Application of UV dissolution imaging to pharmaceutical systems

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Application of UV dissolution imaging to pharmaceutical systems

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Abstract

UV-vis spectrometry is widely used in the pharmaceutical sciences for compound quantification, alone or in conjunction with separation techniques, due to most drug entities possessing a chromophore absorbing light in the range 190 to 800 nm. UV dissolution imaging, the scope of this review, generates spatially and temporally resolved absorbance maps by exploiting the UV absorbance of the analyte. This review aims to give an introduction to UV dissolution imaging and its use in the determination of intrinsic dissolution rates and drug release from whole dosage forms. Applications of UV imaging to non-oral formulations have started to emerge and are reviewed together with the possibility of utilizing UV imaging for physical chemical characterisation of drug substances. The benefits of imaging drug diffusion and transport processes are also discussed.

Keywords: Diffusion, dissolution testing, intrinsic dissolution rate, in-vitro release, parenterals, UV imaging

Abbreviations: API, Active Pharmaceutical Ingredient; ASD, Amorphous Solid Dispersion; UV, Ultra-Violet; CARS, Coherent Anti-Stokes Raman Scattering Microscopy; MRI, Magnetic Resonance Imaging; FaSSGF, Fasted State Simulated Gastric Fluid; FeSSIF, Fed State Simulated Intestinal Fluid; FTIR, Fourier Transforms Infrared; FVM, Focus Variation Microscope; CLSM, Confocal Laser Scanning Microscopy; NMR, Nuclear Magnetic Resonance; THz, Terahertz; NIR, Near Infra-red; XµT, X-ray micro-Tomography; PAT, Process Analytical Tools; SEM, Scanning Electron Microscope; CMOS, Complementary Metal Oxide Semiconductors; CCD, Charged Coupled Devices; GIT, Gastro-Intestinal Tract; DIDR, Disk Intrinsic Dissolution Rate; BCS, Biopharmaceutics Classification System; MCC, Microcrystalline Cellulose; SDS, Sodium Dodecyl Sulfate; PCL, Poly-ε-Caprolactone; SDI, Surface Dissolution Imaging HPC; Hydroxypropylcellulose; LOQ, Limit of Quantification; pI, Isoelectric Point;
1. Introduction

Traditional drug dissolution and release testing methods are based on measurements of the active pharmaceutical ingredient (API) in the bulk solution or in samples withdrawn from the medium, which upon accumulation often is analysed by a UV spectrophotometric technique. The measurement approaches may be intrusive and disturb the release process without offering significant insights into drug release mechanisms. Thus, there has been a drive towards real-time analysis and continuous monitoring methods to determine and better understand dissolution and drug release phenomena. This has led to utilization of fibre optic UV probes and the implementation of advanced imaging techniques in dissolution and release testing.

Advanced imaging techniques applied within the pharmaceutical industry can provide useful real-time information to understand critical behaviour of APIs. Imaging techniques that have found use in drug development include Coherent Anti-Stokes Raman Scattering Microscopy (CARS) (1), Raman spectroscopy and imaging (2), (3), Magnetic Resonance Imaging (MRI) (4), (5), X-ray micro-Tomography (XµT) (6), Fourier Transform Infrared (FTIR) imaging (7), (8), Near Infra-red (NIR) imaging (9), (10), fluorescence imaging (11), Terahertz (THz) imaging (12), (13), Confocal Laser Scanning Microscopy (CLSM) (14), and UV dissolution imaging (15), (16).

Briefly, CARS has been used in the imaging of the solid-state changes of APIs in situ in dissolution to correlate with dissolution rate change (1), (17) the release of API from pharmaceutical tablets (18). Raman imaging has been used to assess excipient distribution within a tablet (19), (20) for chemical imaging of nasal spray suspensions and API distribution within a tablet (21), (22). NMR/MRI has been used to study the internal mechanisms of the in vitro drug release in dosage forms such as the ingress of dissolution media (23), (24), (25), (26), (27), water distribution in extrudates (28), water diffusion coefficients and polymer concentrations (29), (30), (31), (32), water penetration in osmotic pumps (33), homogeneity in powder mixtures (34), the tablet disintegration process (35), internal barriers for zero-order release systems (36), pulsatile capsules (37) and mapping the distribution of porosity in compacts (38). XµT has been used to measure/assess heterogeneity of powder beds (39), (40), density variations in compacts (41), (42) with and without embossed features (43), (44), porosity of hot-melt extrudates (45), compression characteristic of dry coated tablets (46), packing in granular systems (47), (48), excipient homogeneity (49), visualization of tablets after drug release (50). FTIR imaging has allowed dissolution studies of tablet-in-tablet
formulations (51), within-tablet dynamics for extended release formulations (52) and pharmaceutical films (53). The depletion of API and the hydration processes for hydrophilic matrices (54), (55), (56) as well as films (57) and pharmaceutical product characterisation (58) have been visualised using NIR imaging. Fluorescence imaging has been applied to the swelling of polymers to investigate molecular diffusion and interactions between chromophores and solvent molecules (59) (60), (61), (62). Recent developments in semiconductor physics and ultrafast laser technology have made it possible to provide light at terahertz frequencies which now allow the analysis of coating thickness, structure and uniformity and intra-tablet coating uniformity (12), (63), (64), (65), (66), (67), chemical component calibration and density mapping (68), (69), (70), inspection of pharmaceutical tablets and quality control (71), (72), dissolution assay for immediate-release tablets (73), and determining porosity and non-contact weight measurements (74), (75), (76). CLSM has been utilised in a number of different pharmaceutical applications such as the visualisation of the early gel layer formation in tablet matrices (77), (78), visualisation and identification of the percolation threshold (79), visualisation of drug release from milk and fat-rich emulsions (80) and the designing of matrices with improved resistance to dissolved sugar and dietary sugars (78), (81). Optical microscopy has also been used to visualise and determine intrinsic dissolution rates (IDRs) from single drug particles as small as 14 μg (82), (83), (84) These methodologies have wide-ranging applications from the early drug development phase, characterisation of drug delivery systems, process measurements, as process analytical tools (PAT), product performance testing and quality control.

UV dissolution imaging is an emerging technology that provides visualisation of the dissolution of a material and subsequent related events at the solid-liquid interface as well as obtaining a concentration measurement. The insights offered into API behaviour will be useful in providing an understanding for phenomena not usually captured in offline measurements. UV dissolution imaging, sometimes referred to as UV imaging, UV/Vis imaging or surface dissolution imaging, has offered a compound sparing approach to the determination of IDRs. However, additional UV imaging applications in relation to oral as well as non-oral formulations and physical chemical characterization have started to occur. Here we aim to give an introduction to UV dissolution imaging covering the basic principles of the technology as well as current applications to pharmaceutical systems, hoping to spur new ideas and wider uptake in pharmaceutical research. The review addresses visualization of dissolution phenomena in relation to form selection, to determination of intrinsic dissolution rates, and to
drug-excipient compatibility in early drug development. These relatively best-established
applications of UV dissolution imaging, are followed by drug release studies from whole
dosage forms, which have become available due to recent instrumentation developments.
Likewise, applications of UV imaging to non-oral formulations are starting to emerge and are
reviewed together with the utilization of UV imaging for physical chemical characterization of
drug substances. Advantages and limitations of the technology are discussed with a perspective
to the positioning of UV imaging relative to more advanced imaging modalities used in drug
development.

2. Principles of UV imaging

Most pharmaceutically relevant applications of the UV/Vis spectrophotometry is based on light
in the wavelength range 190 nm to 800 nm, with UV light ranging from 190 to 400 nm and
visible light from 400 to 800 nm (85), (86). The ability of drug substances to absorb light forms
the basic principle of UV/Vis spectrophotometry and UV/Vis imaging. The absorption takes
place when an electron is promoted to a higher energy state by the energy of an incident photon
where the extent of the light absorbed by the compound in the UV/Vis spectrum is determined
by the chromophores present in the molecule. The main application of spectrophotometry is
quantification of compounds present in solution. However, different molecular structures will
yield different absorbance spectra, which may be useful as a secondary method in compound
identification (86). The quantification is governed by absorbance of light according to the Beer-
Lambert law (Equation 1) stating that the amount of light absorbed is directly proportional to
the concentration of absorbing species present in the medium.

\[
\log_{10} \left( \frac{I_0}{I} \right) = \log_{10} \left( \frac{1}{T} \right) = A = \varepsilon bc
\]

where \( I_0 \) denotes the incident light intensity, \( I \) is transmitted light intensity, \( T \) specifies
transmittance given by \( \frac{I}{I_0} \), \( b \) is the light path, \( A \) is the absorbance, \( \varepsilon \) is the molar absorption
coefficient, and \( c \) denotes the concentration of the absorbing substance.

The Beer-Lambert’s law only applies to monochromatic radiation. However, in practice,
polychromatic radiation is applied, which may cause deviation from the Beer-Lambert’s law
(85), (86), (87). Other causes for deviation from the Beer-Lambert law relate to the
instrumentation, for instance stray light or may be attributed to properties of the analytes. High analyte concentrations may lead to refractive index changes, chemical reactions, self-association or interactions between molecules (e.g., drug – surfactants interactions) in solution can lead to deviations from linearity. Given the absence of interactions affecting the absorbance spectra, the Beer-Lambert law applies to mixtures of molecules or analytes. This and, in particular, the proportionality, between absorbance and concentration, make UV/Vis spectrophotometry and imaging convenient and relatively simple means for quantification of drugs. Spatially and temporally resolved absorbance measurements are used in the creation of images from UV (and Vis) imaging (16), (88).

2.1. UV Imaging instrumentation

The UV imaging systems encountered in this review utilise a CMOS based image detector to record images. The application of a phosphor coating produces a spectra shift from 200 – 400 nm into the visible range and thus allowing detection (89). Limitations for conventional CMOS detectors include higher noise, lower image quality, lower sensitivity and a limited dynamic range, however the recent development CMOS detectors has addressed many of these limitations, making CMOS detectors suitable for scientific applications, which can perform well in the UV range (90), (91).

