A Modern Approach to the Heckel Equation: The Effect of Compaction Pressure on the Yield Pressure of Ibuprofen and its Sodium Salt

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Abstract

Despite being heavily criticised in the literature the most widely used and accepted compaction equation is the Heckel equation proposed by Heckel in 1961. Its presence in literature for 55 years is due to the ease in which it can distinguish between plastic and brittle materials. Achieving the correct balance of plastic and brittle materials in a formulation is critical to ensure adequate tablet strength is achieved therefore classifying materials using the Heckel equation is attractive. Despite the importance of this understanding, especially in the design and manufacture of direct compression formulations, there are no set analytical testing standards or materials classification guidelines. Instead many workers have attempted to develop techniques for the measurement and classification of a materials deformation but there is still confusion and contradiction present in this field.

In this study we report the effect of compaction pressure on the yield pressure of ibuprofen and its sodium salt. Ibuprofen and its sodium salt were selected as models for study due to the availability of comparative literature yield pressure values. The reported variation in yield pressure can be significant e.g, ibuprofen which has literature values of 21-1139 MPa. This study proposes an approach to determine yield pressure from the Heckel equation using a linear regression method. The full linear regression methodology utilised is described and is used to report the yield pressure of ibuprofen and its sodium salt dihydrate. This technique reveals the most representative compaction pressure in order to derive yield pressure to be 121 MPa.

The yield pressure of ibuprofen and its sodium salt have been shown to increase with increasing compaction pressure. The reported values lie between 52-78 MPa for ibuprofen and 48-75 MPa for ibuprofen sodium dihydrate. The slightly lower reported yield pressure values for ibuprofen sodium suggest increased plasticity which could be attributed to the water contained within the structure acting as a plasticiser.

Keywords: Heckel equation; Deformation; Yield pressure; Linear regression; True density

Introduction

As tablets are the most common dosage platform understanding the deformation behaviour of the individual components is critical. Despite the importance of this understanding, especially in the design and manufacture of direct compression formulations, there are no set analytical testing standards or materials classification guidelines. Instead many workers have attempted to develop techniques for the measurement and classification of a materials deformation but there is still confusion and contradiction present in this field.

Recently many efforts have focused on the study of a materials deformation at single crystal level [1-4]. Techniques such as nanoindentation provide information using minimal quantities of material however the relationship between single crystal and bulk mechanical behaviour is not well understood. This had led to the continued use of compaction equations to classify materials in terms of their deformation.

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Despite being heavily criticised in the literature the most widely used and accepted compaction equation is the Heckel equation proposed by Heckel in 1961. Its presence in literature for 55 years is due to the ease in which it can distinguish between plastic and brittle materials. Achieving the correct balance of plastic and brittle materials in a formulation is critical to ensure adequate tablet strength is achieved therefore classifying materials using the Heckel equation is attractive.

The model proposed by Heckel [5] is represented by the following equation:

$$\ln(1/1-D) = K \cdot P + A$$

(1)

where D is the relative density of the compact, P is the applied pressure, A is a constant suggested to represent particle rearrangement and the reciprocal of K is used to calculate apparent mean yield pressure (P_y). The density pressure relationship is based on first order kinetics and has been defined as the stress at which plastic deformation of a particle is initiated, where low P_y values represent plastically deforming materials and high P_y values represent brittle materials [6]. In the pharmaceutical industry, the deformation of excipients has been most widely characterised [7] and it is accepted that microcrystalline cellulose (MCC) and dibasic calcium phosphate (DCP) form the extremes of the deformation spectrum with MCC (Avicel PH-101) exhibiting a yield pressure of 50 MPa and DCP exhibiting a yield pressure value of 957 MPa [8].

The yield stress values of well characterised materials become important when attempting to classify new materials. Often a comparison to materials whose deformation behaviour is well understood will be made due to the absence of defined standard class limits on yield pressure in terms of a plastic or brittle material. Roberts and Rowe attempted to set limits on these values along with other mechanical descriptors including young’s modulus, hardness and strain rate sensitivity which allowed materials to be categorised [8]. Materials with P_y<80 were classified as soft/plastic materials whereas P_y>80 indicated hard/brittle substances. Although these limits are available they are difficult to adhere to due to variation present in P_y values for the same material between research groups. The variation present is due to many experimental and physical factors (experiment type, compaction speed, compaction pressure and particle size) affecting derived values and no standardisation being set on these.

