pellets exposed to the release medium. Importantly, about straight lines were observed right from the beginning. Thus, water penetration into and through the films was rapid compared to drug diffusion through the polymeric barrier: Very soon, a steady-state was established, with a saturated

drug solution in the donor compartment and perfect sink conditions in the acceptor compartment. Hence, also the dissolution of the theophylline particles was much more rapid than the subsequent diffusion through the thin films. The thickness of the latter was in the range of 17 to 23 μm (thus, in a similar range as the film coatings of the investigated pellets). Furthermore, no major difference was observed when using 0.1 M HCl versus 0.1 M HCl:ethanol 60:40 as bulk fluids. Hence, the mobility of the drug in the film coatings does not seem to be affected by the presence/absence of high ethanol concentrations to a noteworthy extent. Fitting the following solution of Fick's law of diffusion to the experimentally determined drug permeation kinetics allowed determining the apparent diffusion coefficient of the drug in these polymeric films (Crank, 1975):

$$M_t = 2 \cdot A \cdot L \cdot K \cdot c_s \cdot \left[\frac{D \cdot t}{4 \cdot L^2} - \frac{1}{6} - \frac{2}{\pi^2} \cdot \sum_{n=1}^{\infty} \frac{(-1)^n}{n^2} \cdot \exp\left(\frac{-D \cdot n^2 \cdot \pi^2 \cdot t}{4 \cdot L^2}\right) \right] \quad (6)$$

where M_t is the cumulative absolute amount of drug in the acceptor compartment at time t ; A denotes the surface area of the film available for diffusion, L the half-thickness of the film, and c_s the solubility of the drug in the medium; K is the partition coefficient of the drug between the film and the bulk fluid. The solubility of theophylline in 0.1 M HCl and 0.1 M HCl:ethanol 60:40 at 37 °C was experimentally measured and found to be equal to 12.7 +/- 0.5 and 30.2 +/- 1.0 mg/mL, respectively. The partition coefficient, K , of this drug between the investigated ethylcellulose:guar gum films and 0.1 M HCl or 0.1 M HCl:ethanol 60:40 was determined to be equal to 1.8 and 0.5 (for both polymer:polymer blend ratios: 85:15 and 93:7).

As it can be seen in Figure 32, good agreement was obtained when fitting Eq. 6 (curves) to the experimentally determined drug permeation kinetics through the polymeric films (symbols). Based on these calculations, the following apparent diffusion coefficients of theophylline in ethylcellulose:guar gum 85:15 based films could be determined: $D = 3.2$ +/- 0.5 and 5.0 +/- 0.4×10^{-8} cm^2/s upon exposure to 0.1 M HCl and 0.1 M HCl:ethanol 60:40, respectively. In the case of ethylcellulose:guar gum 93:7 blends, the D values were equal to 2.1 +/- 0.4 and 2.6 +/- 0.7×10^{-8} cm^2/s , respectively. Thus, the presence of 40 % ethanol in

the bulk fluid had only a limited impact on drug mobility within the polymeric films. As expected, the theophylline mobility increased with increasing guar gum content, which can probably be attributed to increased water uptake of the films (guar gum being more hydrophilic than ethylcellulose) and/or partial leaching of this compound into the bulk fluid (leading to less dense polymeric networks).

6.2.5. Mathematical modeling of drug release

Based on this knowledge and on the hypothesis that theophylline release is predominantly controlled by drug diffusion through the intact polymeric film coatings (no crack formation), the following solution of Fick's second law of diffusion can be used to quantitatively predict the resulting drug release kinetics from coated pellets (Siepmann and Siepmann, 2012):

$$M_t = \frac{4 \cdot \pi \cdot D \cdot K \cdot c_s \cdot R_o \cdot R_i}{R_o - R_i} \cdot t \quad (7)$$

where M_t denotes the cumulative amount of drug released at time t ; D is the apparent diffusion coefficient of the drug within the film coating; K is the partition coefficient of the drug between the film coating and the bulk fluid; c_s is the solubility of the drug in the core, and R_i and R_o are the inner and outer radii of the device.

Knowing the values on the right hand side of Eq. 7, the resulting drug release kinetics could be quantitatively predicted for arbitrary coating levels. The straight lines in Figure 6 show some examples: The dashed lines illustrate the theoretically predicted theophylline release kinetics in 0.1 M HCl from pellets coated with 10, 15, or 20 % ethylcellulose:guar gum 85:15 (as indicated). The solid lines illustrate the respective release kinetics in 0.1 M HCl:ethanol 60:40. As it can be seen, very similar theophylline release rates are expected in the presence and absence of ethanol in the release medium. Thus, the slightly increased apparent diffusion coefficient of the drug and the increased drug solubility in the presence of 40 % ethanol is compensated by the decreased partition coefficient between the film coating and the bulk fluid. In order to evaluate the validity of these theoretical predictions, the respective pellets were prepared in reality (only after the predictions were made) and the

resulting drug release kinetics were measured experimentally. Looking at Figure 33, it can clearly be seen that the theoretical predictions (lines) and independent experiments (symbols) were in good agreement in all cases: irrespective of the presence/absence of ethanol in the release medium and of the coating level.

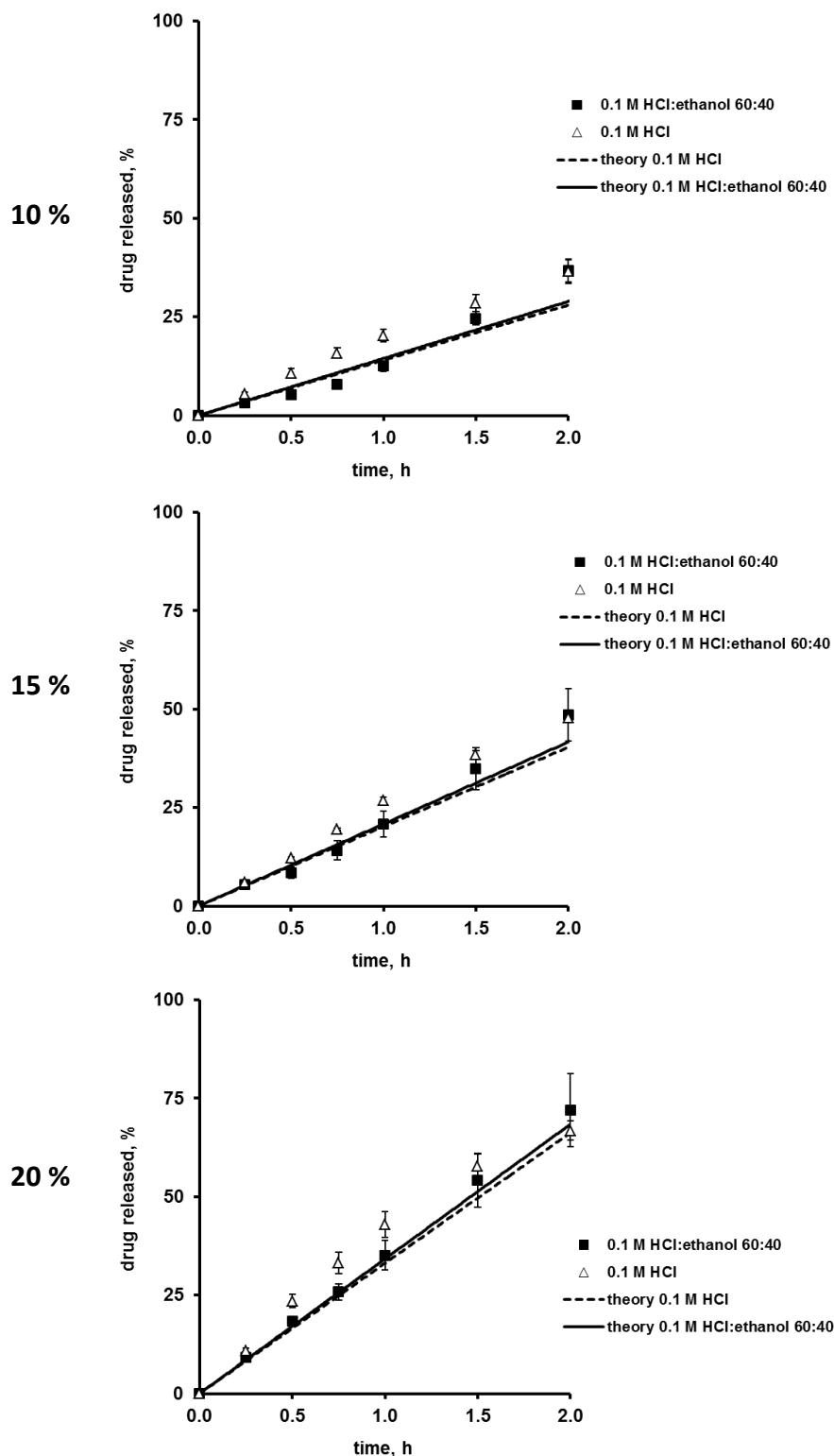


Figure 33: Theoretical prediction (straight lines, Eq. 7) and independent experimental verification (symbols): Theophylline release from pellets coated with 10, 15 or 20 % ethylcellulose:guar gum 85:15 in 0.1 M HCl, or 0.1 M HCl:ethanol 60:40 (as indicated). The guar gum concentration in the total dispersion was 0.7% w/w.