The earliest applications of UV imaging were in the area of separation sciences, electrophoretically mediated microanalysis and capillary electrophoresis, using the Paraytec Actipix D100 detector (92), (93), (94), (95). Quartz cells were developed for the imaging detector allowing development of pharmaceutical applications (96). The design of the imaging system and the relatively small imaging area (7 × 9 mm²) was suited for compound sparing small-scale dissolution studies, leading to IDR measurements to be among the first applications to emerge (97), (98). Recent developments have been in the direction of larger imaging areas and multi wavelength imaging allowing studies of larger and more complex samples (16). The interested reader can get an appreciation of the commercial instruments iterations that have occurred and instrument specifications from Table 1 and the literature (16), (99), (100), (101). Taking into account that different instruments are available and assuming further instrument developments, the description of UV dissolution imaging instrumentation below is limited to a generic level.
Table 1. System overview of the commercially available surface dissolution imaging equipment

<table>
<thead>
<tr>
<th>INSTRUMENT</th>
<th>IMAGING AREA</th>
<th>PIXEL SIZE</th>
<th>RESOLUTION</th>
<th>LIGHT SOURCE</th>
<th>COMMENTS</th>
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<tr>
<td>D100</td>
<td>7 x 9 mm²</td>
<td>7 x 7 µm²</td>
<td>40 µm x-direction 30 µm y-direction (99)</td>
<td>Pulsed xenon flash lamp 190 – 1100 nm via bandpass filter</td>
<td>Standalone CMOS detector</td>
</tr>
<tr>
<td>SDI 300 (ALSO KNOWN AS SDI OR SDI1)</td>
<td>7 x 9 mm²</td>
<td>7 x 7 µm²</td>
<td>40 µm x-direction 30 µm y-direction (99)</td>
<td>Pulsed xenon flash lamp 190 – 1100 nm via bandpass filters</td>
<td>First purpose-built surface dissolution imaging system, using syringe pump</td>
</tr>
<tr>
<td>D200 (VARIOUS CONFIGURATIONS)</td>
<td>28 x 28 mm²</td>
<td>13.75 x 13.75 µm²</td>
<td>30 µm (both direction) (102)</td>
<td>LEDs: 255 nm, 280 nm, 320 nm, 525 nm</td>
<td>Standalone CMOS detector</td>
</tr>
<tr>
<td></td>
<td>11 x 11 mm²</td>
<td>5.5 x 5.5 µm²</td>
<td>ND²</td>
<td>Pulsed xenon flash lamp 180 – 800 nm via bandpass filter</td>
<td></td>
</tr>
<tr>
<td></td>
<td>11 x 6 mm²</td>
<td>5.5 x 5.5 µm²</td>
<td>ND²</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SDI2</td>
<td>28 x 28 mm²</td>
<td>13.75 x 13.75 µm²</td>
<td>ND²</td>
<td>LEDs: 255 nm, 280 nm, 300 nm, 320 nm, 520 nm</td>
<td>Fully integrated system, containing CMOS detector and automated pumps. Compact flow cell and whole dosage cell configurations.</td>
</tr>
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</table>

ND² - Not experimentally determined

The key components of a UV dissolution imaging system are a light source, a quartz cell for containing the sample of interest and an imaging detector (Figure 1). The system has to be shielded from outside light because the phosphor coated CMOS chip is sensitive to light in both the UV and Vis wavelength range. Wavelength selection is achieved by LEDs emitting...
light of a narrow wavelength range or the use of band pass filters in conjunction with a pulsed Xenon lamp. Key distinguishing features between early instrumentation and more recent instruments are the size of the imaging area (currently up to $28 \times 28 \text{ mm}^2$, $2048 \times 2048$ pixels) and the ability to perform multi wavelength imaging (up to 4 wavelengths) or not. In addition, dedicated cells and pump systems for medium handling differentiate the systems. The schematic in Figure 1 shows the most typical configurations for a UV imaging system (Pion SDI2) with two different flow cells, the so called compact cell (Figure 1a) and the whole dose cell (Figure 1b) to image compacts (IDR determinations) and formulated full dosage forms, respectively. The compact cell is specifically designed to image compacts made from API powders to determine the intrinsic dissolution rate (IDR). The design of this cell resembles the classical Nelson and Shah cell (103), providing laminar flow condition across a solid surface of compacted material (3 - 10 mg). The whole dose cell is based on a USP apparatus IV-like flow cell used to image dissolution and release of API from whole dosage forms such as tablets and capsules (104), (105), (106), (107). The cell is cylindrical (Figure 1b) with dimensions $142.5 \text{ mm} \times 27.5 \text{ mm} (H \times D)$ with a conical shape which holds glass beads, has a volume of $60 \text{ mL}$. The imaging area is $28 \text{ mm} \times 24 \text{ mm}^2 (H \times W)$ and optical path length is variable $\leq 28 \text{ mm}$ due to the cylindrical shape (108).
Figure 1. Schematic representations of a UV imaging system for (a) intrinsic dissolution rate (IDR) measurements and (b) whole dosage form imaging. The photographs (I) and (II) show the Pion SDI2 compact flow cell and (b) whole dose set-up, respectively. The inserts show representative UV images obtained during dissolution testing. The effective image area is 28 × 28 mm² and the instrument allows selection of two wavelengths at a time: 255, 280, 300, 320, and 520 nm (108). Reprinted from (105). Copyright (2019) with permission from Elsevier.
3. Small-scale dissolution and intrinsic dissolution rate measurements

Dissolution testing commonly constitutes a significant element of more or less complex setup designed to mimic the physiology of the gastrointestinal tract (GIT). Reviews of existing in-vitro models for predicting in-vivo performance suggest that there is no one method that can fit all needs with respect to the methodology chosen. Methodology chosen can be dependent on factors such as requirements for the development phase and the drug delivery system (109), (110), (111).

Most of the traditional or conventional in-vitro dissolution methods such as the rotating disc, paddle, and basket methods often require relatively larger amounts of dissolution medium, API or formulation. Due to the limited amount of API present during the early stages of drug development, miniaturised or micro-dissolution techniques that require only few milligrams of the API are of interest as they do overcome some of these limitations (97), (111), (112), (113), (114), (115), (116). This drive towards miniaturisation has produced instrumentation that uses either offline or real-time analytics, including the mini-paddle apparatus (117), (118), mini dissolution vessel (119), 96-well plate assay for intrinsic dissolution testing (MINIDISS) (120), transfer method to predict precipitation (121), solvent shift method (122), µDISS using UV fibre optics (112), miniaturised rotating disk intrinsic dissolution test (115), µFlux test (123), Partially Automated Solubility Screening (PASS) assay (124), miniaturized assay for solubility and residual solid screening (SORESOS) (125), SiriusT3 and inForm (111), and UV dissolution imaging (97), (98), (126). The advantage of these miniaturised techniques is that they are classed as compound-sparing meaning each IDR determination typically requires 5-10 mg of sample, or less. This compares favourably to traditional IDR determinations such as the Woods apparatus, which often require in the order of hundreds of milligrams of an API. Thus, IDR determination via a miniaturised method is an attractive option in the early stages of development, where when access to API often is limited.

Solid drugs have been characterised using disk intrinsic dissolution rate (DIDR) for many years (112), (127), (128). As DIDR is a rate phenomenon and not an equilibrium phenomenon, it is thought to correlate more closely with in-vivo drug dissolution dynamics than solubility. Consequently, it has been suggested that DIDRs be used to classify drugs instead of solubilities as used in the Biopharmaceutics Classification System (BCS) (129).

Intrinsic dissolution rate ( IDR ) is defined as the rate of mass transfer per a unit area which is exposed to dissolution media (130). Throughout an experiment, the IDR surface area should
be kept constant, and the use of a constant flow of dissolution media (in the case of UV dissolution imaging) applied to maintain sink conditions.

The pharmacopoeia method described for IDR determinations include the rotating disk system (or Wood’s apparatus). The IDR, which is expressed by the Nernst-Brunner’s adaptation of the Noyes-Whitney equation and commonly obtained from the modified USP I/II set up, takes into consideration the concentration of the API dissolved, sampling time, diffusion coefficient, surface area, volume of the medium, boundary layer thickness, and the concentrations at saturation at the solid surface and in the bulk solution (131).

The Nernst-Brunner and Noyes-Whitney equations constitute a convenient frame for discussion of dissolution processes (occurring in vessels or vials). For flow-through based UV dissolution imaging, however, the convective diffusion drug dissolution model developed by Nelson and Shah (132), (103), (133), serves as the quantitative basis. The dissolution rate ($R$) is represented by Equation 2 according to the convective diffusion dissolution model in case of a circular compact. Using the model of Nelson and Shah and taking into account the two-dimensional images captured by UV imaging, IDR values may be calculated according to Equations 3 and 4 (133).

$$R = 2.157 \cdot D^{2/3} \cdot c_0 \alpha^{1/3} \cdot r^{5/3} \quad \text{Eq.2}$$

$$IDR = \frac{\sum_{z=0}^{H} \left( v_z M c_z W \Delta z \right)}{S} \quad \text{Eq.3}$$

$$v_z = \frac{3Q}{2HW} \cdot \left( 1 - \left( \frac{2z-H}{H} \right)^2 \right) \quad \text{Eq.4}$$

where $D$ is the diffusivity, $c_0$ is the solubility and $r$ is the compact radius, $z$ is the height above the $z$-origin, $v_z$ is the velocity at $z$, $M$ is the molecular weight, $c_z$ is the concentration at $z$, $W$ is the width of flow cell channel, $\Delta z$ is the effective pixel height, $S$ is the surface area of sample, $H$ is the height of flow cell channel in the observation region and $Q$ is the volumetric flow rate (133).

3.1. Characterisation techniques used with UV dissolution imaging

The dissolution behaviour of an API is intimately related to its solid-state properties. A comprehensive characterisation of the solid form properties is essential because drugs may have a propensity to undergo solvent mediated solid form transitions during dissolution.
It is therefore commonplace to characterise the solid-state properties of compound, prior to IDR determination. Most studies achieve this via the use of in-situ or offline Raman spectroscopy, differential scanning calorimetry (DSC), X-ray powder diffraction (XRPD), scanning electron microscopy (SEM) and focus variation microscopy (FVM) (Table 2).

One area of particular success is the ability to assess the interplay between solvent mediated solid form transitions and drug dissolution (97), (134), (135), (136) (Table 2). A few studies have reported the use of in-situ Raman spectroscopy measurements and UV dissolution imaging data to be collected simultaneously (137), (136), (138). Scanning electron microscopy (SEM) or optical microscopy have been used to qualitatively assess the quality of the drug compacts prior to IDR determinations. The need to generate quantitative data describing surface properties of the compacts used for IDR determinations has also spurred the incorporation of focus variation microscopy (FVM) as a tool for characterizing surface topography (139), (140). This technique, which stems from the field of precision engineering, has the ability to accurately measure the surface topography of drug compacts. This allowed for the exploration of how manufacturing conditions affected the surface of the drug compacts and how the dissolution process itself changed the topography of the drug compact (139), (140).

