1967

The Interrelationship of Bulk Density, Granule Density, Tablet Weight, and Weight Variation

Amritkumar Bhandari
University of Rhode Island

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THE INTERRELATIONSHIP OF BULK DENSITY,  
GRANULE DENSITY, TABLET WEIGHT,  
AND WEIGHT VARIATION  
BY  
AMBITKUMAR BHANDARI  

A THESIS SUBMITTED IN PARTIAL FULFILLMENT OF THE  
REQUIREMENTS FOR THE DEGREE OF  
MASTER OF SCIENCE  
IN  
PHARMACY  

UNIVERSITY OF RHODE ISLAND  
1967
ABSTRACT

The effects of granule density and bulk density on tablet weight, weight variation, and tablet density were determined for tablets compressed on a Colton Model 216, rotary tablet press. Hardness, thickness, and disintegration times were also determined for the tablets produced. Standardized granulations were made from lactose and mixtures of lactose and bismuth subcarbonate, with gelatin as a binder. Granule density and bulk density of the granulations were varied by changing the concentrations of bismuth subcarbonate in the formula. All tablets were made at a fixed rate of tableting using standardized settings of the fill, pressure, and overload adjustments. The interrelationship between granule density and bulk density was found to be almost linear, as was also the relationship between granule density, bulk density, and tablet weight.
MASTER OF SCIENCE THESIS
OF
AMRITKUMAR BHANDARI

Approved:

Thesis Committee:
Chairman
Dean of the Graduate School

UNIVERSITY OF RHODE ISLAND
1967
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INTRODUCTION

The compressed tablet has become one of the most widely accepted dosage forms for the administration of orally effective therapeutic agents. It provides a convenient and, if well made, an efficient form of solid dosage for oral administration. The advantages of this type of pharmaceutical preparation are well known and include accuracy of dose, economy, stability, portability, concentration, elegance, and convenience in dispensing and shipping.

The invention of compressed tablets is usually attributed to Brackedon, who in 1843 obtained an English patent on a simple device for compressing dry powders. By 1874 tablets for almost every known disease were being sold on European and American markets. At one time the preparation of tablets was based more on empirical considerations than on sound scientific principles. In recent years, however, there has been a trend to move from the 'art' in making tablets toward the science of tableting.

Tablets first became official in the 9th revision of the United States Pharmacopeia, 1926, and have been official in each succeeding revision.

The National Formulary has included an increasing
number of tablets since their introduction into NF V in 1926.

Both the United States Pharmacopeia and National Formulary set certain standards for the tablets listed therein, including:

1. Identity and purity of active ingredients
2. Quantitative limits for the active ingredients
3. Limits on disintegration time
4. Limits on weight variation.

Of these four standards for good tablets, the first two are mainly applicable to the purity and physical-chemical standards of the ingredients to be used for the required therapeutic action from the tablets. The standards for disintegration time limits and those for the variation in the weight of tablets are of great importance in the formulation and compression of granules in the preparation of tablets as dosage forms.

Most materials can not be made into tablets directly, but must be granulated; that is, they must be made into granules before they can be successfully compressed. The characteristics of the granulation such as granule size, bulk density, granule density, and porosity all have a direct effect on the tablets produced.

The qualities of the finished tablets are influenced by the physical and chemical properties of the granulations,
which, in turn, depend upon the physical properties of the ingredients used to make them. These properties, e.g., density, will most likely not be the same for each ingredient being used; and the density of granules prepared from such a variety of ingredients will not be the same for different types of tablet formulations. Therefore, overall granule density of different tablet formulations will change from one to another, and these variations should have some measurable effect on the weight of tablets produced. In addition, the bulk density is related to granule density; therefore, if granule density will vary from one granulation to another, the bulk density will also vary.