Thus, theophylline diffusion through the intact polymeric film coatings seems to be indeed the dominant mass transport mechanism controlling drug release from these dosage forms. Figure 34 shows that this was also true for 93:7 ethylcellulose:guar gum blends. The dashed and solid lines show the theoretically predicted theophylline release kinetics from pellets (coating level = 15 %) in 0.1 M HCl or 0.1 M HCl:ethanol 60:40, respectively. The symbols illustrate the independent experimental results. The observed good agreement further confirms the hypothesized drug release mechanism. These examples also demonstrate the great practical benefit of this type of mathematical modeling: *In silico* simulations offer the possibility to replace time-consuming and cost-intensive series of trial-and-error experiments during product optimization. The film composition and coating level required to achieve a specific, desired drug release profile can be theoretically predicted.

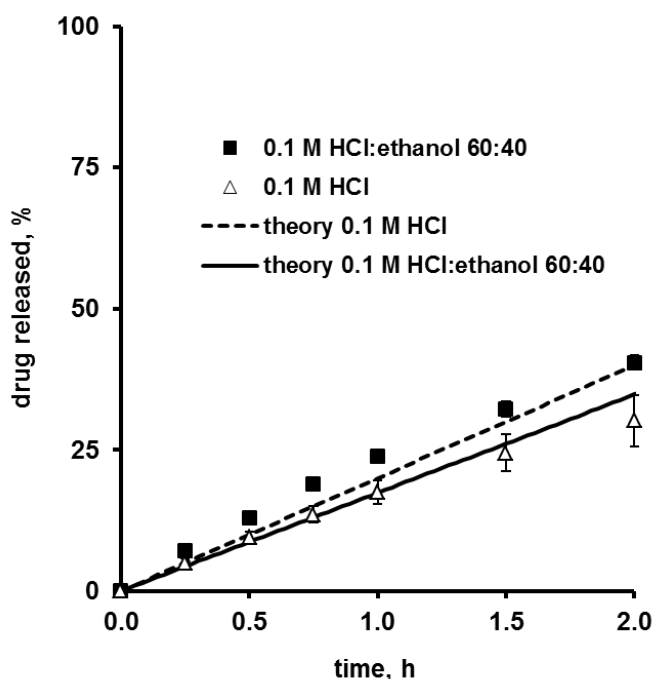


Figure 34: Theoretical prediction (straight lines, Eq. 7) and independent experimental verification (symbols): Theophylline release from pellets coated with 15 % ethylcellulose:guar gum 93:7 in 0.1 M HCl, or 0.1 M HCl:ethanol 60:40 (as indicated).

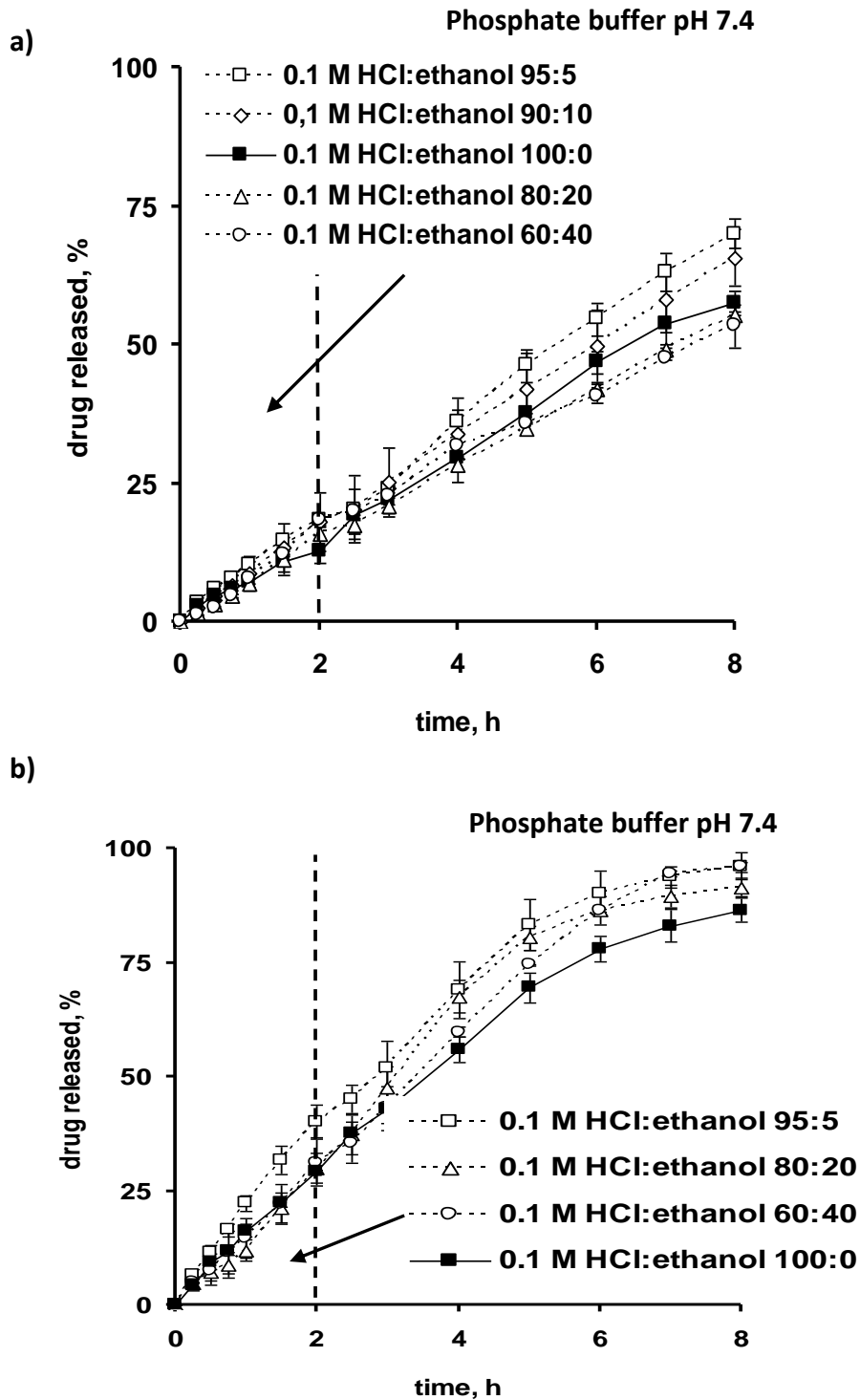


Figure 35: Impact of intermediate ethanol concentrations in the release medium: Theophylline release from pellets coated with 20%: **a)** 93:7 ethylcellulose:medium η guar gum (guar gum concentration in the total dispersion = 1.0% w/w), or **b)** 85:15 ethylcellulose:high η guar gum (guar gum concentration in the total dispersion = 0.7% w/w). The release medium was a 100:0, 95:5, 90:10, 80:20, or 60:40 blend of 0.1 M HCl and ethanol (V/V) (as indicated) for the first 2 h, followed by phosphate buffer pH 7.4 for the subsequent 6 h.