3.2. UV dissolution imaging and IDR determination

A typical image generated by a UV imaging dissolution instrument is depicted in Figure 2. The dissolution of indomethacin is apparent as the increase in absorbance at 320 nm. The indomethacin dissolved from the compact is carried with the flow exiting the flow cell on the right hand side (Figure 2). The flow conditions appear to be laminar. The swelling and dissolution-precipitation phenomena that some substances are subject apparently did not occur under the experimental conditions applied. As mentioned above, quantitative information may be extracted from the images captured. Data analysis software typically allows for correction of changes in medium absorbance (reference zone), measurement of the drug concentration immediately above the compact surface using the so-called surface absorbance zone, and quantification of the amount of drug dissolved per area and time unit from the IDR absorbance zone (Figure 2). The software also allows additional absorbance measurement zones to be positioned in any area within the image field. The IDR is calculated from the model and
equations presented in section 3. Consequently, IDR measurements by UV imaging are prone to error when the hydrodynamic model does not match the actual flow conditions, for instance, due to density effects, gelling or swelling of the API or precipitated material affecting the hydrodynamic pattern, or when the Beer-Lambert law is not adhered to (141), (136).

Figure 2. Image from the analysis software of the UV dissolution imaging instrument (Pion SDI2). The image shows the dissolution of indomethacin from solid compact into phosphate buffer (pH 7.2) at a flow rate of 2 mL/min at 37 °C. Detection at 320 nm. Modified and reprinted from (105). Copyright (2019) with permission from Elsevier.

### 3.2.1. Polymorphs, salts and form selection

The application of flow-through UV dissolution imaging for dissolution studies was first proposed by Wren and Lenke (96). The first study to report intrinsic dissolution rates was performed on amlodipine besylate (97). Two amlodipine besylate forms (amorphous and dihydrate) as well as the free base form were subjected to UV dissolution imaging in combination with off-line Raman spectroscopy measurements. The IDRs were in the order amorphous amlodipine besylate > amlodipine besylate dihydrate > amlodipine free base. The decrease in the IDR of the amorphous form observed was due to a solvent mediated solid form
transition as confirmed by Raman spectroscopy. During dissolution, API swelling or recrystallization at the compact surfaces (the amorphous and free base forms) was visualized and recorded by UV imaging in real time (97) (Figure 3 I and II).

Li and co-workers expanded the application of UV imaging dissolution to pharmaceutical cocrystals using carbamazepine as the API of interest and nicotinamide as the cocrystal former (134), (142). Qiao et al. utilised UV imaging to gain an understanding of the intrinsic dissolution behaviour of carbamazepine-nicotinamide cocrystals (134). For comparison, dissolution studies were conducted on carbamazepine dihydrate, carbamazepine forms I and III, nicotinamide as well as a physical mixture of carbamazepine and nicotinamide. Each sample was subjected to SEM analysis at various points with attempts made to visualise the dissolution effects on the sample surface topography. Offline Raman spectroscopy was used to evaluate solution mediated solid form transformations; this showed that the cocrystal, the two polymorphic forms and the physical mixture all underwent conversion to carbamazepine dihydrate during dissolution and that this was associated with a decrease in IDR as determined via UV imaging. Of interest was the slower IDR decrease for the cocrystals as compared to the pure drug and physical mixture. This was confirmed to be due to a slow recrystallisation of carbamazepine cocrystal to carbamazepine dihydrate during dissolution. Qiao et al. demonstrated the ability to monitor dissolution processes over prolonged periods of time (up to 3 h). Most other UV imaging-based dissolution studies followed the dissolution process for 10-20 min (97), (98), (135), (143). The influence of sodium dodecyl sulfate (SDS) and Tween 80 addition to the dissolution medium below and above the critical micelle concentration on the dissolution of carbamazepine–nicotinamide cocrystals, carbamazepine III, and carbamazepine–nicotinamide physical mixture was investigated (142). SDS was found to significantly increase the IDR, whereas Tween 80 addition lead to a decrease of carbamazepine IDR from the co-crystal. Characterisation of the solid-form showed solution-mediated solid form transformation into carbamazepine dihydrate to occur during the dissolution process.
Figure 3. (I) UV absorbance maps of dissolution at static conditions of (a) amlodipine besylate dihydrate, (b) amorphous amlodipine besylate and (c) amlodipine free base. Contour lines: blue, 400 mAU, green, 800 mAU, black, 1200 mAU and light blue, 1600 mAU. (II) Light microscopy photographs of the compacts containing (a) amlodipine besylate dihydrate, (b) amorphous amlodipine besylate and (c) amlodipine free base prior to and after imaging. The red arrows indicate crystal growth beyond the boundaries of the cylindrical compact. (d) Representative photo of compact within the channel flow cell showing crystal growth of amorphous amlodipine besylate upon dissolution imaging at static conditions. (III) Selected absorbance maps for the dissolution of sodium naproxenate in 0.01 M HCl containing 5.35 × 10⁻⁵ M thymol blue studied by dissolution imaging at 1.0 ml min⁻¹ and 550 nm. Intense red color indicates high absorbance and the contours represent the iso-absorbance/iso-pH lines. Reprinted in modified form with permission from (97). Copyright (2011) American Chemical Society. Reprinted from (144). Copyright (2014) with permission from Elsevier.
Hulse et al. showed how a range of parameters such as the polymorphic form and particle size affected the dissolution of indomethacin, theophylline and ibuprofen (98). UV imaging was able to successfully distinguish between the dissolution behaviour of alpha and gamma indomethacin, anhydrous and monohydrate theophylline and ibuprofen sodium and free base. Theophylline anhydrate exhibited a higher dissolution rate when compared to its monohydrate form. Sodium ibuprofen was also shown to exhibit a higher rate of dissolution as compared to ibuprofen in phosphate buffer (pH 7.2). When the effects of micronisation of indomethacin were studied, the unmicronised powder, which had larger particles, resulted in a faster dissolution for the alpha indomethacin as compared to the micronized powder. This phenomenon was explained as the change in surface energy due to micronisation bringing about an increase in the proportion of hydrophobic surfaces. X-ray powder diffraction, offline Raman spectroscopy and dynamic vapour sorption (DVS) were used to confirm the solid-state forms of the drugs.

Østergaard and coworkers pioneered the use of UV imaging with in-situ Raman spectroscopy for characterising solvent-mediated solid form transitions during dissolution testing (136), (137), (138). The dissolution behaviour of naproxen sodium and theophylline anhydrate and monohydrate with the simultaneous use of UV imaging and in situ Raman spectroscopy, showed that decreases in their dissolution behaviour could be attributed to form changes where naproxen sodium and theophylline anhydrate both converted to their stable forms. The form conversion for theophylline anhydrate as detected by Raman spectroscopy occurred within 5 min (136). For the naproxen system, two intermediate forms were detected prior to formation of the stable form, which would readily have been overlooked by offline Raman spectroscopy. The use of in-situ Raman spectroscopy in combination with UV imaging seems to offer an interesting approach, which may be beneficial in preformulation and early drug phase development as it facilitates the understanding of basic dissolution behaviour and solvent-mediated solid form transitions.

Salt formation is an attractive option for improving solubility and bioavailability for ionisable drug candidates (145), (146), (147). The solubility of twenty different Danirixin salts was determined with the hydrobromide salt giving the largest solubility improvement (148). The effects of reduced gastric pH were investigated using simulated gastric fluid at pH 1.6, sodium phosphate, pH 3.0, and potassium citrate at pH 4.0. Dissolution studies showed improved dissolution rate for the hydrobromide salt as compared to the free base. This was most pronounced at pH 4.0, where a 6-fold increase was observed. The clinical study confirmed
these findings with the hydrobromide salt having a greater bioavailability when compared to the free base (148). A proof of concept setup was developed for mapping the local or microenvironmental pH change in relation to the dissolution of sodium naproxenate in 0.01 M and 0.10 M hydrochloric acid using the dissolution imaging a wavelength in the visible range and a pH indicator (thymol blue) (144) (Figure 3 III). The dissolution of the sodium naproxenate in 0.01 M hydrochloric acid showed that sodium naproxenate could significantly change the microenvironment pH within the vicinity of the solid salt compact. The in-situ pH measurement is conceptually simple but requires some tailoring during method developments in terms of dye selection (pK$_a$, pH, concentration and wavelength). The dissolution performance of gemfibrozil and various salts with different alkyl chain lengths cyclopropylamine (CPROP), cyclobutylamine (CBUT), cyclopentylamine (CPENT) and cyclohexylamine (CHEX) were investigated showing all salts to provide increased IDR relative to its free form (the free base). The results indicated that the longer alkyl chain length of the counter ion led to lower dissolution rates as compared to their shorter chain length counterparts (149).

Etherson et al. conducted an inter-laboratory investigation to assess variability in IDR determinations of drugs with low (tadalafil, bromocriptine mesylate), medium (carvedilol, indomethacin) and high (ibuprofen, valsartan) solubility using UV imaging (130). The dissolution experiments were conducted in FaSSIF and blank FaSSIF at pH 6.5. These experiments were specifically conducted on the surface dissolution instrument (SDI) with Actipix™ Technology from Sirius Analytical, Forest Row, UK (now Pion Inc.). This study found that in general the rank order of mean IDR reported across the six participating sites for each drug was similar to the solubility rank in both FaSSIF and blank FaSSIF. The variability in measurements, however, ranged from 33% to 130%, with the highest variability observed for the lowest solubility drugs. IDR data for indomethacin were selected for further analysis with the mean IDR data from a single operator compared to the site IDR data. A large reduction in the standard deviation could be achieved if a single operator was used for data analysis. Similar high variability related to IDR measurements was also found in another inter-laboratory study using the miniaturised dissolution instrument, the µDiss profiler (150). These studies illustrate that, irrespective of standardised protocols for data collection the generation of repeatable dissolution data is far from trivial.
Biorelevant dissolution media had little effect on the dissolution rate of furosemide (135). It was observed that some samples had pronounced irregular upward shifts of the contour lines in UV absorbance maps, which may be an indication of dissolution followed by precipitation associated with solid form transformation or swelling of the solid material. In-situ Raman spectroscopy measurements were conducted and together with X-ray diffraction confirmed that this was not due to solvent mediated solid form transformation (showing the difficulty associated with inferring form changes from dissolution rates and visual characteristics). These observations warranted further studies of the dissolution behaviour of amorphous furosemide sodium salt and amorphous furosemide (acidic neutral species) in simulated bio-relevant intestinal medium at pH 6.5 (151) along with its corresponding crystalline forms (152). In-situ Raman spectroscopy measurements were employed to gain an insight into solvent mediated form changes. This study also included IDR determinations conducted on a µ-Diss profiler for comparison purposes. IDR determinations for the four furosemide samples showed that the two salt forms had a higher dissolution rate when compared to the acid forms, with the highest dissolution rate observed for the amorphous salt form. Fast conversion of the two amorphous forms to trihydrate forms was shown by both a decrease in dissolution rate and changes in Raman spectra.