The character or qualities of granules are influenced not only by the materials going into the granulation but also by the technique of manufacture of the granulation. A granulation made largely from bismuth subcarbonate (density 6.86 gm/cc) is likely to have a higher bulk density and higher granule density than one containing largely calcium carbonate (density 2.92 gm/cc). In addition, a granulation with a wide range of granule sizes and a large proportion of "fines" is likely to have a higher bulk density than a granulation with a narrow granule size distribution.

Since, during the compression of a particular granulation, the weight of a tablet is adjusted by varying the volume of the die cavity, the packing characteristics of a
granulation are important. If the die cavity in the operating tablet machine is filled by predominantly large granules, there will be large void spaces. But if the same die cavity is filled by smaller granules, the space will be more compactly filled and there will be less void space. The tablets compressed from the two granulations will not have the same weight. Thus variations in the granule size ultimately may cause variation in the weight of tablets.

The effects on tablet weight of the variables of granule size, granule fluidity, punch and die size, tablet base formula, and tableting speed have all been studied and reported in the literature. However, there appears to be no report in the literature concerning the effects of the variables of bulk density and granule density on tablet weight. Since the weight or dose of medication in a tablet ultimately depends on the volume of granulation entering a die cavity, it is obvious that the density of the volume of granulations measured out by the machine will influence the final weight of the tablets produced.

This study is, therefore, primarily concerned with the relationship of the quality of density, including bulk density and granule density, to tablet weight. Since bulk density and granule density are interrelated, and since tablet weight variation is an important adjunct to tablet weight, these relationships are also considered.
Variation in the weight of compressed tablets was reported in the pharmaceutical literature as early as 1926; in this study Liverseege (2), an Analyst of Birmingham, England, examined forty-two samples of 5 grain tablets of four different kinds, including sodium salicylate, aspirin, calcium lactate, and sodium citrate. He analyzed more than 1700 individual tablets to determine the uniformity in weight of the tablets, uniformity in the amount of medicinally active substance, and uniformity in disintegration time of tablets. He found that the weight of 91.8 per cent of the tablets did not vary more than 5 per cent from the mean weight of the tablets of the sample; 7.3 per cent of the tablets ranged from 5 to 10 per cent from the mean; and 0.9 per cent showed a deviation of greater than 10 per cent from the mean weight of tablets. Based on his observations of the average weight of tablets, he suggested that, "It is necessary that the mixture (i.e. granules mixed with talcum, starch, etc.) should be in a uniform state of division, or equal volumes will not correspond to the equal weights, and therefore some tablets will weigh more than others." In this report he did not pinpoint any other factors causing weight variation, but stated only that the
granulation should be of uniform size. It has been only since 1948 that further detailed investigations of tablet uniformity have been reported. Arambulo and Deardorff (4 and 5) studied weight variation in tablets, using sodium chloride as a basic granulation. In this study, sodium chloride granules were separated into narrow size ranges from 4380 to 81 microns and compressed into tablets on a single punch tablet press. The press adjustments were set and remained constant during the compression of all the tablets.

These workers found that, as the particle size decreased, the average tablet weight at first increased, reaching a maximum at 150 - 350 microns, and then decreased. They also found that, as the particle size decreased, the percentage of physically perfect tablets increased, the tablets became more glossy, and in general more satisfactory. This trend also passed through a maximum, followed by production of tablets decreasing in quality.

Akoi and Fukuda (9), two Japanese workers, have studied several factors causing variation in the weight of tablets. In one paper they reported a study of the relationships between both granule size and tablet diameter and tablet weight variation. Their conclusion on this aspect of the investigation was, "In tableting of tablets of a definite size, the larger the diameter of granules, the larger became
the weight variation, the ratio of its increase being linear
and in tableting of tablets of different sizes with the
granules of the same size, the greater the weight of tab-
lets smaller became the weight variation, the ratio of
decrease being linear."