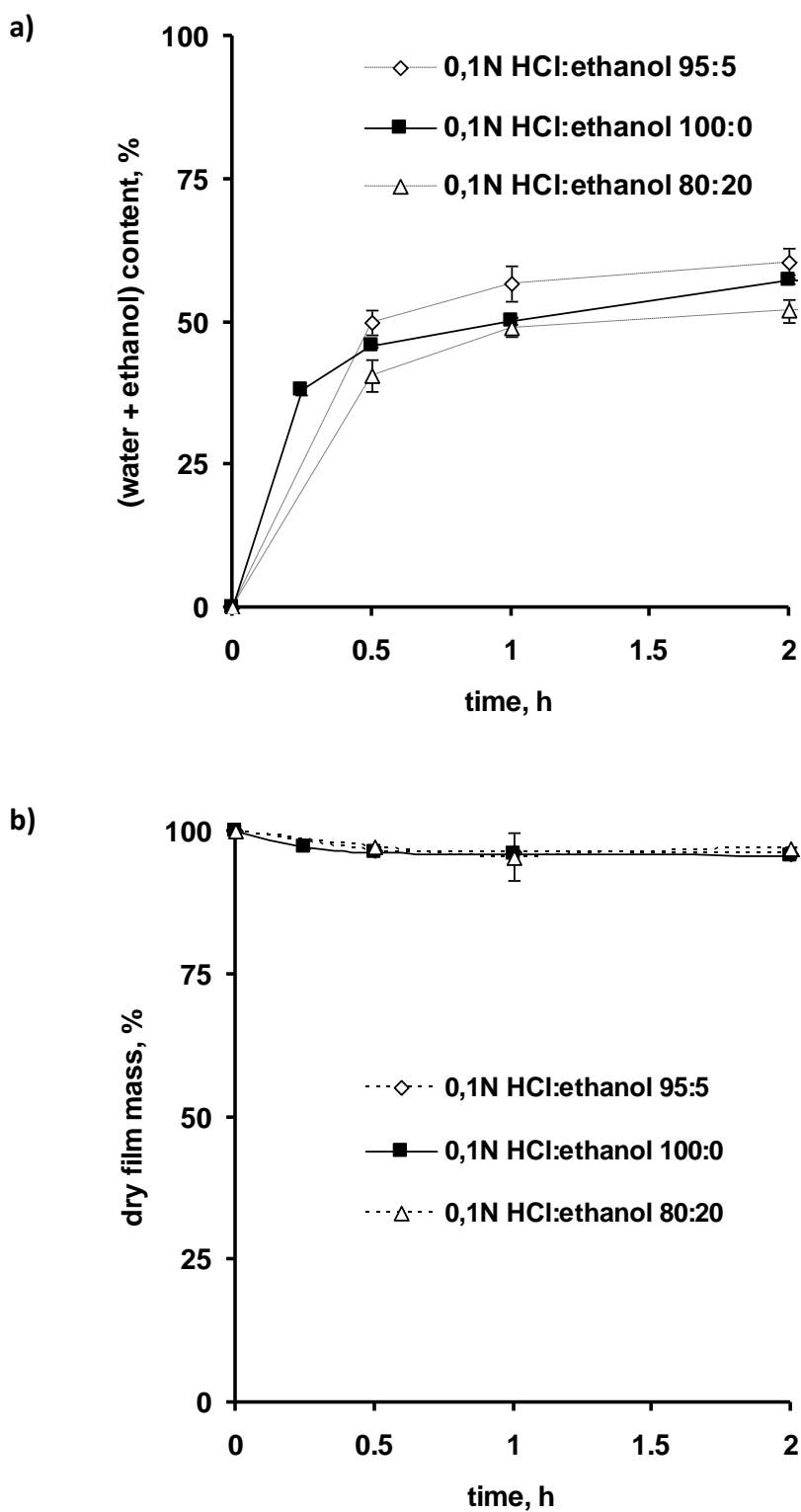


Figure 36: Impact of intermediate ethanol concentrations in the release medium on changes in the: **a)** (water + ethanol) content, and **b)** dry mass of thin, free (cast) 93:7 ethylcellulose:guar gum films upon exposure to a 100:0, 95:5, or 80:20 blend of 0.1 M HCl and ethanol (V/V) (as indicated).

6.2.6. Intermediate ethanol concentrations

All results presented so far were obtained upon exposure to 0.1 M HCl, or 0.1 M HCl:ethanol 60:40, respectively. However, in practice the ethanol concentration in the contents of the gastro intestinal tract of a patient is more likely to be in-between 0 and 40 % upon co-consumption of alcoholic beverages. Thus, it was important to see whether intermediate ethanol concentrations in the release medium might eventually affect the pellets' performance in a different way and/or alter the underlying drug release mechanism. Figures 35a and 35b show the experimentally measured theophylline release kinetics from pellets coated with 20 % 93:7 ethylcellulose:*medium* η guar gum, or 85:15 ethylcellulose:*high* η guar gum, respectively. The release medium was a 100:0, 95:5, 90:10, 80:20, or 60:40 blend of 0.1 M HCl and ethanol (V/V) (as indicated) for the first 2 h, followed by phosphate buffer pH 7.4 for the subsequent 6 h. Clearly, the impact of varying the ethanol concentration between 0 and 40 % was limited in all cases, irrespective of the investigated ethylcellulose:guar gum blend ratio and guar gum viscosity. These results were consistent with the experimentally measured water and ethanol uptake kinetics of thin, free films of identical composition as the film coatings: Figures 36a and 36b show the limited impact of varying the ethanol concentrations in the release medium on the resulting changes in the "water + ethanol" content and dry mass of thin 93:7 ethylcellulose:guar gum films. In all cases, the presence of the guar gum effectively minimized undesired ethylcellulose dissolution (and *vice-versa*) and the "ethanol and water" uptake kinetics were similar. Hence, neither the ethanol-resistance, nor the underlying drug release mechanism was altered at intermediate ethanol concentrations (corresponding for instance to the co-consumption of beer or wine, instead of high amounts of hard liquors).

6. 4. Conclusion

Drug diffusion through the intact polymeric film coatings seems to be the dominant mass transport mechanism controlling drug release from the recently proposed ethanol-resistant polymeric film coatings, based on ethylcellulose: guar gum blends. Hence, Fick's law of diffusion can be used to facilitate product optimization and to avoid time-consuming and

cost-intensive series of trial-and-error experiments. Depending of the type of drug, eventually also other phenomena might have to be taken into account.

7. Effects of film coating thickness on in-vitro drug release about sustained-release coated pellets: Using terahertz pulsed imaging

7.1. Materials and Methods

7.1 1. Materials

The following materials were used: metoprolol succinate (Salutas Pharma GMBH, Germany); nonpareil sugar starter cores (diameter 710-850 μm , NP Pharma SR, France); polyvinyl acetate (Kollicoat SR 30 D; BASF, Germany), poly(vinyl alcohol)-poly(ethylene glycol) graft copolymer (Kollicoat IR; BASF, Germany); hydroxypropyl methylcellulose (HPMC, Methocel E5; Colorcon, United Kingdom); triethyl citrate (TEC; Morflex, USA); and talc (Luzenca Val Chisone, Italy).

7.1 2. Preparation of the Pellets

Pellets were prepared by layering an aqueous drug-binder solution (20% metoprolol succinate, 1% HPMC) onto sugar starter cores (diameter = 710–850 μm ; Boire, France) until a 10% (w/w) drug load using a fluidized bed coater equipped with a Wurster insert (Strea 1; Aeromatic-Fielder, Switzerland). A spray rate of 2–3 g/min, a spray nozzle of 1.2 mm diameter and an atomisation pressure of 1.2 bar was used to apply the drug-binder solution. The inlet temperature was 40 ± 2 °C and the product temperature 38 ± 2 °C. The method used to apply the drug layer onto the sugar starter cores resulted in a visually smooth surface.

Kollicoat[®] IR (polyvinyl alcohol–polyethylene glycol graft copolymer) was dissolved in purified water and blended with plasticised Kollicoat[®] SR 30 D (an aqueous polyvinyl acetate dispersion) (overnight stirring with 5% triethyl citrate, w/w based on the polymer content) 30 min prior to the coating process. The polymer:polymer blend ratio was 25:75 (w/w referring to the dry mass). Talcum (1.5%) was added (w/w; based on the total solids content) and the dispersion was gently stirred throughout the coating process. The process parameters were as follows: inlet temperature 38 ± 2 °C, product temperature 35 ± 2 °C, spray rate 2–3 g/min, atomisation pressure 1.2 bar, nozzle diameter 1.2 mm. After coating, the pellets were further fluidised for 10 min and subsequently cured in an oven for 24 h at 60 °C. The metoprolol succinate loaded cores were coated until a coating thickness of

approximately 40, 60 and 100 μm (estimated based on weight-gain) was achieved. The overall minimum, maximum and the average diameter (measured in x and y directions for each pellet) from 10 pellets of each batch were determined using an electronic calliper (TESA Technology, Switzerland). The results were as follows: 830 μm , 1250 μm and 969 (± 95) μm for batch I; 940 μm , 1160 μm and 1043 (± 58) μm for batch II; and 990 μm , 1390 μm and 1144 (± 104) μm for batch III.

7.1.3. Terahertz Pulsed Imaging (TPI)

A total of ten pellets in each batch were imaged individually using a TPS Spectra3000 (TeraView Ltd, Cambridge, UK) in reflection mode. The single pellets were fixed on a glass slide and the terahertz incident beam was manually focussed on the highest point of the pellet surface by moving the x-y stage. The instrument set-up involved mapping a 1.2 x 1.2 mm area with a step-size (point-to-point mode) of 0.05 mm, and an axial penetration depth of 1.5 mm. The resulting spatial resolution with this analytical technique lies between 30-40 μm in the lateral direction and 200 μm in the axial direction (Zeitler et al., 2007a; Zeitler et al., 2007b). Data analysis was carried out using TPIView TVL imaging software version 3.0.5. The following equation was used to directly derive the CT from the terahertz waveform:

$$2d_{\text{coat}} = \Delta t c / n$$

where d_{coat} is the thickness of the coating, Δt is the time difference between the reflection of the terahertz pulse off the coating surface and the reflection from the interface of the coating and the drug layer; c is the speed of light; and n is the refractive index of the coating material (Ho et al., 2007). The refractive index (RI) used to calculate the CT was 1.5 (Ho et al., 2009) and the RI used to obtain the drug layer thickness was 1.81. The method of RI determination using terahertz pulsed spectroscopy in transmission mode has been previously described (Ho et al., 2007). In brief, the coating excipients and the drug were diluted with high density polyethylene at 5%, 10 % and 20% (w/w) concentrations and compressed into 13 mm diameter pellets under 2-ton pressure. Dilution was carried out to avoid signal saturation. Three pellets for every concentration were prepared and the RI spectra were generated using the TPS Spectra3000 (TeraView Ltd) in transmission mode. The

RI used for TPI was the average of the spectral RI over the bandwidth of the terahertz pulsed imaging system (Ho et al., 2007).