Niederquell and Kuentz determined the IDR of celecoxib, ketoprofen, naproxen, and sulfathiazole in simulated gastric biorelevant media at pH 6.5 (143). High and variable initial drug dissolution rates observed by the UV imaging dissolution, which was followed by decreasing dissolution rates with time. This was attributed to sample heterogeneity. High and variable initial dissolution rates have also been observed in other studies (97), (98), (135). Niederquell and Kuentz modelled the heterogeneity effects using double logarithmic plots of the dissolution rate versus time and fractal-like dissolution behaviour as the profiles adhered to a power law relation. Although the initial phase was dominated by effects of surface heterogeneity, the ensuing phase could be viewed as a pseudo-equilibrium meaning an intrinsic drug dissolution rate could be inferred from this range (143). The sample heterogeneity was confirmed using SEM (Figure 4 I and II). This study depicted how heterogeneity-related effects could affect dissolution experiments on small-scale dissolution systems. Other authors have used similar models to evaluate solvent-mediated changes of APIs; in solid dispersions systems, dosage forms as well as surfaces of nanocrystals (153), (154), (155), (156) (157). Sample surface heterogeneity has been characterised quantitatively to provide information on
the surfaces for compacts prior and post UV dissolution using focus variation microscopy (FVM) (139), (140) (Figure 4 III and IV). Focus variation microscopy was employed with the view of establishing a relationship between drug compact surface properties and dissolution performance (139). The surface parameter of Sdr, which gives an estimation of changes in surface area was of particular interest and showed a general decrease of the surface area by the dissolution process (this however was a decrease from the “post flush” to the end of the dissolution process). The IDR also decreased as the experiment continued, with higher IDR values reported during the first 10 min as opposed to IDR values reported after 20 min as observed previously (143). Such studies show that drug compact surface properties may impact IDR determination. An investigation into the influence of compaction force on three different API (ibuprofen, ketoprofen and paracetamol) compacts compressed against different surfaces showed that generally, higher pressures resulted in a flatter, smoother compacts as reported by decreases in the Sdr parameter. When compacts were made on a manufactured relatively smoother surface for both the ibuprofen and paracetamol, their compacts however experienced increases in the Sdr parameter indicating a larger surface area to be associated to the compacts. This suggested that the ability of the drugs to form smoother drug compacts may be API dependent (140). The change in surface properties however, did not seem to significantly affect the dissolution rate, suggesting the solubility of the drug to be a predominant factor in the dissolution process (140).
Figure 4. Scanning electron microscopy images (×500 magnification) of (I) celecoxib and ketoprofen before and after 5 min of dissolution in biorelevant medium (FaSSIF) at 37 °C. (II) naproxen and sulfathiazole before and after 5 min of dissolution in biorelevant medium (FaSSIF) at 37 °C. (III) Focus variation (Alicona™) 3D data set images of representative compacts of IBU (a) post compaction, (b) post flush, (c) after 10 min run on SDI, (d) after 20 min run on SDI. Note: diameter of sample holder in image is 2.5 mm (measured using a 150 mm digital calliper gauge from Draper Tools Ltd, UK). (IV) 3D images of the roughness of the original (a), and the manufactured plate insert surfaces (b), both used in the production compacts for IDR determinations. 3D images of the surfaces of IBU compacts at varying compressions of forces using the original press surface (c). Reprinted from (139, 140 and 143). Copyright (2014, 2017, 2020) with permission from Elsevier.

3.2.3. Co-amorphous systems

Co-amorphous systems present an attractive approach for stabilising amorphous compounds which can be prone to recrystallization, especially when stored in humid conditions or above their glass transition temperature (107), (158), (159). These systems as compared with their respective crystalline forms have no long-range crystallographic order and higher internal energy (160), (161), (162). Recently, UV dissolution imaging with in-situ Raman spectroscopy was used to investigate the solvent-mediated transformation in the co-amorphous system composed from naproxen and naproxen sodium during dissolution (138). Solid-state analysis using XRD, DSC and FTIR showed that the 1:1 naproxen and naproxen sodium salt co-amorphous system was stable for 2 months at 40 °C. For comparison, dissolution studies were also conducted on a physical mixture, sodium naproxen and naproxen (free acid). UV imaging showed that the dissolution rate of the co-amorphous system was between that of the crystalline free acid and the sodium salt. Although a co-amorphous was successful produced, it somewhat surprisingly did not lead to any improvement in dissolution rate over the pure sodium salt. In-situ Raman spectroscopy measurements revealed initial phase separation of the co-amorphous form to naproxen free acid and sodium naproxen followed by disproportionation of the sodium salt and crystallization to free acid naproxen. This to date has been the only study of its kind, however, UV imaging with simultaneous in-situ Raman spectroscopy may provide further detailed insights into these systems.
3.2.4. Single crystal and nanocrystal dissolution

An early UV imaging study visualized the dissolution of lidocaine from a single crystal in stagnant phosphate buffer, pH 7.4 (126) (Figure 5). Imaging at 254 nm revealed dissolved lidocaine falling towards the bottom of the quartz cell due to natural convection caused by the local density gradients formed from dissolution of the highly soluble lidocaine. Quantitative estimates of the amount of lidocaine dissolved from the crystal were obtained from the UV images. The natural convection can be efficiently suppressed by imbedding the crystal in a gel matrix; symmetrical diffusion patterns around similar lidocaine crystals placed in agarose gels have been reported (99). Single crystal dissolution and, in particular, face specific dissolution of single crystals may provide fundamental knowledge regarding the mechanism of crystalline API dissolution. Apparently, it has proved difficult to progress from the initial studies cited above to face specific dissolution measurements using UV imaging. Among the challenges encountered are crystal size, resolution and difficulty to resolve crystal faces from dissolved compound. At present, other approaches seem more promising as relates to face specific dissolution (163), (164), (165), (166), (167). The key interest from pursuing UV/Vis imaging in this area should relate to the ability to attain information about both the dissolved drug (UV wavelength) and the solid (Visible wavelength) in a single measurement.

One of the techniques to improve the bioavailability of poorly soluble drugs is the use of the nanocrystal approach (168), (169). UV imaging alongside a traditional flow-through dissolution method showed dissolution improvements of indomethacin nanocrystals prepared using a top-down wet milling technique with poloxamer F68, poloxamer F127 and polysorbate 80 as stabilizers, as compared to unmilled indomethacin and their physical mixtures (170). The initial UV and Vis imaging, however, showed significant absorbance increases during the flow-through dissolution at 265 nm and 550 nm. This was explained by nanocrystals liberating from the surface of the compact into the dissolution medium scattering and absorbing light at 550 nm. A hydrogel matrix was used in the suppression of these particles “escaping” from the compact surface. This resulted in the nano-crystal compacts having a significantly higher dissolution rate as compared to unmilled indomethacin (five times the solubility in the immediate vicinity of the nanocrystal compact, which may be due to supersaturation). It is important to note that the poloxamer F68 and F127 might also have local solubilising effects on indomethacin.
Figure 5. (I) (A) Microscope photograph (the image is $8.3 \times 6.4 \text{ mm}^2$) and (B) absorbance maps of the lidocaine crystals arranged in a quartz cell obtained by the Sirius SDI imaging system with pixels binned $4 \times 4$ at 254 nm. The image is $10.4 \times 4.6 \text{ mm}^2$. (C) Absorbance – y-distance profile of the lidocaine crystal placed in the x-direction (●) and absorbance – x-distance profile for crystal placed in the y-direction (●) for determining the width of the crystals from the absorbance maps and (D) normalized absorbance – distance profile of one side of a lidocaine crystal placed in the x-direction (●) and y-direction (●) for estimating the resolution of the Sirius SDI imaging system. . (II) Time-dependent absorbance contour maps of the dissolution of lidocaine crystals in (A) 0.5% (w/v) agarose gel, pH 7.4 and (B) 0.067 M phosphate buffered solution, pH 7.4. The dissolution was performed in quartz cells with 1 mm light path using the Sirius SDI UV imaging system with the pixels binned $4 \times 4$ at 254 nm and the sensor head placed in the upright position. Reprinted from (126). Copyright (2016) with permission from Elsevier.
Polymeric strip-films constitute a versatile drug delivery platform that has recently received significant attention and are capable of incorporating active substances in thin polymer matrices that allow its delivery upon oral administration (171), (172), (173). The feasibility of using a drug-loaded hydrophilic (HPMC) polymeric layer sandwiched between two hydrophobic poly-ε-caprolactone (PCL) layers for improving film drug load whilst achieving sustained release of fenofibrate was investigated (174). Thinner films followed Fickian diffusion, and thicker films followed non-Fickian anomalous diffusion as modelled from drug release rates obtained using the USP IV flow-through cell and a UV dissolution imaging system. Changing the thickness of the middle layer allowed manipulation of drug load without the need for altering the formulation composition. Fenofibrate nanocrystals were incorporated into oral strip-films of variable thickness and compared to a marketed formulation (Tricor) (175). *In-vivo* pharmacokinetic properties of the different thickness oral films were also investigated using rabbits. The bioavailability of the nanocrystal films was improved by a factor of 1.4 to 1.8 fold as compared to the marketed formulation. Solid-state analysis suggested that fenofibrate nanocrystals were successfully incorporated into the oral strip-films and the crystalline state remained unchanged and stable. UV dissolution imaging results qualitatively showed the swelling associated with the films as well as visualisation of the drug release. Quantitatively, the UV dissolution imaging showed that the fastest dissolution occurred using the thinnest film and was comparable to the dissolution studies using the USP-IV dissolution apparatus.

3.3. Drug-excipient compatibility

Although limited structural information can be obtained from UV-Vis spectra as compared to other chemical imaging techniques, UV dissolution imaging may allow for visualisation of dissolution phenomenon such as drug-polymer interactions, differentiation between polymer grades, viscosity effects of formulated products and ranking of salts. Here the applications of UV imaging involving excipients is discussed.