The Heckel equation can be fitted to data collected from ‘in-die’ or ‘out-of-die’ compaction experiments. These different experiment types have been shown to affect yield pressure measurements due to the contribution of elastic energy [9,10]. Ilić et al., measured the yield pressure of MCC (Avicel PH 101) in-die and out-of-die and the respective yield pressure values were 61.7 MPa and 98.3 MPa. This difference of 36.6 MPa between experiment types is one of many factors contributing to discrepancies in yield pressure data published from different research groups.

As well as experiment type, it is also well documented that experimental conditions can affect yield pressure. It has been shown by numerous workers that compaction speed and final compaction pressure can affect yield pressure [11-13]. Roberts and Rowe proposed a term, strain rate sensitivity [14], which describes the relationship between yield pressure values collected at two different compaction speeds. Since then it has been shown that plastic materials are more sensitive to strain rate compared to brittle materials [15]. This means that when reporting the yield pressure of plastic materials, the compaction speed will alter the yield pressure value.

The variation seen in yield pressure is not only due to experimental conditions but can be attributed to the physical effect of particle size. A materials yield pressure is an intrinsic property and therefore should not be affected by physical properties such as particle size however there is contradicting information in the literature. Fell and Newton [16] described early on that the yield pressure of lactose was influenced by particle size. However, in 1982 a study by McKenna and McCafferty [17] revealed that during their calculation of yield pressure, particle size was irrelevant for lactose. Another study by Patel et al. [18] found that the yield pressure of paracetamol was dependant on particle size, where increasing particle size resulted in higher yield pressure values. Recent studies have suggested that as the yield pressure is derived from the density pressure relationship this value is actually an indication of powder densification rate that may be effected by particle size [19]. Nevertheless, yield pressure calculated from the Heckel equation is still derived to provide information on bulk deformation behaviour and particle size is one of the many variables affecting the outcome.

One area which may account for variability in the yield pressure value, but has been neglected in the literature, is the way in which K from the Heckel equation is derived. This is due to the fact that although it is accepted that yield pressure is derived from the gradient of a linear slope, Heckel plots are not completely linear and contain deviations at low and high pressures. These deviations are due to particle rearrangements at low pressures [6], and elastic deformation as pressure increases [9]. It is the responsibility of the analyst to select the most appropriate region in which to calculate yield pressure. This method is susceptible to user subjectiveness and has the potential to cause error in derived values which could lead to mistakes in material classification. Despite this, there are no standards or recommendations in the literature to define what area should be used in order to calculate yield pressure.

The type of variation in yield pressure that can be seen for a material, due to the factors discussed, can be significant e.g. ibuprofen which has reported values of 21-1139 MPa. The compressibility of ibuprofen has been studied by many workers [20-23] and a table listing the reported yield pressure values is shown in Table 1. The small variations present in studies, 4 MPa for Di Martino et al. [21] and 3 MPa for Liu et al. [23], are due to particle habit and particle size differences, respectively. The study by Patel et al. [22] reveals large differences accounted for by the effect of compaction pressure. This would suggest, according to the limits set by Rowe and Roberts, [8], that ibuprofen is changing from a plastic to brittle material. In an attempt to further understand these variations within the literature this study will focus on the effect of compaction pressure on ibuprofen yield pressure and the method to determine the linear region.

### Materials and Methods

#### Materials

Ibuprofen (40 µm grade) was supplied by Pfizer Ltd. Ibuprofen sodium dihydrate was purchased from Sigma Aldrich, Dorest, UK.

#### Methods

**Scanning electron microscopy:** Scanning Electron Microscopy (SEM) was used to qualitatively assess particle size and shape. Electron micrographs were captured using a Zeiss SUPRA 40VP (Carl Zeiss Microscopy GmbH, Cambridge, UK). The samples were mounted onto an aluminum pin stub containing sticky carbon tabs and sputtered...
coated with platinum. A voltage of 3.0 kV and working distance of 10 mm were used.