Due to the size and spherical nature of the pellets and the experimental set-up, the impact of pellet curvature on the observed thickness had to be considered and its effect minimised. The detected TPI parameter, terahertz electric field peak strength (TEFPS), measures the intensity of the reflected terahertz pulse from the surface of a sample normalised to the initial terahertz pulse intensity and is presented in % (Ho et al., 2008). It is a function of angle of incidence of the terahertz pulse with respect to the pellet surface, with it being a maximum when the pulse propagates in a direction normal to the pellet surface. However, due to the size of the pellets used in this study and the manual imaging set-up, the pellet curvature leads to the edges of the visible region being out of focus of the terahertz pulse and some terahertz radiation is also scattered away from the detector. This leads to significant signal loss towards the edges of the visible surface area of the pellets similar to the signal loss identified in a previous study on biconvex tablets (Ho et al., 2010), where the authors showed that the curvature of the centre band of a tablet can cause a signal loss that is reflected in the TEFPS values derived from the TPI maps. In addition, signal scattering and therefore signal loss was also demonstrated in another study on granulated materials in a size range comparable to the wavelength of the terahertz radiation (83 μm to 5 mm, Shen et al., 2008). A decrease in TEFPS due to pellet curvature with the current measurement set-up is associated with an increased risk of incorrect layer thickness results. Thus, the TEFPS values (minimum two thirds (3.0%) of the highest intensity (4.5%) of the reflected terahertz signal) were used to determine a circular area (0.13 mm^2) in the TPI maps, where the average thickness values could be accurately determined.

7.1.4. Dissolution Testing

Drug release measurements were carried out on the same pellets after TPI investigation. Pellets were placed individually into 2 mL Eppendorf-tubes filled with 1.5 mL phosphate buffer pH 7.4 for 8 h. The tubes were agitated in a horizontal shaker (80 rpm; GFL 3033, Gesellschaft für Labortechnik, Germany). At pre-determined time points (0, 0.5, 1, 2, 3, 4, 5, 6, 7, 8 h) the medium was completely removed and replaced. Analysis of the dissolution

medium was carried out using a UV-visible spectrophotometer (UV-1650, Shimadzu, France) at a detection wavelength of 222 nm. The average drug release of ten pellets was calculated and one-way analysis of variance (ANOVA) was used to compare drug release of the three batches at every time point of dissolution.

7.1.5. Scanning Electron Microscopy (SEM)

The TPI results obtained were compared with scanning electron microscopy (SEM) images (Hitachi S-3400, Hitachi High Technologies America, Inc., USA) of one pellet from each group. Images were taken of a cross-section and coating surface of the pellets at magnifications of 70× and 300×.

7.2. Results and discussion

The film coated pellets prepared in this study consisted of a sugar starter core coated with a drug layer (10 % (w/w)) and then different amounts of sustained release coating. The approximate coating thickness for the different coating amounts as estimated by the weight gain was 40 µm (batch I), 60 µm (batch II) and 100 µm (batch III). SEM micrographs taken of the surface of a pellet from every batch confirmed a smooth coating surface for all batches (see Figure 4.1 for an example). Prior to TPI investigations, 10 pellets were randomly chosen from every batch and the mean size was determined.

Using a pellet from batch II as an example, Figure 37 shows the 2D TEFPs map with the respective SEM image in which circles indicate the analysed area. The SEM image (figure 37b) revealed the surface of the pellets and hence the investigated area within the white circle was reasonably smooth in the entire visible area.

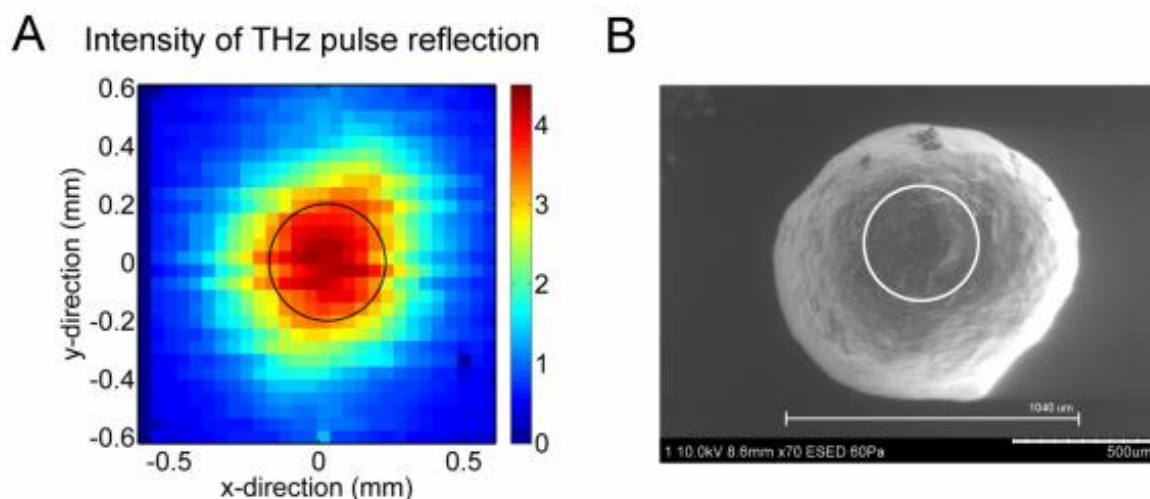


Figure 37. TEFPS map of a pellet from batch II is shown in image (A). The black circle indicates the pixels (measurement points) used for the calculation of the coating and drug layer thickness. The colour scale is in % reflected terahertz (THz) pulse intensity. The SEM micrograph (B) with the white circle also indicates the same area on the surface of a pellet from batch II.

At every pixel within the defined area a TPI waveform was recorded and interfaces between layers were resolved, providing information on the inner structure of the pellets. The interfaces between film coating and drug layer, and between drug layer and starter core were detected and visualised in the corresponding virtual cross-sectional images (images in the z direction; terahertz B-scans). A TPI B-scan together with a representative terahertz waveform of a pellet from every batch is shown in Figure 38. In the cross-sectional images (Figure 38a-c), the large red band located at a depth of 0 mm corresponds to the air/coating interface, followed progressively further to the right by a yellow/light blue band representing the coating/drug layer interface and a light blue band indicating the drug layer/core interface. The interface reflections are also visible in the waveforms as maxima of different intensities from left to right (Figure 38d-e).