UV imaging was used to visualise the swelling behaviour of compacted 15 cP and 50 cP viscosity grades of hydroxypropyl methylcellulose (HPMC or hypromellose), a hydrophilic polymer widely used to the control API release in oral formulations under flow as well as under stagnant conditions (176) (Figure 6 I). UV imaging at 214 nm was able to detect polymer entanglement, swelling and eventual gel formation instrumental in controlling drug release.
The experimental set-up provided sufficient sensitivity for differentiating between the two viscosity grades of HPMC, showing a faster and more pronounced swelling with HPMC 50 cP compared to 15 cP. The swelling behaviour of 3D printed cellulosic tablets matrices for disintegration visualized by UV imaging revealed that cellulose matrix swelling was between 0.2 – 0.4 mm. The spacing gaps around the 3D printed cellulosic tablets matrices would therefore be ineffective once exposed to dissolution medium as they would be closed off as the polymer swelled. Results from the USP apparatus II testing supported the swelling closed gap hypothesis observed from the UV dissolution imaging, as the 3D tablets with the narrowest gaps of 0.2 mm showed the slowest drug release and longest time to fragment (177).

Amorphous solid dispersions (ASDs) constitute a widely used approach for improving the solubility and bioavailability of poorly soluble APIs. Here, the crystalline drug is amorphisized or molecularly dispersed within a synthetic or naturally water-soluble polymer such as polyethylene glycol (PEG), soluplus, polyvinylpyrrolidone (PVP) or hydroxypropyl methylcellulose (HPMC) (178), (179), (180), (181), (182), (183), (184) (185), (186), (187). ASDs of nilotinib were prepared with the polymer hydroxypropyl methylcellulose phthalate (HP55) (as well as with HPMC). The produced amorphous nanodispersions were investigated using UV dissolution imaging with in-situ Raman spectroscopy (137). Two nilotinib nanoparticle dispersions of 20 % and 40 % drug load and physical mixtures were investigated while invoking a pH change from pH 2.2 to 6.4 during the UV dissolution imaging. Extensive physicochemical testing on the dispersions produced showed that the nilotinib drug load had little effect on the physical state of the API but had significant effects on the dissolution properties, with the higher drug load showing increased dissolution rate when compared to the ASDs with 20 % drug load, the pure drug and both physical mixes. This enhancement was attributed to the stabilising effects of hydrophobic and H-bond interactions, which, particularly at higher drug loads prevented the formation of crystalline nilotinib. Dissolution experiments were also conducted using USP apparatus II and it was observed that a marked decrease in dissolution rate at 30 min was observed when the pH change occurred. This was, however, less profound in the UV imaging testing (137). Gautschi et al. investigated whether increases with respect to IDR could be achieved by solid dispersions prepared by the addition of either mono or diacyl lecithin phospholipid to a series of poorly soluble drugs (BCS class II and IV) (188). Phospholipid:drug 1:1 molar ratios were investigated. The ability of the drug substances to form solid dispersions with the phospholipids were related to the API’s physical chemical parameters such as the frontier orbital energy, enthalpy of fusion, and log P. UV imaging results
showed an improvement in IDR when using the monoacyl lecithin phospholipid as compared to the diacyl lecithin phospholipid and pure drug.

Figure 6. (I) Images of HPMC 15 cP in stagnant medium (A) and flow of 0.5 ml/min (B), and HPMC 50 cP in stagnant PBS (C) and in presence of flow (D) at time points of 0, 15, 50 and 90 min. The white colour represents absorbance caused by the polymer, the contour line corresponds to an absorbance threshold of gel point, i.e. 3.2% (w/w) for HPMC 15 cP, and 2.3% (w/w) for 50 cP, and the rectangle designate the area, in which the analysis was performed. (II) Images captured of the web-like phenomenon within the IDR flow through cell taking place in the UV experiments. Absorbance values (Abs) of the studied superdisintegrant types and brands as a function of distance from the centre of the sample cup (mm) in (III) 0.1 N HCl pH 1 (SDI1, 0 mL/min, 37 °C, 254 nm) presented up to 2.5 min. (IV) in phosphate buffer
pH 6.8 (SDI1, 0 mL/min, 37 °C, 254 nm) up to 2.5 min. Reprinted from (176, 105 and 189).

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Indomethacin-Soluplus ASDs prepared by spray drying, freeze drying and using a homogenising-freeze drying technique were also assessed using UV dissolution imaging (105). It was observed that the samples with the lower polymer content had improved drug release rates as revealed by flow-through UV dissolution imaging. The higher ratios of Soluplus resulted in the formation of a web-like pattern of polymer swelling within the cell (Figure 6 II). These formed web-like strands, which attached to the top of the flow cell and may have resulted in the decrease in drug liberation from the compacts.

The UV dissolution imaging effects of pure or amorphous forms of cefuroxime axetil and itraconazole using FaSSGF and FaSSIF-V1 biorelevant dissolution media and simulated gastric fluid without pepsin (SGF) and fasted state simulated intestinal fluid without pancreatin (SIF) were investigated (190). The drug formulations Zinnat tablets (cefuroxime axetil) and Sporanox capsules (itraconazole) were used to provide the amorphous forms of the drug substances and were crushed and made into a compact in preparation for UV imaging testing. This, however, proved challenging for Zinnat as the presence of a superdisintegrant resulted in a physical blockage of light in the flow cell. The Sporanox formulation contained HPMC, which swelled and thereby had an effect on drug release from the HPMC polymer matrix. For the compacts under investigation, the higher flow rates resulted in an increased drug release from the compact surfaces. The pure drug, itraconazole compacts exhibited slightly increased drug release from the surface of its compacts over the amorphous form in the Sporanox formulation, however the excipients in this formulation may have affected these results. Hiew et al. investigated the impact of excipient shielding on release of acetylsalicylic acid (191). A number of compacts were produced, containing different ratios of drug and the excipient microcrystalline cellulose (MCC). Offline Raman spectroscopy was used to establish the distribution of MCC on the surface of each drug-excipient compact. Two different acetylsalicylic acid particle size fractions were investigated. This resulted in IDRs, which were broadly similar and could only be statically distinguished at lower acetylsalicylic acid to MCC ratios. A flow rate of 0.1 mL min\(^{-1}\) was deemed preferable as it achieved IDRs with the lowest standard deviation. This study showed that MCC at higher concentrations (75 % and over), resulted in initial suppression of acetylsalicylic acid dissolution and lower maximum IDR.
values (191). Madelung et al. utilised UV imaging to investigate the dissolution enhancement of the incorporation of sodium dodecyl sulfate (SDS) into griseofulvin compacts (192). This was achieved by production of a griseofulvin-SDS solid dispersion disc or the addition of SDS into the dissolution media. SEM-EDX prior to dissolution showed heterogeneity with respect to the distribution of griseofulvin and SDS. The initial high dissolution rate was attributed to the loose particles on the surface of the compacts as revealed by SEM analysis. Results showed that higher SDS concentrations either in the buffer or incorporated into the disc showed the largest improvement in griseofulvin dissolution rate (192). A key excipient used in immediate release formulations are superdisintegrants as they promote fast tablet disintegration. Zarmpi et al. utilised UV imaging to investigate the impact of two superdisintegrants (Figure 6 III and IV) on the dissolution of paracetamol and carbamazepine. The viscosity of sodium starch glycolate (SSG) was varied by applying three brands of the superdisintegrant (Explotab - low viscosity, Glycolys - low viscosity and Glycolys - high viscosity). Particle size distribution was also investigated using two brands of croscarmellose sodium (CCS), (AcDiSol - small particle size and Primellose - large particle size). All drug compacts containing superdisintegrants resulted in a higher dissolution rate when compared to the pure drug compact. This was attributed to the enhanced wetting of the compacts or compact disintegration. Pronounced dissolution rate enhancement was achieved by the low viscosity SSG and small particle size CCS and was more effective for the higher solubility drug paracetamol, as compared to carbamazepine. Paracetamol showed a faster dissolution in neutral solution than at acidic pH. This was largely attributed to increased excipient hydration. The opposite was true for the poorly soluble drug carbamazepine. This was attributed to the swelling of excipient on the surface of compact, which impeded drug release (189).

Table 2. Application of UV dissolution imaging in intrinsic dissolution rate (IDR) measurements and drug-excipient compatibility studies.
<table>
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<th>Comments</th>
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<th>Complementary assessment</th>
<th>Reference</th>
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*signifies not an imaging technique
4. Whole dose imaging

The introduction of the USP type IV-like whole dose cell in dissolution imaging (SDI2) facilitated studies on tablets and capsules not possible with the early smaller UV imaging instrumentation. A typical image generated by the UV imaging dissolution instrument for a whole dosage form is depicted in Figure 7. The dissolution of carbamazepine is apparent as the increase in absorbance at 280 nm. The carbamazepine dissolved from the tablet is carried with the flow exiting the flow cell at the top side. Quantitative information may be extracted from the images captured. Measurement of the drug concentration above the tablet is conducted using the so-called drug measurement zone, and subsequently used for calculation of the dissolution rate (Figure 7). The apparent carbamazepine concentration distribution in the cell reveals turbulent flow conditions.

Figure 7. Image from the analysis software of the UV dissolution imaging instrument (Pion SDI2). Image shows the dissolution of carbamazepine from a tablet in 1% SDS solution at a flow rate of 8.2 mL/min at 37°C. Detection at 280 nm. (Asare-Addo. Unpublished data).

The first mentioning of whole dose UV dissolution imaging showed the simultaneous release of metformin from (255 nm) and the swelling (520 nm) of a commercial Glucophage SR tablet (16). In addition to the larger imaging area allowing the assessment of larger dosage forms, the dual wavelength imaging capability of the recent UV/Vis imaging instrumentation importantly provided the possibility to separate absorbance due to released drug substance from the signal.
due to the formulation or excipients. This feature has also improved the possibility of relating drug dissolution to physical changes in the formulation or delivery system (16).

The oral drug delivery route still remains the route of choice over other delivery systems (195), (196). By formulating a drug to include a hydrophilic polymer matrix, modification of the kinetics of drug release from orally administered solid dosage forms can be pursued. Hydrophilic swellable polymers often used include HPMC, poly(ethylene oxide) and hydroxypropylcellulose (HPC) (27). Upon exposure to water or biological fluids, the hydrophilic polymer becomes hydrated, swells and forms a gel layer around an initially dry core (184), (195), (196), (197), (198). This gel layer hydration process is dynamic. The inward migration of fluid or ingress of fluid means that the gel layer grows over time. The thickness of the gel layer can delay the diffusion of an incorporated drug (195), (196). To this end, Ward et al. (106) developed a method to allow the measurement of the developed gel layer from hydrophilic matrices. Tablet measurement zones, absorbance thresholds and medium flow rates were investigated to determine the appropriate parameters required to facilitate repeatable measurements of swelling in hydrophilic matrices using the Vis imaging capability of the dissolution instrument. Using the CR and DC grade of hypromellose, the authors reported similar swelling for both grades despite the difference in morphology and differences in porosity as a result (106). The images produced, showed denser gels to form for the CR grade potentially as a result of the porosity differences, which may be missed by compendial methods. Li et al. (107) investigated the effect of Methocel K15M, K100M and microcrystalline cellulose (MCC) effects on theophylline release. Simultaneous imaging of drug release at 255 nm and swelling at 520 nm showed the distinctly different behaviour of the Methocel polymers relative to the MCC matrix demonstrating the potential of the imaging system to characterise excipients as well as potential drug-excipient interactions. The early onset of the gel layer in hydrophilic matrices is of importance especially for the controlled release of soluble drugs to ensure dose-dumping does not occur (199), (200).