In order to study the relationship between the fluid-
ity of granules and weight variation of tablets during tab-
leting (8;11) these same workers coated the granules with
Vaseline, thereby producing granules of different fluidity.
They found that the greater the coefficient of friction of
the granules, the greater becomes the weight variation. The
coefficient of static friction was found to increase with
increasing amount of powder (including lubricant) in the
granules. From their results they concluded that it seems
advantageous to have the granules as small as possible.
They also studied the influence of a number of different
lubricants, including potato starch, Carbowax 400, stearic
acid, Boric acid, talcum, magnesium stearate, calcium stear-
ate, lycopodium, and Clerosil on the static friction co-
efficient. From this study they concluded that the lubri-
cant powder effects a decrease of the coefficient of static
friction of sticky granules, but such ability differs with
different grades of lubricant. They suggested that it would
be useful to measure the static friction coefficient of
granules in checking the quality of batches of granules in
plant production.

Raff, Arambulo, and Deardorff (10) have also reported weight variation in tablets in their study of internal flow of granulation during compression. They used statistical quality control charts for tablet weight, hardness, and compressional pressure as an aid to study the internal flow of a granulation in the single punch tablet press.

In their study they used lactose-starch, sodium chloride, and aspirin-lactose-starch granulations. The granulations were prepared by a wet granulation method using starch paste as a binder. In the case of the lactose-starch granulation, granules were passed through a U. S. Standard Sieve #10 and retained on a #50. In the case of the aspirin-lactose-starch granulation, aspirin was passed through a #70 sieve and retained on a #100, and it was mixed with lactose-starch granulation. In the case of sodium chloride, two collections were made; one granulation was passed through a #10 sieve and retained on a #12; the other was passed through a #45 sieve and retained on a #50. Talc and magnesium stearate were used as lubricants for all granulations. Compression was carried out in the Stokes Model E single punch tablet press, using 11/32-inch punches. The speed of the machine was adjusted to produce approximately 100 tablets a minute.
In this study these workers found that, during the first part of the run, the weight of the tablets usually increased, but sometimes decreased, depending upon the nature of the granulation. In the case of the aspirin-lactose-starch granulation a substantial decrease in the weight of the tablets was observed. Results of studies in other fields indicate that, in order to avoid too rapid a flow of fines through coarser particles and consequent variation in weight of tablets, one should not have too wide a disparity in particle size, and further, the active ingredient should be of such a size as to fill substantially the average void space of the basic granulation. They also point out that their experiments confirmed the usual belief that it is important to check the tablet weights most closely during the first and last quarters of the run.

Hasegawa (12) has reported a study on tablet weight variation in which he considered four factors causing weight variation in tablets. The four factors studied were:

(1) the effect of three different tablet bases;
(2) the effect of punch diameter (7 m.m., 10 m.m., and 13 m.m. diameter punches);
(3) the effect of granule size (sieve size No. 12, No. 14, No. 16, No. 18 and No. 20);
(4) the effect of speed of tableting (42 tablets a minute and 76 tablets a minute).

The study was conducted using a Kimura Model KT2 tablet press. Potato starch was used as a binder, and 2% talcum was used as a lubricant.

From the results obtained during this study, he concluded that:

(1) significant differences were not recognized among different tablet bases;

(2) reduction of punch diameter increased weight variation of tablets quadratically;

(3) reduction of granule size decreased weight variation of tablets, and the relationship was linear; the degree of decrease of weight variation was different for each punch;

(4) significant differences were not recognized between the two tableting rates i.e. 42 and 76 tablets a minute.

Uniformity of drug dosage in compressed tablets was studied by Moskalyk, Chatten, and Pernarowski (14), who weighed and assayed individual tablets in order to determine the actual variability in drug dosage. The variations in potency found in a batch of tablets were as a rule greater than those indicated by the uniformity of weight test. Deviations were found greatest in the lightest
weight tablets within a batch. These investigators believe that, should this prove true for all tablets, then a test based on the analysis of a small number of lightest weight tablets or groups of tablets could be adopted as a control procedure to assure uniformity of drug dosage. Used in conjunction with the uniformity of weight test, it would provide a more effective control over excessive dosage variability in tablets.
III