**Laser diffraction**: Laser diffraction was utilised to measure the particle size of the ibuprofen samples. Dry dispersion measurements using 200 mg (±5 mg) of sample were made at a pressure of 1 bar using a Sympatec HELOS laser diffraction system with an R5 lens in conjunction with a RodosM disperser and an Aspiros feeder controlled by WINDOX software (Sympatec GmbH, Germany). Samples were measured in triplicate.

**True density**: The true density of 2.5 g (±0.2 g) samples were measured in duplicate using a Pentapyc 5200e helium pycnometer (Quantachrome UK Ltd., Hook, UK).

**Compression studies**: A Gamlen GTP-1 single punch bench top tablet press which has a uni-axial saw tooth displacement profile (Gamlen Tableting, United Kingdom) was used to complete the compression experiments. The deformation of the punches was corrected by compacting an un-deformable metal disk.

Tablets of 100 mg (±5 mg) were produced using a 6 mm flat-faced punch and die. Compression events were made at a speed of 0.033 mm second⁻¹, following the same methodology described by Roberts and Rowe [11].

In order to investigate the effect of compression pressure on yield pressure ibuprofen and ibuprofen sodium were compacted to five different compaction pressures (52, 87, 121, 156 and 173 MPa). Six individual measurements were made at each compaction pressure for both materials, to enable the variation in the yield pressure for each material to be determined.

**Results and Discussion**

**Scanning electron microscopy**

Scanning electron microscopy was conducted to visually assess and qualitatively determine particle size and shape information. The electron micrographs of ibuprofen and ibuprofen sodium, shown in Figure 1a and 1b respectively, reveal a difference in size and shape between the two materials. The majority of ibuprofen particles are smaller than 100 µm and have a size of around 40 µm whereas the ibuprofen sodium batch contains both small (around 40 µm) and larger particles (around 200 µm). Differences in particle shape are also revealed, with ibuprofen consisting of smooth lath shaped particles compared to ibuprofen sodium which has both rough prismatic and lath shaped particles.

**Particle size distribution**

Ibuprofen and ibuprofen sodium were characterised using laser diffraction to enable a quantitative measure of particle size. The volume based particle size distribution for both materials are shown in Figure 2. The distributions reveal that ibuprofen contains a primary mode centring around 40 µm, which is in agreement with the visual observation. The distribution for ibuprofen sodium reveals that the main mode centres around 200 µm and has a wider distribution compared to ibuprofen. The cumulative underscore values are presented in Table 2 and shows that 50% of particles are <39.3 ± 0.2 µm for ibuprofen whereas this value is almost doubled at 71.7 ± 2.4 µm for ibuprofen sodium.

**True density**

Helium pycnometry was used to measure the true density of the materials. The true density values of ibuprofen and ibuprofen sodium were 1.12 g/cm³ and 1.17 g/cm³, respectively. The standard error of measurement for each material was calculated and determined to be 0.0013 g/cm³ (0.11%) for ibuprofen and 0.007 g/cm³ (0.61%) for ibuprofen sodium.

It has previously been reported that Heckel plots are sensitive to variations in measured density, 1% error in density can lead to 10% error in yield pressure estimate [19], therefore it is important to report density values used and include the standard error in the measurement for transparency. Since the error values are low it is predicted that this will have minimal effect on derived values however the impact or error in density will be investigated during yield pressure calculations.

**Yield pressure determination**

**In Die Heckel calculation**: In-die Heckel plots were obtained by calculating the relative density (D) in-die using the true density and compact thickness (derived from corrected punch displacement). The ln(1/1-D) was determined for each measurement at each compaction pressure and these were plotted against each other to enable yield pressure to be determined.
Linear regression fitting: To enable accurate determination and eliminate subjective selection of the linear region of the plot, linear regression analysis was performed. A qualitative assessment to select the most central part of the linear region was made. The regression coefficient for this selected region was calculated. The linear region was then extended in both directions and on each occasion the regression coefficient recalculated. This was achieved by extending the linear range, one x coordinate at a time, in approximately 10 MPa steps. Once the region which contained the highest regression coefficient (specific to x coordinate) was established the step size was decreased until an accuracy of 1 MPa was reached.