The $Smr2$ parameter from the focus variation microscope, which reflects the ratio of the area of the material (compact) at the intersection line, which separates the dales from the core surface to the evaluation area was used as a proof of concept predictive tool in determining wetting and initial hydration from a natural (xanthan gum, XG), semi-synthetic (hypromellose, HPMC) and synthetic (polyethylene oxide, PEO) polymer using the whole dose cell. $Smr2$ can also be used to calculate the percentage of ‘valleys’ that will retain lubrication and was applied to determine the percentage of the compact surface available for wetting during the initial
hydration process. The porosity of the compacts produced were in the order XG > HPMC > PEO. From the porosity perspective, it was therefore expected for the initial wetting and gel layer to be in the self-same way. The Smr2 parameter, however, from the FVM analysis suggested the tablet surfaces able to retain more lubrication and therefore wet quicker to be in the order of XG > PEO > HPMC. The whole dose imaging of the wetting and early gel layer formation concurred with the Smr2 prediction confirming it was possible to predict initial wetting therefore suggesting differences in porosity was not a major indicator for water ingress (201) (Figure 8).

The rate of drug release from matrices can be influenced by formulation variables as well as by the composition of the surrounding medium it is exposed to after ingestion, such as pH, enzymes, electrolytes and surfactants (202), (203), (204). The liquisolid technique has been used as a means for either slowing down (high solubility APIs) or increasing (poorly soluble APIs) drug release (205). Propranolol hydrochloride release from liquisolid compacts of sesame gum polysaccharide was successfully imaged qualitatively and quantitatively using the UV dissolution imaging (104).

The pharmaceutical industry utilizes film coatings for various reasons which includes but not limited to aesthetics, trade marking, taste masking of bitter APIs, product stability, drug release manipulation through retardation and providing resistance to the harsh conditions experienced in the stomach(206). Gaunø, et al. utilised UV dissolution imaging to investigate the release of 5-aminosalicylic acid from single extrudates, with an ethyl cellulose layer coating into a 0.5 % (w/v) agarose hydrogel (207). The effect of varying coating thicknesses on drug dissolution was investigated. The thickness of ethyl cellulose layer coating was determined by percentage weight gain, four different variations were used (0.5 %, 1.0 %, 1.5 % and 2.0 % weight gain). Dissolution testing was also performed using a conventional paddle dissolution system. The same rank order of dissolution performance for different coated extrudes was reported in each dissolution method, with the highest dissolution rate reported in the thinnest coated extrude and the lowest from the thickest coat. Studies were conducted to establish the impact on damages to the extrudate coating, this showed a faster dissolution rate from the damaged extrudates thereby exhibiting the technique for potential trouble shooting. Alqahtani et al. (208) investigated the use of low quantities of functional additives to control drug release from hot-melt extruded solid dispersions of carbamazepine (extrudates). The UV imaging aspects corroborated with the conventional dissolution studies (basket method). The disintegration
process of the surface of the extrudates containing gelucire was also visualised which was missed by the conventional basket method.
Figure 8. Optical images obtained from the dissolution imaging system at the 520 nm wavelength: (a) XG, (b) PEO, and (c) HPMC CR. Note: Images are taken from the 2 h experiment at time intervals of 1, 2, 5, 20, 30, 60, 90 and 120 min. XG is xanthan gum, PEO is polyethylene oxide, and HPMC CR is hypromellose K100M CR grade. Reprinted from (201). MDPI (Open Access).

Zarmpi et al. used a QbD approach to investigate the effect of superdisintegrants, viscosity of binders, particle size and lubricants on the release of a soluble (paracetamol) and poorly-soluble (carbamazepine) model compound from compacts made using blends of API and varying concentrations of superdisintegrants (209). It was observed from the UV dissolution imaging process that for the highly soluble drug, a high viscosity binder and smaller particle size lubricant caused a quicker disintegration process whereas for the poorly soluble drug, a slower disintegration was observed with an increase in the viscosity of the binder as compared to the control compacts of pure API (Figure 9). The UV imaging thus provided a visualisation of the disintegration and dissolution process.

The whole dose dissolution imaging has been used in the evaluation of several salt forms of the carboxylic acid gemfibrozil Asare-Addo et al. (194). The formulated salts exhibited increased dissolution over the parent drug gemfibrozil and correlated with the rank order using the IDR compact cell as reported earlier. The whole dose dissolution imaging used in the assessment of the release of indomethacin from ASDs with Soluplus showed that the solid dispersions regardless of the ratio of Soluplus or preparation method had an increased dissolution rate as compared to its parent drug (indomethacin only). Here, a different outcome to that of the rank order produced by UV dissolution imaging compact cell was observed (105). UV and Vis images depicted the tip of a capsule opening for the liberation of the indomethacin drug. UV imaging has also been used to provide insights into the dissolution behaviour of 3D printed formulations. Arafat et al. investigated sodium warfarin release from 3D printed dosage forms at a wavelength of 255 nm (210). The visible wavelength of 520 nm was used to visualise the erosion of the methacrylate used in the 3D printing of the matrix. The ability of the dissolution method to simultaneously monitor the sodium warfarin release and the erosion of the methacrylate showed that the bulk of the drug release to occurred via a diffusion process prior to polymer erosion at a rate of 16.4 and 15.2 µm/min in the horizontal and vertical planes, respectively.
Figure 9. Real-time surface dissolution UV images for a. PRC and b. CBZ tablets in 0.1 N HCl pH 1. Reprinted from (209). Copyright (2020) with permission from Elsevier.

5. In vitro release testing - non-oral routes of drug delivery

Most studies involving UV imaging have concerned the assessment of dissolution behavior in the context of oral drug delivery, e.g., intrinsic or tablet dissolution. However, one of the very first UV imaging applications was the release of nicotine from transdermal patch samples (211). The ability to image concentration gradients and drug transport is of potential value for elucidating release processes relating to most routes of administration and formulation approaches. In general, UV imaging may be suitable for assessment of drug release when the API possesses a chromophore and the release medium or matrix is sufficiently transparent at the relevant wavelength(s). The physical form and geometry (size) of delivery vehicle, complexity of the formulation as well as the release matrix are additional parameters that should be considered in relation to image area and resolution prior to embarking on UV imaging-based release testing. In this section, the application of UV imaging for characterization of parenterals is covered; special applications directed at physicochemical characterization of the active substance by UV imaging are discussed in section 7.

5.1 Injectables

Similar to in-vitro dissolution testing for oral products, in vitro release or drug release testing is a key tool for characterization of injectable sustained release formulations. In vitro release
testing for injectable parenterals is of importance throughout formulation development and in quality control. The use of the compendial methods, developed mainly for oral drugs or transdermals, has been endorsed and utilized with variable degrees of success (212), (213). Relative to oral drug delivery this field is less well developed; pharmacopeial reference methods remain to be established for injectables (212), (214). A standardized in vitro release testing method for injectables is recognized as highly desirable but also a difficult to reach objective given the diversity and complexity of injectable formulations (213), (214). Typically, in vitro release tests for injectables are classified as sample and separate, dialysis membrane or continuous flow methods. A coverage of the general features of the three types of methods and the variability among the classes can be found in several reviews, e.g., (215), (216), (217), (218), (219).

Somewhat surprisingly, given the commercially promoted flow cells for dissolution testing, UV imaging continuous flow methods for injectables have not yet been reported. Membrane-based methods in connection with UV imaging also appears not to have been explored in relation to injectables, albeit to permeability studies (220). The use of UV imaging in relation to injectables has been connected to the use of gel matrixes as the receiver medium (221), (222), (223), (224), (102), (225). A distinct and useful feature of UV imaging for in vitro release testing is the possibility of performing the release studies in a non-intrusive manner and without the need for compound labeling, thus, perturbation of the system or release process may be avoided. On this basis, it may be argued that the UV imaging methods do not belong to the group of sample and separate methods either. However, the ability to separate the absorbance signal due to the released drug from the absorbance of the formulation and matrix is required in UV imaging-based release studies.

Various hydrogel constructs have attracted interest as soft tissue mimicks in relation to drug release testing (226), (227), (228), (229), (230), (231), (232), (233), (234), (235). In relation to UV imaging studies, primarily agarose gels have been applied due to their robustness and transparency (236). Ye et al. studied piroxicam release from medium chain triglyceride (MCT) into either 0.5 % (w/v) agarose or 25 % (w/v) Pluronic F127-based hydrogels serving as simple subcutaneous tissue models (221). Vegetable oils, like MCT, serve as delivery vehicle for depot injectables comprising, for instance, steroids or antipsychotics (219). Using a simple geometry limiting drug transport to one dimension, the diffusion and partitioning of piroxicam was characterized as a function of matrix (agarose and Pluronic F127) and pH of the aqueous gel phase utilizing expressions based on Fick’s second law (221), (237). Changing pH from 4.0 to 7.4 appeared to have little impact on the diffusivity of piroxicam in the gel phase, however, a
larger effect was observed on the distribution coefficient depending on the vehicles. This was attributed to medium polarity affecting the degree of piroxicam ionization. These proof-of-concept studies demonstrated the ability of UV imaging to detect and monitor small changes, however, the ability to monitor distribution of highly lipophilic studies remains to be demonstrated. The distribution of piroxicam upon injection of small volumes (3 μl) of aqueous and MCT solution into agarose gels was also assessed (237).

Sun et al. initially assessed the feasibility of UV imaging instrumentation to study the phase separation of in situ poly (lactide-co-glycolide acid) (PLGA) forming implants (225). The PLGA was dissolved in 1-methyl-2-pyrrolidinone (NMP) or triacetin (TA) without model drug. The solutions were subsequently pipetted into a 2 mm wide cavity in the centre of an agarose gel. In this small-scale format, the effect of PLGA concentration (20 % and 40 % (w/w)) and type of cosolvent on the rate of phase separation was investigated at 550 nm. Using changes in absorbance or light intensity, slower phase separation with PLGA-TA than PLGA-NMP solutions was observed in line with existing literature on slow and fast phase separating systems. Interestingly, SEM and light microscopy studies showed that the agarose concentration had a significant impact on the morphology of the formed implants (225).