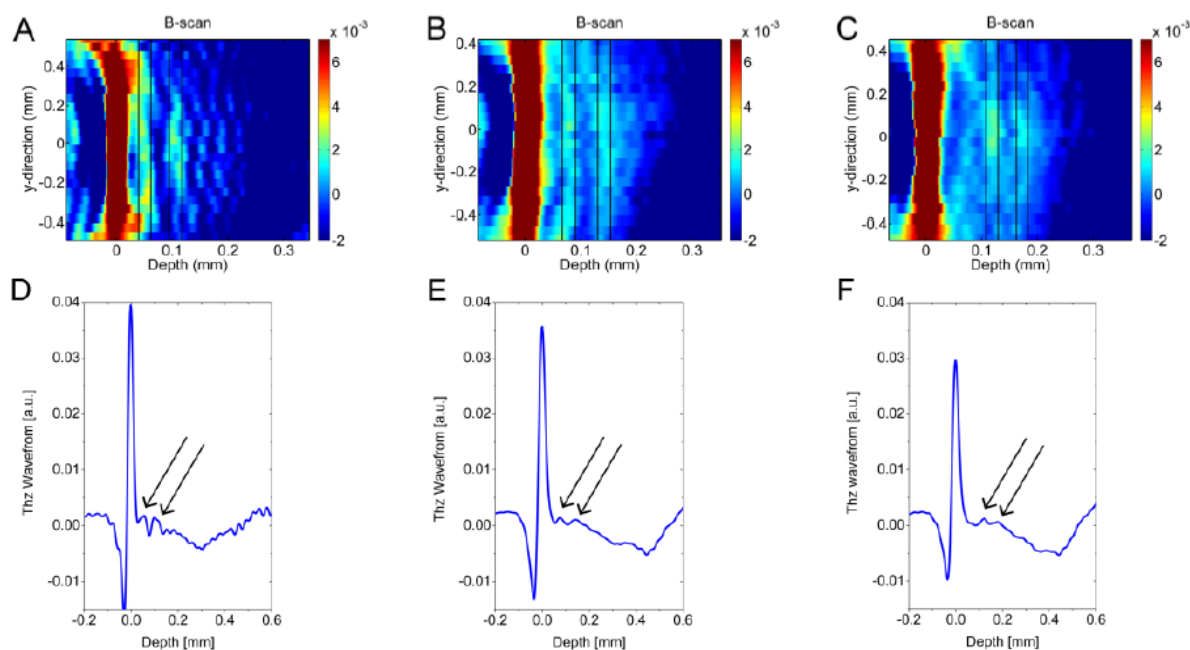


Figure 38: Terahertz cross-sections in the depth direction (B-scan) of a pellet from batch I (A), batch II (B) and batch III (C) with the respective terahertz waveforms (D, E and F). From the left in the cross-sections, the red band represents the air/coating interface, the yellow/light blue band the coating/drug layer interface (indicated by two blacklines), followed by a light blue band representing the drug layer/sugar core interface (indicated by two black lines). The first intensive positive maximum in the terahertz waveforms (D, E and F) relates to the air/coating interface, whilst the two black arrows indicate the maxima in the terahertz waveform corresponding to a signal reflection at the coating/drug layer and the drug layer/sugar core, respectively. The colour scale bar is the intensity of the terahertz signal in a.u.

Table 3.

Coating and drug layer thicknesses for all batches obtained by TPI. The mean thickness values and standard deviations (SD) are shown for the individual pellets (P1–P10) from every batch, with the overall mean thickness and SD for each of the batches below.

Batch I	Coating thickness (μm)	Drug layer thickness (μm)	Batch II	Coating thickness (μm)	Drug layer thickness (μm)	Batch III	Coating thickness (μm)	Drug layer thickness (μm)
P1	45 (±9)	55 (±8)	P1	74 (±6)	52 (±10)	P1	108 (±12)	56 (±10)
P2	40 (±12)	57 (±10)	P2	73 (±8)	57 (±10)	P2	108 (±11)	56 (±9)
P3	48 (±12)	53 (±10)	P3	70 (±8)	52 (±9)	P3	133 (±11)	49 (±9)
P4	46 (±11)	55 (±9)	P4	70 (±7)	54 (±9)	P4	122 (±14)	52 (±11)
P5	45 (±10)	54 (±11)	P5	69 (±7)	56 (±11)	P5	113 (±12)	54 (±10)
P6	50 (±10)	54 (±9)	P6	71 (±7)	51 (±9)	P6	108 (±13)	55 (±10)
P7	42 (±11)	55 (±9)	P7	73 (±8)	55 (±9)	P7	116 (±9)	53 (±9)
P8	44 (±10)	55 (±9)	P8	69 (±7)	53 (±10)	P8	113 (±14)	54 (±10)
P9	50 (±11)	52 (±10)	P9	70 (±6)	58 (±11)	P9	112 (±12)	57 (±10)
P10	49 (±10)	53 (±9)	P10	70 (±7)	52 (±9)	P10	111 (±14)	55 (±10)
Average	46 ± 3	54 ± 1	Average	71 ± 2	54 ± 2	Average	114 ± 8	54 ± 3

The results of the coating and drug layer thickness measurements from the corresponding TPI maps for all pellets are detailed in Table 3.

The average coating thickness for pellets from batches I, II and III was $46 \pm 2 \mu\text{m}$, $71 \pm 3 \mu\text{m}$ and $114 \pm 8 \mu\text{m}$ respectively. Two tailed, unpaired *t*-tests indicated statistically significant coating thickness differences between all batches (all *P* values well below the null hypothesis, $\alpha = 0.05$). In contrast, there was no difference in the observed drug layer thicknesses, with average thicknesses derived from the TPI maps of $54 \pm 1 \mu\text{m}$, $54 \pm 2 \mu\text{m}$ and $54 \pm 3 \mu\text{m}$ for batches I, II and III, respectively. These differences may also be identified by studying the terahertz B-scans and waveforms introduced above (Figure 38). The depth of the coating/drug layer interface varied between batches according to the thickness difference of the film coating. The depth of the following drug layer/core interface varied due to the different coating thicknesses even though the distance between the coating/drug layer interface band and the drug layer/core interface band visible in the B-scan images revealed similar thicknesses of the drug layer between batches.

The coating and drug layer thickness values both varied across individual pellets (high standard deviation (SD); Table 3). Such variation is likely to have been induced by the shape/surface morphology of the starter core. An uneven core surface can affect the drug layer uniformity, which in turn affects the coating uniformity (Heinicke and Schwartz, 2007).

Previously, it has been shown with TPI that the coating surface morphology and coating thickness is affected by the drug layer uniformity in large pellets (Ho et al., 2009). Layer thickness 2D maps constructed of the coating and drug layer thickness demonstrate the intra-pellet variations as well as the overall differences between batches (Figure 39). As mentioned previously, the waveforms recorded outside the circles may not be accurate and

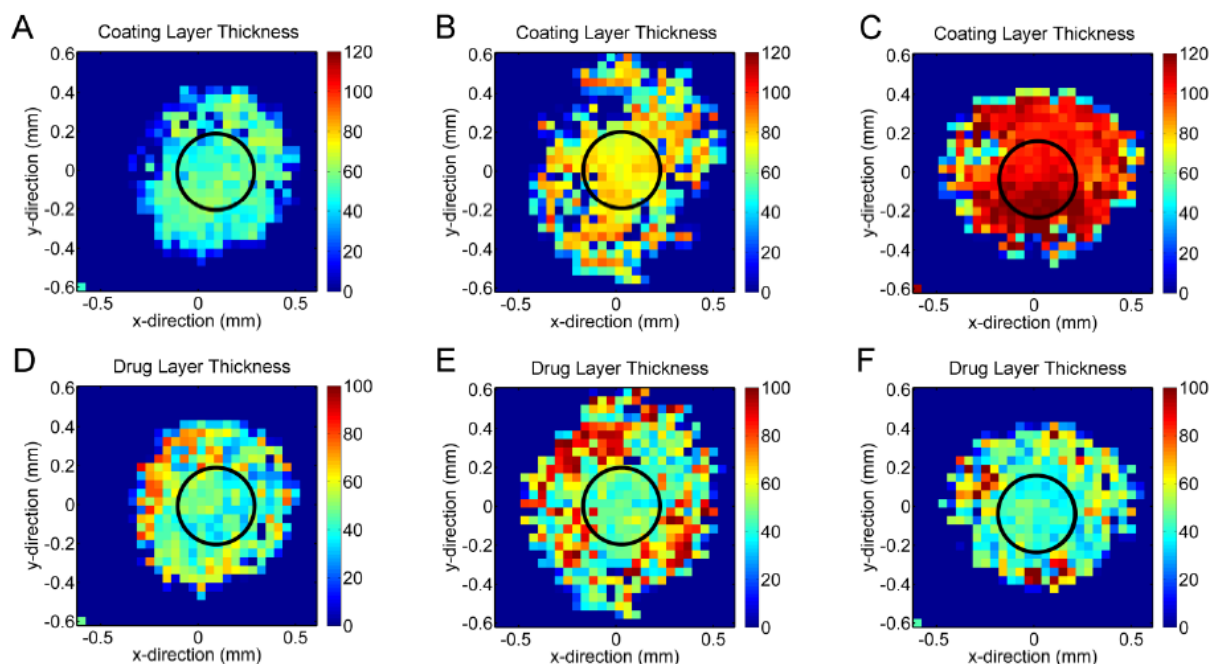


Figure 39: Coating (top) and drug layer (bottom) thickness, 2D maps of a pellet from batch I (A and D), Batch II (B and E) and batch III (C and F). The black circles indicate the areas used for coating and drug layer thickness calculation. Dark blue pixels shown in the CT 2D maps correlate with measurement failure and do not comprise a signal related to the coating/drug layer interface thus, they were excluded from the CT and DLT calculation. The colour scale bars are in μm .

Further indication for intra-pellet thickness differences and indication of variation of the starter core shape was obtained with SEM imaging of pellet cross-sections (Figure 40). The film coating was clearly visible in the micrographs, directly followed by the drug layer and the starter core. Although the drug layer/core interface was more difficult to identify using SEM, TPI investigation readily determined the interface. Nonetheless, the variation of the core shape can be seen in the SEM images hence, coating and drug layer thickness inconsistencies across individual pellets can be explained by irregular coating owing to the

starter core shape. However, the average coating and drug layer thickness for individual pellets within the same batch did not vary significantly with P values of 0.968, 0.987 and 0.356 (null hypothesis, $\alpha = 0.05$) for the CT and P values of 0.999, 0.994 and 0.994 (null hypothesis, $\alpha = 0.05$) for the DLT for batches I, II and III, respectively.