EXPERIMENTAL PROCEDURE

A. General Order of Experiments

Preliminary experiments were carried out to establish whether conclusions from previous work could be applied to the tabletting set-up available for testing the hypothesis that tablet weight and weight variation is related to granule density and bulk density of granules. Thus, in the first series of experiments, the effects of granule size, tableting rate, and die size on tablet weight and weight variation were studied and the results compared with those in literature. The results of these experiments allowed the selection of a normal operating region for further experiments concerning the main object of this thesis, i.e., the interrelationship between granule density, bulk density, tablet weight, and weight variation. In the controlled density experiments, density of a basic granulation was varied by the addition of bismuth subcarbonate; the size of granules was controlled within a narrow range; bulk and granule density were determined, and the relationships among tablet weight, weight variation, and granule-bulk density were found, by compressing the granules on a rotary tablet machine using the same machine settings for all the granulations.
B. Preliminary Experiments to Test Applicability of the Literature Data

The following preliminary experiments were carried out to determine the applicability of the literature data.

1) Granules of three different sieve sizes were compressed on the tablet machine, maintaining constant speed of machine, pressure ejection, capacity, etc.

2) Speed of the tablet machine was increased, keeping pressure and capacity settings constant.

3) Two sets of standard concave dies and punches (3/8" and 3/16") were used on the tablet machine; the other settings were kept constant during tablet compression.

Granulation. The general formula for preparing granules was

| Lactose USP | 300.0 gms  |
| Starch Soln. 5% | 650 ml |

Preparation of Starch Solution. Starch paste was used as a binding agent to form the granulation. Commercially available corn starch, 50 gms, was put in a suitable container, one liter of cold water was added to it slowly, and with continuous stirring to avoid the formation of lumps. This mixture was slowly heated with continuous stirring, until the solution began to boil and a translucent paste resulted.
Preparation of Granules. The weighed amount of lactose was placed in a Hobart* mixer and starch paste was added gradually to the contents of the mixer while the mixer was rotating. Starch paste was slowly added and the mixture stirred until the mass readily cohered to a ball but equally readily broke up when rubbed between the fingers. This damp mass was then taken out of the mixer and granulated with an Erweka* wet granulator. The wet granules were collected on brown paper, put on trays, and finally dried in an oven at 70° for four hours.

The dried granules were then reduced to a smaller size using an Erweka dry granulator and separated using U. S. Standard No. 16, 20 and 40 sieves. The granules referred to as #16 mesh were passed through a No. 16 Standard sieve and were retained on a No. 20 sieve. In the case of 20 mesh granules, all granules were passed through a No. 20 and were retained on a No. 40 sieve. In the case of No. 40 granules all granules passed through #40 sieve and were retained on a #60 sieve. The granule and bulk densities were determined. (See III C - granule and bulk density experiments.)

*Hobart Manufacturing Company, Troy, Ohio

Lubrication. All granules were lubricated with 0.5% magnesium stearate prior to compression.

Tablet Compression. A Colton Model 216* rotary tablet press was used for the compression of all of the granulations. The settings and operating conditions for the compression of each granulation lot were varied in order to control certain variables.

Granule Size Experiments. In order to study the effects of granule size on tablet weight and weight variation, three different sizes of granules prepared and separated were used. Three-eighths inch standard concave dies and punches were set on the tablet machine. The tablet weight was adjusted to 450 mgms. by hand-turning machine when loaded with 16 mesh granules. Pressure roller settings and weight control settings were not changed during the compression of the other two granulations. The tableting rate was kept constant at 582 tab./min. during the compression of all three granulations. To determine the tableting rate, tablets produced during one minute of operation were collected and counted; and the count was repeated three times.

Samples of the tablets used to study the weight and weight variation were collected as follows: ten samples

of ten tablets each were collected during the run of tablet machine. Each tablet was weighed individually and the average weight of each group was calculated.

**Tableting Rate Experiments.** The effect of speed of compression on tablet weight variation of the three different size of granulations was determined.