Concomitant UV imaging, Vis imaging and light microscopy were subsequently applied to monitor the in-situ formation of PLGA-NMP and PLGA-TA implants containing piroxicam or α-lactalbumin upon exposure to an aqueous medium in the form of 1.0 % (w/v) agarose gels (238). The measurements were performed over 24 h using the so-called small-scale format, i.e., in 1 mm light path cuvettes holding the in-situ forming implants (20 μl preformulation), which were moved between the three measurement units. Implant formation was assessed using Vis imaging and light microscopy and initial drug release was quantified by UV imaging. α-Lactalbumin invoked an accelerated phase separation of the PLGA-solvent systems as compared to piroxicam, highlighting the interplay between (model) drug, polymer and cosolvent (238). Recently, the initial (early time frame of 3 h) release of leuprolide acetate from in-situ forming PLGA-NMP implants was investigated by UV-Vis imaging (102). Figure 10 shows the schematic UV-Vis imaging setup applied and selected UV and Vis images recorded at selected time points. In contrast to previous studies, a larger imaging area of 25 × 25 mm² (and dual-wavelength capability) facilitated the injection of a dose and volume (3.0 mg of leuprolide acetate and 100 μl, respectively) comparable to a clinical setting. The study addressed the significance of different injection conditions and matrices applied for emulating the subcutaneous space (102). Figure 10B shows the effect of applying hyaluronic acid solution
as the release matrix as compared to an agarose gel with or without an imbedded reservoir to define the geometry of the in situ forming implant. The increase in absorbance at 280 nm near the implants was due to the release and subsequent diffusion of the drug and represents a characteristic for UV imaging-based release studies. The visible wavelength (525 nm) was used for identifying the depot-matrix interface, which was subsequently applied to extract an estimate of implant swelling. The application of hyaluronic acid, a constituent of the extracellular matrix and a proposed subcutaneous tissue surrogate for in vitro release testing (239), led to lower leuprolide acetate release as compared to the release studies encompassing agarose gels. UV-Vis imaging was sufficiently sensitive to monitor the initial or burst release of leuprolide acetate from the long acting PLGA implant. Supplementary imaging measurements identified an interaction between the drug and the polyelectrolyte hyaluronic acid (102) (cf. section 6), in addition to identifying the importance of the surrounding matrix on the swelling behaviour of the implants. Implant swelling and drug release have previously been reported to be affected by the interstitial pressure and compressive forces of the surrounding matrix (235).

In relation to injectables, dissolution studies of various insulins from compacts into agarose gel provided the following rank order: human insulin > bovine insulin > protamine insulin at pH 7.4 (223). The dissolution of amorphous human insulin (pI ~5.4) was investigated at pH 3.0, 5.4 and 7.4. The dissolution rate was correlated to the apparent solubility of the peptide in the order pH 3.0 > 7.4 > 5.4. Surprisingly, the dissolution profiles exhibited different shapes in the stagnant system; a linear relationship between the amounts of human insulin dissolved and time was observed at pH 3.0 and 7.4, whereas at pH 5.4 the dissolution rate decreased with time consistent with $\sqrt{t}$ - kinetics. Based on “surface” insulin concentration measurements and in situ local pH measurements (the pH at the solid surface changed with time in the experiments performed at pH 3.0 ad 7.4) performed using the UV imaging instrumentation; the anomalous dissolution behaviour could be related to the ionization properties of insulin and the buffer capacity of the solution phase in the gel (223). Jensen et al. further compared the release of insulin from drug loaded Sterotex implants into agarose gel by UV imaging (224). The effect of drug load and implant porosity was assessed in a non-intrusive manner by imaging, however, the most striking result was the widely different release profiles obtained upon comparison to release profiles generated using another “gel sampling” method and an agitated phosphate buffer solution as the release media (224).
It should be noted that the prototypic release setups utilizing gels or stagnant solution phases discussed above work under non-sink conditions in contrast to what is normally recommended for dissolution and release testing. For the characterization of long acting injectables by UV-Vis imaging beyond the initial phase, incorporation of a convective contribution to drug transport is considered a necessity. To this end, Sun et al. applied a custom-made cell similar to the one depicted in Figure 10A as a flow cell (240). Piroxicam dissolved in 1-methyl-2-pyrrolidinone was injected into the flow cell being perfused with phosphate buffer at pH 7.4. UV imaging in combination with light microscopy and Raman spectroscopy was applied to monitor and document the initial local piroxicam supersaturation, precipitation occurring in the flow cell and subsequent drug dissolution. The powerful setup provided apparent drug concentrations and the parallel Raman measurements identified the solid form of the precipitated piroxicam (240). The utility of such a setup for in situ forming suspensions or other depot type injectables can easily be envisioned. However, the study also highlighted the sensitivity of such systems to small changes, e.g., with respect to hydrodynamic conditions. The simple alteration of the flow cell from a horizontal to an upright position fundamentally changed the outcome of the experiment due to the interplay between density and hydrodynamic conditions. This should be taken into consideration during the course of developing new in vitro release testing methods for injectables (and other formulation types). Even subtle changes may make comparison to work performed in other labs challenging. Similarly, the change of release medium and drug–matrix interactions discussed above may offer an opportunity to attain important new insights on formulations under development, but is likely to constitute a hindrance in a quality control setting (unless thoroughly characterized). The current UV-Vis imaging-based in vitro release testing methods for injectables, in their present form, cannot be expected to accurately model or simulate the entire in vivo release and absorption process upon, for instance, injection into subcutaneous or muscular tissue. The UV imaging setup rather lends itself to testing the importance individual physiological or physical chemical factors or processes. From a set of consecutive experiments involving UV imaging and other techniques, the entire release and absorption process may then be simulated. In its present form, UV imaging must be anticipated mainly to find use in early formulation screening rather than quality control and batch-to-batch release testing. Although the main area of use for UV imaging is aiding decision making in the early stages of formulation, due to its ability to produce a visualization of dissolution phenomena, it could have potential applications in quality control checks of finished formulations should they fail QC testing.
Figure 10. Schematic representation of the D200 UV-Vis imaging system for monitoring release of leuprolide acetate from in situ forming PLGA implants (A), representative absorbance maps obtained by UV imaging at 280 nm and vis imaging at 525 nm in 0.5% (w/v) agarose gel and 10 mg/mL HA solution upon injection of the PLGA/NMP pre-formulation containing 3% (w/w) LA (B), and photographs of the corresponding implants removed from the release medium after the experiments (C). Reprinted in modified form with permission from (102). Copyright 2020 American Chemical Society.
5.2 Miscellaneous formulations

As alluded to above, nicotine release from a transdermal patch was a first application of UV imaging (211). As is probably the case for most of the applications discussed in this review, the study had the character of a feasibility study aiming to characterise UV imaging as a tool for performing in vitro drug release studies. Nicorette patch samples were mounted in a flow cell, the cell was filled with phosphate buffer, which was periodically replenished. The work mainly focussed on the analytical aspects of performing UV imaging measurements, e.g., calibration curve and LOQ. However, the release of nicotine as a function of time was quantified and found to adhere to Higuchi kinetics ($\sqrt{t}$ – kinetics) (211). In relation to transdermal drug delivery, Fazili et al. investigated the permeation of ibuprofen over a silicone membrane in a 3D printed Franz type cell compatible with UV imaging (220). The experimental setup facilitated real-time measurement of ibuprofen permeated to the acceptor phase. In the context of wound healing, Preem et al. studied the release of chloramphenicol from electrospun nanofiber mats using UV imaging (241) (Figure 11). In line with previous studies, an agarose gel was applied as the release matrix assumed to reflect better the in vivo conditions as compared to a buffer solution. This setup allowed a qualitative comparison to antibacterial tests conducted on agar plates and a bioreporter disc diffusion assay. There was a considerable difference with respect to chloramphenicol release into phosphate buffer solution between mats made exclusively from polycaprolactone (PCL) and those comprising PCL and 2 \% (w/v) polyethylene oxide (PEO). The faster drug release from the latter was attributed to the hydrophilic nature of PEO. The same rank ordering, but with a much less pronounced difference, was observed upon chloramphenicol release into the hydrogels, indicating that the matrix constituted a diffusion barrier minimizing burst release effects (241). He et al. designed HPMC-based buccal films containing the basic drug saquinavir ($pK_a$ 7.1). With the aim of increasing the pH dependent solubility and release of saquinavir, the organic acids malic acid, citric acid or succinic acid were incorporated into the film matrix (242). In vitro release testing at pH 6.8 showed that addition of an organic acid increased the release of saquinavir from the buccal films. Vis imaging was applied to probe the change in microenvironmental pH during drug release. The pH indicator bromothymol blue ($pK_a$ 7.2) was dissolved in the phosphate buffer (pH 6.8) used for preparing the 0.5 \% (w/v) agarose gel used as the release matrix. Saquinavir release from the films was initiated by placing the films on the gel in the imaging system. The local pH, or microenvironmental pH, was assessed by imaging at 610 nm where bromothymol blue has a pH dependent absorbance (242). The apparent transient pH changes
were the result of drug and excipient transport processes and the buffer capacity of the system. Further studies are, however, needed to assess how relevant the pH maps or images created are relative to the *in vivo* situation. However, the setup provides a simple approach for estimating local pH changes and potentially provides an indication on the importance of altering formulation parameters, i.e. pH modification.

Figure 11. (A) Representative images of CAM release from fibers and diffusion into 0.5% (w/V) agarose hydrogel at 37 °C. Top is PCL/PEO/CAM, middle is PCL/CAM and bottom is PCL/Blank fibers (control). The zone used for the quantification of the CAM absorbance is shown in the middle row at 30 min. Representative concentration-distance profiles for PCL/PEO/CAM (B) and PCL/CAM (C) fibers diffusing into 0.5% (w/V) agarose hydrogel at 37 °C. The red, black and blue line is for 0 min (immediately after starting the experiment), 30
min and 180 min, respectively. Key: CAM, chloramphenicol; PCL, polycaprolactone; PEO, polyethylene oxide. Reprinted from (241). MDPI (Open Access).

6. Characterization of drug substances (and delivery systems)

The dissolution and release behavior of a drug substance is closely related to the physical chemical properties of the compound. Given the ability to monitor drug concentrations and concentration gradients, UV imaging may also be suited for characterization of physical chemical properties such as solubility, diffusion, partitioning, and binding or interaction phenomena in a spatially resolved manner.