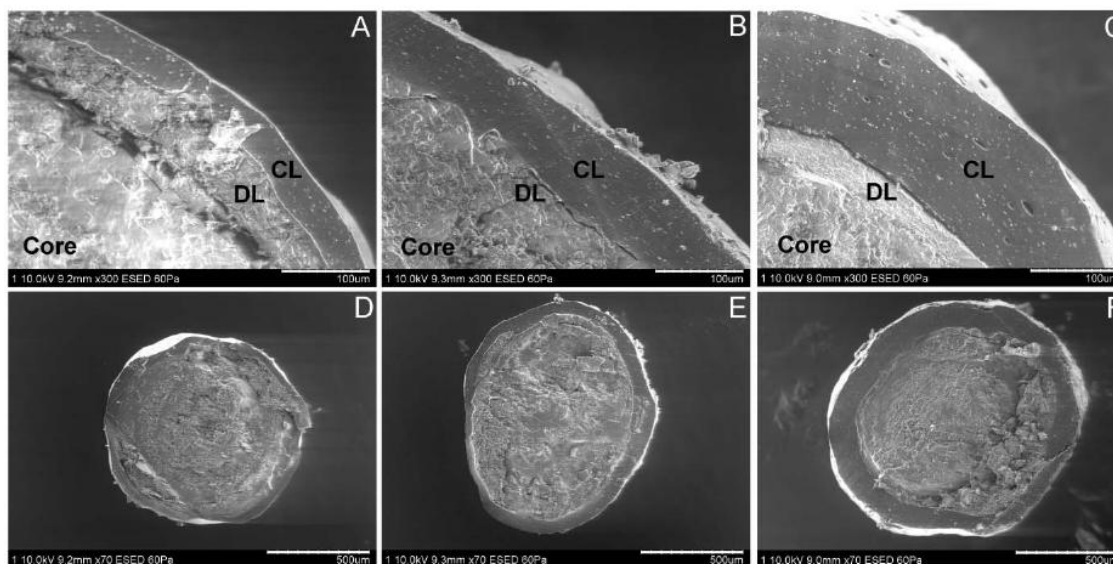


Figure 40: SEM micrographs of pellet cross sections showing the coating (CL) and drug layer (DL) (A, B and C) and the respective global images (D, E and F). Images were taken of a pellet from batch I (A and D), batch II (B and E) and batch III (C and F) and indicate differences in the coating thickness (CL) between batches.

Most importantly, correlation between the coating thickness and the drug release behaviour was determined. With increasing coating thickness of the pellets the drug release rate decreased (Figure 41). A one-way analysis of variance (ANOVA) of drug released from pellets in all three batches at every time interval was carried out. The average amounts of drug released after 0.5, 1, 2, 3, 4 and 5 h were significantly different between batches ($P < 0.05$). After 5 h, over 95% drug was released from all pellets under investigation and differences in the amount of drug release between batches were no longer significant. These results contrast with those from a previous study in which larger pellets containing theophylline were imaged with TPI (Ho et al., 2009) and differences in drug release rate as a function of coating thickness were not observed. However, in that study the drug release behaviour was controlled by the dissolution of the drug in the polymer coating and not its diffusion through

the coating. In the present study, the drug release was diffusion-controlled since the drug release behaviour is directly related to coating thickness.

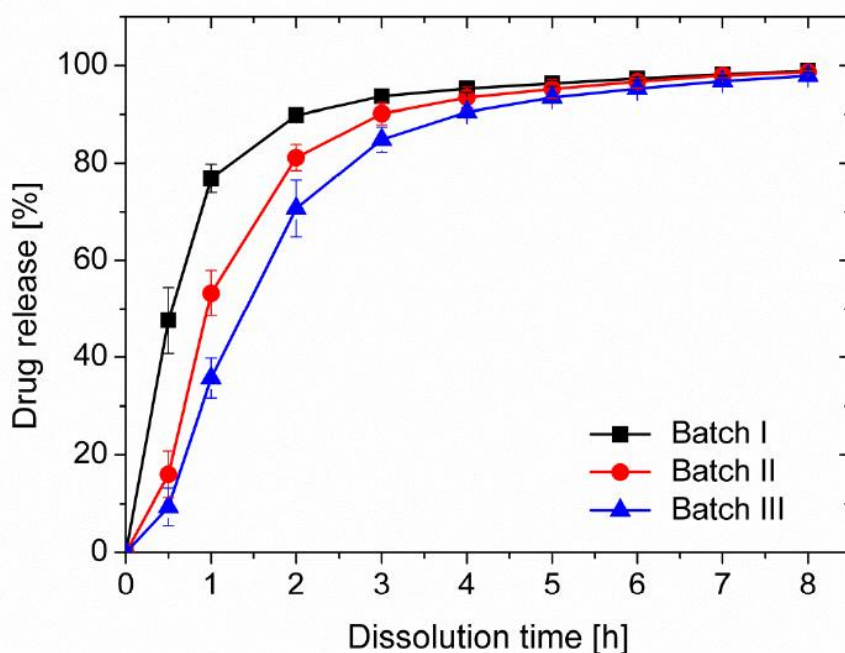


Figure 41. Average drug release profiles of 10 pellets from each batch. One-way ANOVA revealed that the drug release rate at 0.5, 1, 2, 3, 4 and 5 was statistically different between the batches.

7.3. Conclusion

TPI investigations were successfully carried out to image layers in standard size pellets for the first time. Despite the high curvature of the pellets due to their small size it was possible to analyse the drug and film coating layer thicknesses across individual pellets non-destructively. TEFPS values were used to determine a circular area in the TPI maps in which the pellet surface curvature did not adversely affect the accuracy of the coating thickness results. The coating and drug layer thickness across individual pellets was not uniform, but the mean coating and drug layer thickness was not significantly different between pellets within the same batch. Coating thickness differences were observed between batches (with different amounts of coating applied), whilst the thickness of the drug layer between batches was the same. These results were supported by SEM analysis. Moreover, the

thickness differences of the film coating between batches were correlated to the subsequent drug release behaviour of the pellets. Overall, the technique showed potential to non-invasively evaluate film coating characteristics of standard size pellets. A future fully automated measurement set-up may enable investigation of a larger number of standard sized-pellets.

8. Summary

In this PhD thesis, advanced drug delivery systems for time-controlled oral drug delivery have been investigated. Both, matrix as well as coated dosage forms have been prepared, thoroughly characterized and the underlying drug release mechanisms have been elucidated. In particular, lipid based matrix tablets, novel ethanol resistant polymeric film coatings and an advanced characterization method for film coatings have been addressed, as briefly described in the following.

A mechanistically realistic mathematical model is presented allowing for the quantification of niacin release from lipid tablets, based on glyceryl dibehenate. The systems were prepared either by direct compression or via hot-melt extrusion/grinding/compression. The model assumptions are based on a thorough physico-chemical characterization of the tablets before and after exposure to the release medium. Importantly, the model allows for the first time for the quantitative prediction of the effects of the composition, dimensions and type of preparation method of the tablets on the resulting niacin release kinetics. These quantitative theoretical model predictions were confirmed by several sets of independent experimental results. Furthermore, *in-silico* simulations revealed the fundamental importance of limited niacin solubility within the lipid tablets: During major parts of the release periods, very steep concentration gradients exist and net vitamin flux is restricted to specific regions within the tablets.

Furthermore, recently reported ethylcellulose:guar gum blends have been studied to provide ethanol-resistant drug release kinetics from coated dosage forms: Theophylline release from pellets coated with the aqueous ethylcellulose dispersion Aquacoat[®] ECD 30 containing guar gum was virtually unaffected by the addition of 40 % ethanol to the release medium. This is because the ethanol insoluble guar gum effectively avoids undesired ethylcellulose dissolution in ethanol-rich bulk fluids. However, the importance of crucial formulation parameters, including the minimum amount of guar gum to be incorporated and the minimum required guar gum viscosity, was not yet clear. The aim of this thesis was to identify the most important film coating properties, determining whether or not the resulting drug release kinetics is ethanol-resistant. Theophylline matrix cores were coated with various Aquacoat[®] ECD 30:guar gum blends, varying the polymer blend ratio, guar gum viscosity and degree of dilution of the final coating dispersion. Importantly, it was found that more than 5 % guar gum (referred to the total polymer content) must be incorporated in the

film coating and that the apparent viscosity of a 1% aqueous guar gum solution must be greater than 150 cP to provide ethanol-resistance. In contrast, the investigated degree of coating dispersion dilution was not found to be decisive for the ethanol sensitivity. Furthermore, all investigated formulations were long term stable, even upon open storage under stress conditions for 6 months. To elucidate the mass transport mechanisms controlling drug release from these systems, drug release from single pellets and ensembles of pellets was measured in various release media. Changes in the systems' morphology, composition and mechanical properties were monitored using SEM, gravimetric analysis and a texture analyzer. Based on the obtained experimental results a mechanistically realistic mathematical model was identified and used to quantitatively predict drug release from coated pellets in ethanol-free and ethanol-containing bulk fluids. Drug diffusion through the intact polymeric film coatings was found to be likely the dominant mass transport mechanism in the investigated systems, irrespective of the ethanol content in the surrounding environment. An appropriate solution of Fick's law could be used to quantitatively predict theophylline release from pellets coated with different ethylcellulose:guar gum blends at different coating levels. Importantly, independent experiments confirmed the theoretical predictions. *In silico* simulations can help facilitating the optimization of the novel ethanol-resistant polymeric film coatings, avoiding time-consuming and cost-intensive series of trial-and-error experiments. The presence/absence of ethanol does not affect the underlying drug release mechanisms.