The effect of the regression coefficient on the gradient of the slope and hence the yield pressure is highlighted using an example of ibuprofen compacted to 121 MPa (repeat 1) in Figure 3. This reveals the importance of selecting the most representative linear region as selecting only a small portion of the plot, where r² is 0.984, gives rise to a yield pressure value of 85 MPa compared to a larger portion of the plot (r² is 0.999) which gives rise to a yield pressure value of 76 MPa, a difference of around 10%. According to Rowe and Roberts, [8] this would classify ibuprofen in two different categories, changing from hard/brittle to soft/plastic.

Effect of compaction pressure on derived yield pressure: Heckel transformations for ibuprofen at the different compaction pressures analysed are shown in Figure 4. The plots produced at each compaction pressure overlay suggesting that in principal the same linear region for each pressure could be selected and therefore deriving the same yield pressure results. However, the plot also reveals that the full linear range of the Heckel plot may only become apparent when higher final pressures are used.

Linear regression was performed on six Heckel plots at each compaction pressure for ibuprofen and ibuprofen sodium. During analysis it became apparent that differences in yield pressures arise due to the range of the linear region available for selection at different compaction pressures. Figure 5 shows an example where ibuprofen compacted to 52 MPa returns a yield pressure value of 48 MPa compared to 76 MPa for ibuprofen compacted to 121 MPa. The visually subtle difference in gradient due to the compaction pressure used causes a significantly large difference in yield pressure, 28 MPa. Compacting ibuprofen to higher compaction pressures extends the rate of powder densification which in turn lowers the gradient of the slope and explains the increase in yield pressure values.
Investigating the effect of error in true density: As discussed previously, it is known that error in measured true density can affect derived yield pressure values. To investigate this further a worst case scenario of ibuprofen sodium compacted to 52 MPa was chosen for investigation, due to ibuprofen sodium containing 0.5% higher error in density compared to ibuprofen. This measurement gave a yield pressure of 50 MPa and when the error in density was included returned values of 46 and 53 MPa which is a difference of around 14%. This large error associated with density is unavoidable due to the logarithmic transformation of the Heckel equation; therefore alternative methods to reduce error are of upmost importance.

Comparison of Ibuprofen and Ibuprofen Sodium: The effect of compaction pressure on yield pressure for ibuprofen and ibuprofen sodium is displayed in Figure 6. The derived value for ibuprofen ranges from 52-78 MPa and ranges from 48-75 MPa for ibuprofen sodium. At the very high compaction pressures (>156 MPa) it is observed that the yield pressure values for both materials reaches a plateau where the standard deviations overlap. This would suggest that at these compaction pressures the highest possible rate of powder densification has been reached however examination of Figure 3 reveals that in order to utilise the greatest range of linearity, final compaction pressures of 121 MPa should be used to report yield pressure values.

The comparison of ibuprofen yield stress to its sodium salt dihydrate reveals that at all compaction pressures ibuprofen sodium exhibits lower yield stress values. It would be expected from the results of previous studies that ibuprofen sodium would exhibit a smaller particle size distribution which would account for the difference [18]. However, examination of the particle size data (Figure 2) reveals that the salt contains a larger particle size. Therefore regardless of particle size, ibuprofen sodium dihydrate possesses an increased powder densification rate suggesting higher plasticity compared to the free acid. The water contained within the crystal structure of the sodium salt could be acting as a plasticiser allowing the easier slip to take place. This could explain the lower yield stress and therefore plasticity of ibuprofen sodium dihydrate compared to ibuprofen free base.

Conclusion

A new approach utilising linear regression has been developed to analyse Heckel plots. Examination of regression coefficients of the linear region of the plot has allowed the selection of the most representative area in which yield stress can be derived. The variation due to the linear region selection has been reviewed and revealed that variations of 10% are possible dependant on which part of the plot is selected.

This linear regression method has been effectively applied to ibuprofen and ibuprofen sodium dihydrate and highlights that increasing compaction pressure results in the calculation of higher yield pressure values. This phenomena reaches its peak at very high compaction pressures (>156 MPa) however the greatest range of linearity is observed when pressures of 121 MPa were used in this study. Therefore it is proposed to enable valuable comparison of yield pressure, high compaction pressures (around 121 MPa) should be used for analysis. We hope to extend this strategy to other pharmaceutical materials.

It is also proposed from this research that when reporting yield pressure values from the Heckel equation that all experimental conditions and linear regression methods should be clear and transparent in order to reliably classify materials.
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References