According to the Noyes-Whitney model, the concentration of the dissolving substance at the solid surface is equal to the solubility. To this end, Lu and Li proposed a method for simultaneous measurement of drug solubility and diffusion coefficients based on fitting a numerical solution of Fick’s second law to experimental drug concentration distributions determined from disk dissolution experiments performed under static conditions by UV imaging (193). Satisfactory agreement to literature results was observed for the investigated model compounds carbamazepine and ibuprofen. UV imaging is also frequently denominated surface dissolution imaging. However, at least with the first generation instruments it has proved difficult to get good concentration measurements at the solid surface due to optical and instrument artefacts, cf. (141) and (99). Thus, the approach of Lu and Li (193) deserve merit.

Diffusion, the transport of matter due to Brownian motion of molecules and a chemical potential gradient (concentration gradient) is one of the most important process in drug research. UV imaging provides an interesting approach to the study of drug diffusion properties. Methods such as dynamic light scattering, nuclear magnetic resonance and Taylor dispersion analysis are well suited for simple solutions (236), (243) UV imaging, and other optical methods, are of particular interest in relation to more complex solutions and gels. Diffusion studies based on UV imaging have been performed in agarose gels (99), (221), (236), Pluronic F127 gel (222), composite gels of agarose and polyelectrolytes (237) and hyaluronic acid solution (102). UV imaging diffusion measurements were sufficiently sensitive to capture the concentration and pH dependent self-association behavior of insulin attributed to the peptide monomers, dimers and hexamers (236). A prerequisite for the use of UV imaging in diffusion studies is that natural convection due to density gradients can be adequately suppressed. Fortunately, the imaging approach itself constitutes a tool for detecting the
occurrence of such phenomena as reported previously (99), (126). An additional prerequisite is that the matrix is sufficiently transparent. The diffusion studies are relatively easy to conduct.

At University of Copenhagen, ~200 undergraduate students per year use the UV imaging setup for visualizing diffusion processes and determining diffusion coefficients during a 4 h lab exercise in the pharmacy program. Provided two different phases, i.e. two different gels or an oily and a gel phase are brought in contact with each other in the diffusion cell, UV imaging can also be used to visualize and assess partitioning effects. The distribution coefficients for the partitioning of piroxicam between MCT and aqueous gel phases were determined 11, 28.

Along similar lines, Ye et al. investigated the role of electrostatic interactions as related to analyte diffusion in pure 0.5 % (w/v) agarose gel, 0.5 % (w/v) agarose gel containing 0.5% (w/v) chondroitin sulfate (CS), and 0.5% (w/v) agarose containing 0.5% (w/v) hyaluronic acid (HA) (244). H-Ala-β-naphthylamide, H-Lys-Lys- β-naphthylamide, lysozyme and α-lactalbumin were selected as the model compounds. The incorporation of the negatively charged polyelectrolytes chondroitin sulfate and hyaluronic acid into the gel phase greatly reduced the apparent diffusion coefficients of the net positively charged molecules, whereas the diffusivity of α-lactalbumin was almost unaffected as compared to the diffusivities obtained in the neat 0.5 % agarose gel. The diffusivity of the cationic compounds was highly dependent on the ionic strength, with higher ionic strengths leading to a higher apparent diffusivity, indicating the effect of electrostatic interactions on apparent diffusivity (244). A similar setup, identified the interaction of leuprolide acetate with hyaluronic acid (102) (Table 3). Such studies may be useful to assess the importance of drug physicochemical properties in relation to various drug release applications. UV imaging-based diffusion studies in gels are relatively well established, whereas further experiments are required to explore the wider applicability for assessing partition coefficients and interactions.

Table 3. Overview of UV imaging-based studies related to non-oral routes of administration and physical chemical characterization
UV dissolution imaging is interesting in formulation development due to its ability to monitor and relate physical changes to drug release rates, as discussed above (section 4). The IDR and

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<td>Piroxicam</td>
<td>Diffusion of drug from oil into hydrogel</td>
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<tr>
<td>Piroxicam</td>
<td>Influence of pH on drug diffusion and partitioning</td>
<td>Agarose and Pluronic F127 hydrogel</td>
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<tr>
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<td>IDR imaging from in situ ophthalmic gels</td>
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<td>Insulin</td>
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<td>Insulin</td>
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7. Instrument performance
whole dose set-up of the UV dissolution process presents an interesting bridge between
preformulation and compatibility studies and dosage form evaluation of used later in
formulation development. Visualization is one of the key benefits of the UV imaging system
with the measurement of drug concentrations in the immediate vicinity of the compacts or
delivery system providing additional insights to the drug dissolution process (247). UV
dissolution imaging is complementary to many other imaging techniques, such as CARS, MRI,
NMR imaging, XµT, FTIR imaging, NIR imaging, fluorescence imaging, and THz imaging,
which provide information mainly related to the solid phase or the vehicle. A limitation of the
UV/Vis imaging technique is the lack in chemical selectivity as compared to ATR-FTIR,
Raman mapping and CARS and limited ability to monitor internal changes in internal structure
such as the MRI and XµT.

Insights into the critical steps related to various applications start to emerge. A key issue is the
establishment of best practices and protocols particularly in relation to data collection and data
analysis to reduce sources of error. Approaches for characterizing instrument performance, for
instance in relation to sensitivity, resolution, dynamic range and repeatability, are still under
development (99), (130), (141), (102) (211). The resolution is depending on the
instrumentation used and the samples applied, but certainly larger than the pixel size ~30 µm
for the Actipix D200 system (102) effectively limiting work with nano- and microparticles.
The linear range observed is wavelength and matrix specific, and according to the experience
of the authors in general less than for conventional UV/Vis spectrophotometers. Calibration
curves should be prepared on the instrument used for the imaging measurements (99). Having
only one or a few wavelengths available, care should be exercised when attribution the
measured absorbance to dissolved API or particulate or solid material scattering of blocking
transmission of light. UV imaging has the unique advantage to offer qualitative data in the form
of images of dissolution events and, thus, may still prove useful if accurate measurement of
drug concentrations is not possible.

The relative simplicity of UV/Vis imaging technique in relation to its operation and data
interpretation is likely to facilitate its uptake and use in formulation development labs. As USP
dissolution methods are well established in QC and batch releases, it is unlikely that UV
imaging will be used as a tool in these settings in the near future. The one sample at a time and
lack of procedures for method suitability and instrument qualification are main obstacles.
8. Conclusion

The first applications of UV imaging in relation to dissolution and release testing appeared in 2010 (211), (96). Since, a number of studies have appeared, mainly in the area of intrinsic dissolution, form selection and drug-excipient interaction. Other studies have addressed drug release in relation to non-oral routes of administration or physical chemical characterization. Recently, with the advent of larger area imaging systems, whole dose studies related to tablets, capsules as well as injectables have started to occur. The key feature of UV imaging is related to the fact that most drug substances absorb light in the UV range, allowing drug concentrations and concentration gradients to be visualized and quantified at the surface of solid materials during dissolution or near the delivery vehicle during drug release. This opens for detailed insights into dissolution and release processes difficult to extract using conventional dissolution methods.

UV dissolution imaging ties in with at least two of the current trends in pharmaceutical sciences. One is miniaturization, to accommodate efficient characterization of new active molecules with the limited amounts of material available in the early stages of drug development. The miniaturisation comes in parallel with a high degree of analytical automation or a real-time analytical methods. Here the non-intrusive monitoring of the dissolution process makes UV imaging an attractive option for both the preformulation and formulation.

Another trend is the increasing application of high information content imaging modalities. The use of UV imaging to study pharmaceutical systems in-vitro is still at an early stage when compared to other imaging techniques such as CARS, MRI, NMR imaging, XµT, FTIR imaging, NIR imaging, fluorescence imaging, THz imaging and CLSM. The relative ease of operation and simple data analysis according to the Lambert-Beer law makes UV dissolution imaging an easily accessible imaging approach in formulation development. UV imaging is complementing most imaging techniques by monitoring mainly the solution phase. Some structural information on delivery vehicles may be obtained but in this sense UV imaging is generally much less selective. The micro-environment and structure of dosage forms characterized with respect to monitoring local pH changes, polymer swelling, peptide self-association and electrostatic interactions have been reported.

UV imaging is still an emerging technology but appears promising for drug dissolution research. Many published studies have the character of method feasibility studies. This is likely to change in coming years. To this end, studies such as the danirixin (148) case is particularly
encouraging bridging physical chemical characterization using UV imaging with clinical studies. Additional promising applications is the detailed elucidation of solid-state transformations possible by combining UV imaging with simultaneous in-situ Raman spectroscopy and the emerging whole dose applications.

9. Challenges and outlook

A range of small-scale dissolution methods for IDR determinations are available, most of which are capable of running multiple samples in parallel. In comparison to these, UV dissolution imaging is likely to a have lower throughput, but may also be associated with higher variability on the IDR values because quantification is relying on the suitability of the hydrodynamic model of the prevailing flow conditions and the image analysis. To what extent this limitation is counter balanced by the additional information generated by the images may depend on the specific application. Similar considerations may apply to the whole dose applications.

More work is needed for establishing method qualification and validation procedures for UV imaging. Media exhibiting high UV absorbance can also be problematic because of the limited linear range of the detector. Other challenges encountered in practical UV imaging are storage and analysis of large amounts of data captured. This is particularly challenging for sustained release formulations where prolonged monitoring is desirable. An unknown also, in relation to this area is the advances in UV imaging sensors.

The use of compendial dissolution testing instrumentation is generally endorsed and the proliferation of in-vitro release testing methods considered undesirable. However, the continued interest in developing new release testing methods most likely reflects an unmet need. With the current commercially available instrumentation, UV-Vis imaging is unlikely to replace the pharmacopeial methods as relates to QC, batch-to-batch, and biowaivers. The lack of multiplexing, or only one sample at a time, is a limitation, as is the lack of established instrument qualification procedures. UV imaging is more likely to find use in the area of formulation development shedding light on drug dissolution and release processes and mechanisms, i.e. method proliferation is likely to continue. Here the ability to rapidly design and make customized cells and holders compatible with the UV imaging instrumentation, e.g., by additive manufacturing (220), (240) can be effectively exploited. It is envisioned that dedicated cells can be used to create systems emulating to a larger extent human physiology at
the site of administration, which will be helpful in elucidating the rate limiting parameters controlling drug release, or to shed light on specific transport mechanisms.

UV/Vis imaging is a very versatile tool in drug research, the simplicity in terms of operation and (initial) data interpretation will favor continued use.

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Conflict of interest

NONE

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