In addition, terahertz pulsed imaging (TPI) was employed to explore its suitability for detecting differences in the film coating thickness and drug layer uniformity of multilayered, sustained-release coated, standard size pellets (approximately 1 mm in diameter). Pellets consisting of a sugar starter core and a metoprolol succinate layer were coated with a Kollicoat[®] SR:Kollicoat[®] IR polymer blend for different times giving three groups of pellets (batches I, II and III), each with a different coating thickness according to weight gain. Ten pellets from each batch were mapped individually to evaluate the coating thickness (CT) and drug layer thickness (DLT) between batches, between pellets within each batch, and across individual pellets (uniformity). From the terahertz waveform the terahertz electric field peak strength (TEFPS) was used to define a circular area (approximately 0.4 mm²) in the TPI maps, where no signal distortion was found due to pellet curvature in the measurement setup

used. The average CTs were 46 μm , 71 μm and 114 μm , for batches I, II and III respectively, whilst no DLT difference between batches was observed. No statistically significant differences in the average CT and DLT within batches (between pellets) but high thickness variability across individual pellets was observed. These results were confirmed by scanning electron microscopy (SEM). The CT results correlated with the subsequent drug release behaviour. The fastest drug release was obtained from batch I with the lowest CT and the slowest from batch III with the highest CT. In conclusion, TPI is suitable for detailed, non-destructive evaluation of film coating and drug layer thicknesses in multilayered standard size pellets.

9. Résumé

Dans cette thèse, les avancées concernant les systèmes oraux à libération contrôlée de principe actif ont été appréhendées. Aussi bien les systèmes matriciels que les formes enrobées ont été préparés, caractérisés et les mécanismes impliquant les phénomènes de libération du principe actif ont été élucidés. C'est le cas pour des comprimés à base de lipide, ainsi que d'un nouveau type de film d'enrobage montrant une résistance à l'éthanol et finalement une nouvelle méthode de caractérisation pour des enrobages polymériques a été mise au point, comme décrit brièvement ci-dessous.

Un modèle mathématique mécanique réaliste permettant de quantifier la libération de vitamines à partir de matrice lipidique, le glyceryl behenate a été élucidé. Deux techniques différentes de formulation : la compression directe et une suite d'extrusion en phase chauffante/ broyage/ compression directe ont permis la préparation de comprimés à base de Compritol 888. L'acide nicotinique a été utilisé comme principe actif modèle hautement soluble dans le milieu environnant. Un modèle théorique a été appliqué après l'observation des comprimés avant et après mise en contact avec le milieu de libération. La base de la production de comprimés, est d'une concentration identique en lactose et stéarate magnésium pour chaque lot et un taux variable en principe actif de 10 à 50%. Les profils de libération ont montré l'influence du taux de principe actif sur les cinétiques obtenues lié à un épuisement graduel de la vitamine et à une augmentation des chemins de diffusion. Les observations de Guse en 2006 montrent que les mécanismes impliqués lors de la libération à partir de la matrice lipidique sont liés au phénomène de diffusion. Par application d'un modèle mathématique basé sur la diffusion comme mécanisme de libération et sur la solubilité du principe actif, il est possible de prédire l'impact des paramètres de formulation et de fabrication sur les cinétiques de libération. Par fitting des résultats expérimentaux il est possible de déterminer le coefficient de diffusion noté D , représentant la mobilité du PA au sein de la porosité du comprimé. Connaissant ce coefficient de diffusion il est alors possible de prédire les cinétiques de libération, et ce pour des comprimés à taux de chargement variable, de tailles différentes et dépendamment de la méthode de préparation (figures 42 et 43).

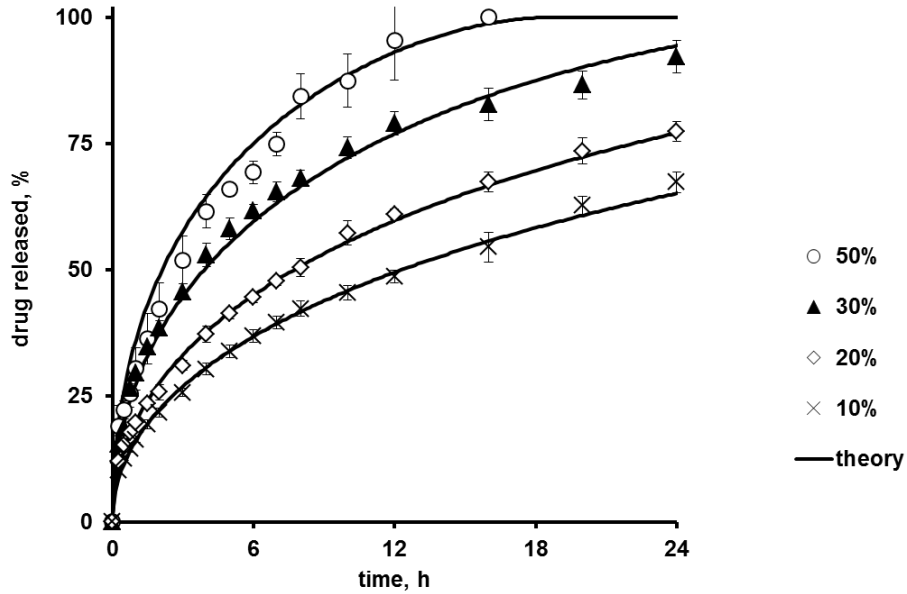


Figure 42 : Cinétiques de libération dans l'HCl 0.1N pour les comprimés obtenus par compression directe (taux de chargement de 10 à 50 %) Les symboles représentent les données expérimentales et les lignes pleines les théories prédites.

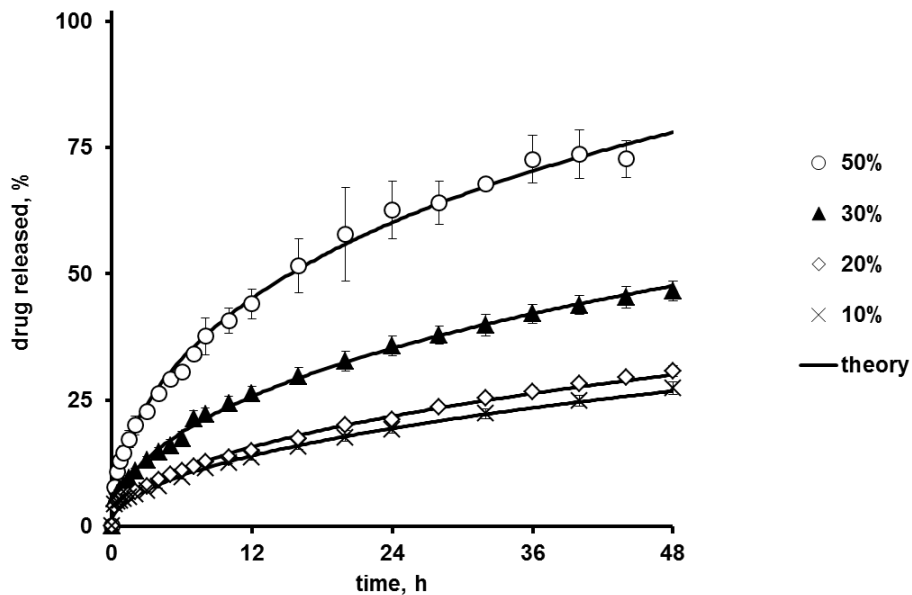


Figure 43 : Cinétiques de libération dans l'HCl 0.1N pour des comprimés obtenus par compression/broyage/compression directe (taux de chargement de 10 à 50 %). Les symboles représentent les données expérimentales et les lignes pleines les théories prédites.

Il a été récemment introduit qu'un mélange d'éthylcellulose: gomme guar permettrait une éthanol résistance. Des minigranules de théophylline enrobées par une dispersion d'éthylcellulose Aquacoat ECD 30 contenant une fraction de gomme guar ont montrées des libérations inchangées en contact d'un milieu avec une concentration d'alcool atteignant les 40% en volume les protégeant de tout contact avec l'éthanol présent dans le milieu de dissolution. Toutefois l'importance cruciale des paramètres de formulation, notamment la quantité minimale de gomme guar à ajouter et la viscosité minimale requise n'est pas claire. Il a donc été d'utilité de caractériser ce mélange dans le but d'avoir une concentration et une viscosité minimale de gomme guar pour éviter les effets de « collage » de ce dernier lors des processus d'enrobage, tout en permettant une concentration en gomme guar suffisante pour les profils souhaités mais surtout pour conserver l'éthanol résistance.

Des minigranules de théophylline ont ainsi été enrobées par plusieurs mélanges de type Aquacoat ECD 30 : gomme guar, variant tant au niveau du ratio du mélange, qu'au niveau de la viscosité et du degré de dilution dans la dispersion d'enrobage finale. Il a été trouvé que plus de 5% de gomme guar (par rapport au contenu polymérique total) doit être incorporé dans le film d'enrobage et une viscosité apparente pour une solution de gomme guar à 1% doit être supérieure à 150 cPs pour induire une éthanol résistance. A fortiori, les investigations concernant la dilution de la dispersion à enrober n'ont pas montrées d'influence sur le sensibilité à l'éthanol. L'ensemble des formulations ont montré une longue stabilité, cela même dans des conditions de stockage sous stress pendant 6 mois.

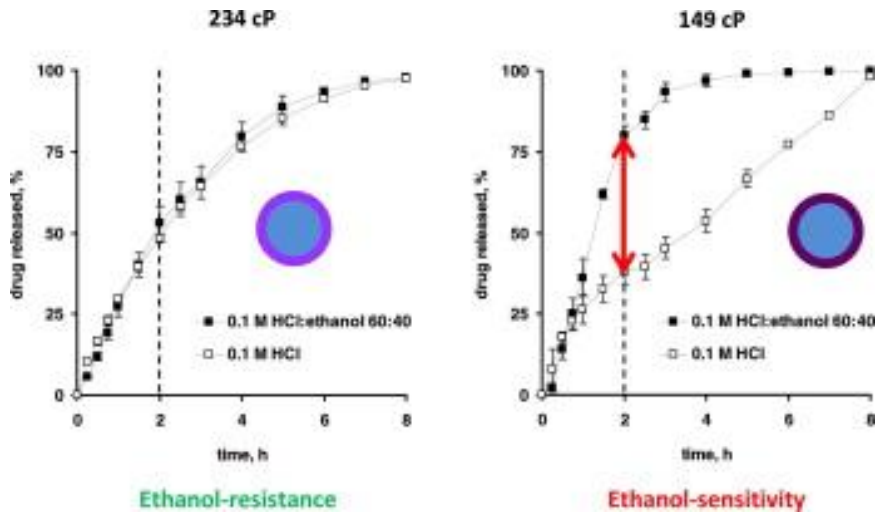


Figure 44 : Importance de la viscosité apparente de la gomme guar (234 cP: éthanol résistante vs 149 cP éthanol sensible) utilisée pour des enrobages de type 90 :10 éthylcellulose/gomme guar sur les profils de libération de théophylline des minigranules dans un milieu HCl 0.1N avec ou sans éthanol durant les deux premières heures suivi par le tampon phosphate pH 7.4 (20 % degré d'enrobage).

Pour élucider les mécanismes de transport de masses contrôlant les profils de libération à partir de ces systèmes, une libération à partir d'une minigranule ou d'un ensemble de minigranules a été mesuré dans différents milieux de libération. Des changements dans la morphologie des systèmes, dans la composition et dans les propriétés mécaniques ont été mesurés par l'utilisation de la MEB, des analyses gravimétrique et d'un analyseur de texture. Basé sur ces observations expérimentales, un modèle mathématique mécanistique et réaliste a été identifié et utilisé pour prédire quantitativement la libération de principe actif à partir de minigranules enrobées dans un milieu enrichi en éthanol et dans un milieu dépourvu en éthanol. Les phénomènes de diffusion au travers du film d'enrobage intact a été prouvé être le mécanisme de transport de masse dominant dans le système étudié, indépendamment de la présence d'éthanol dans le milieu environnant. Une solution issue des lois de Fick pourra être utilisée pour prédire quantitativement la libération de théophylline à partir de minigranules enrobées avec différents niveaux d'enrobage pour une gamme de mélange éthylcellulose :gomme guar variable. Les données expérimentales indépendantes confirment les prédictions théoriques. Les simulations *In-*

silico peuvent alors aider dans l'optimisation des films d'enrobages résistants à l'éthanol, éviter une consommation de temps et d'argent susceptibles d'intervenir par l'inutilité de formulations inadaptées.

L'investigation de l'impact des paramètres de formulation, à savoir l'influence du niveau d'enrobage sur les cinétiques de libérations et d'une autre part la détermination d'un indice permettant de déterminer l'homogénéité de répartition de l'enrobage à la surface des minigranules (TEFPS) a été mise au point. Dans ce but, des noyaux de sucrose sont préalablement enrobés d'une couche de principe actif le succinate de métoprolol, suivi d'une deuxième enrobage polymérique composé d'un mélange 75 :25 de Kollicoat SR :Kollicoat IR pulvérisé sur différents temps pour obtenir trois lots de minigranules, avec chacun une épaisseur d'enrobage différente en accord avec la prise en poids. 10 minigranules venant de chaque lot sont alors visualisées par Spectroscopie à Balayage afin de visualiser la morphologie des minigranules, et sur des coupes déterminer la taille de 1) l'enrobage de principe actif (DLT), 2) l'enrobage polymérique (CT). La même observation par l'imagerie Terahertz a permis l'évaluation de la CT et DLT au sein d'un même lot et entre les minigranules issues d'un même lot pour attester de l'uniformité. A partir des longueurs d'ondes terahertz, le TEFPS (terahertz electric field peak strength) est utilisé pour définir une aire circulaire (approximativement $0,4 \text{ mm}^2$) dans la carte de TPI. Aucune distorsion du signal n'a été rencontrée due à la haute courbure de la minigranule dans l'équipement de mesure utilisé.

La moyenne pour l'épaisseur de la couche polymérique est de $46 \mu\text{m}$, $71 \mu\text{m}$ and $114 \mu\text{m}$, pour les lots I, II and III respectivement, alors qu'aucune différence dans l'épaisseur de la couche de principe actif n'a été recensée. Au sein d'un même lot, aucune différence significative dans l'épaisseur de la CT et DLT n'a été mesurée, mais une haute variabilité dans l'épaisseur entre chaque individu a été observée. Les mesures de la taille du film polymérique pour les différents lots ont été corrélés avec les comportements de libération. Une rapide libération du principe actif a été obtenue pour le lot 1 avec une faible épaisseur de CT et une plus haute retenue avec l'épaisseur la plus importante. Pour conclure, TPI est une technique efficace dans l'évaluation non destructive de film d'enrobage et de couche de principe actif dans des systèmes multicouches de tailles standard.

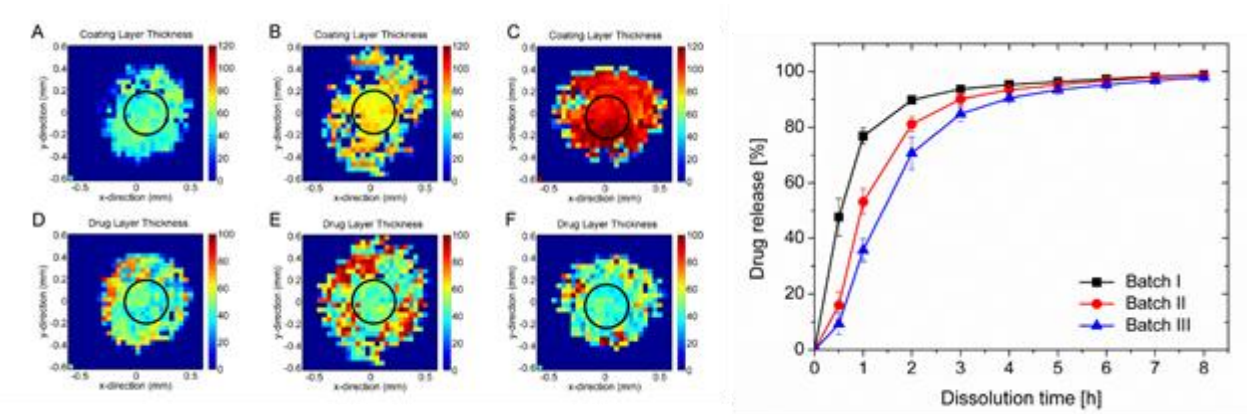


Figure 45 : Corrélation entre les données obtenues par TPI : épaisseur et homogénéité de l'enrobage de surface et celles obtenues par dissolution classique : profils de libération.