**Compression of 16 Mesh Granules.** The granules passing through #16 and retained on #20 were lubricated with 0.5% magnesium stearate. The standard concave 3/8" dies and punches were used for this compression, and the weight of tablet was adjusted to 400 mgms. by hand-turning the machine. The tablets were compressed at five different machine speeds:

The tableting rates were increased gradually and evenly by rotating the speed control wheel six full rotations each time in clockwise direction. Tablets produced during three 1-minute intervals were counted and the average speed of compression was computed from the three counts.

**Compression of 20 Mesh Granules.** Granules passing through a #20 and retained on a #40 sieve were lubricated
as before. The tablet weight was adjusted to 450 mgms. by
hand-turning the machine, and the tablets were compressed,
using exactly the same settings of machine as for the #16
mesh granules. Samples were collected exactly as above.

Effect of Die Size. To determine the effect of die
size on tablet weight, the granules, prepared as described,
passing through a #20 mesh sieve and retained on #40 mesh
were used. The two different die sizes and punches were
used (3/8" and 3/16").

First, 3/8" dies were set on tablet press and the
#20 mesh lubricated granules, were compressed into tablets
at two different speeds. Speed of compression was deter-
mined by counting the number of tablets produced in one
minute (each three times) and then determining the average.
Three samples were collected at each speed and the average
weight was determined by calculation from the weight of
the individual tablets.

Similarly, 3/16" dies were set on tablet press and
same #20 mesh granules were compressed at three different
speeds and samples were collected as before, and percentage
of weight variation was determined at each speed of com-
pression.

C. Granule and Bulk Density Experiments

Preparation of Granules. For the preparation of
granules, the following ingredients were used:

Lactose U.S.P. Malinorodt G. 91R  
Bismuth subcarbonate U.S.P. Amend Drug Co.  
Gelatin U.S.P.

Method of Variation of Density of Granules. The density of the basic lactose granulation was increased by the addition of bismuth subcarbonate in concentrations of 10%, 20%, 30%, and 40%. Table I gives the actual formulas used to prepare each of these granulations.

Preparation of the Binder Solution. Gelatin binder solution was prepared by covering 10 gms. of gelatin U.S.P. with 40 ml of water and allowing the mixture to stand until the gelatin became hydrated. The mixture was then heated gently with frequent agitation until the gelatin was dissolved. A sufficient quantity of water was then added to make final volume measure 100 ml.

Mixing. Weighed quantities of lactose and bismuth subcarbonate powders were placed in a Hobart mixer, and the dry powders were mixed to uniformity, for about 3 - 5 minutes.

Granulation. To the uniform mixture of these powders in Hobart mixer warm gelatin solution was added

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Hobart Mfg. Co., Troy, Ohio
TABLE I

GRANULATION FORMULAS

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<tr>
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<tr>
<td>% Bismuth sub-carbonate</td>
<td>0%</td>
<td>10%</td>
<td>20%</td>
<td>30%</td>
<td>40%</td>
</tr>
<tr>
<td>Ingredients</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lactose</td>
<td>3000 gms</td>
<td>2700 gms</td>
<td>2400 gms</td>
<td>2100 gms</td>
<td>1500 gms</td>
</tr>
<tr>
<td>Bismuth sub-carbonate</td>
<td></td>
<td>300 gms</td>
<td>600 gms</td>
<td>900 gms</td>
<td>1200 gms</td>
</tr>
<tr>
<td>Gelatin Solution</td>
<td>375 ml</td>
<td>390 ml</td>
<td>405 ml</td>
<td>425 ml</td>
<td>460 ml</td>
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gradually, while mixer was rotating, and the ingredients mixed until the mass readily cohered to a ball but equally readily broke up when rubbed between the fingers. This damp mass was then taken out of the mixer and granulated with an Erweka wet granulator. The wet granules were collected on brown paper, put on trays, and dried in an oven at 70°F for four hours.

Separation into Size Fractions. The dried granules were then reduced to smaller size by hand forcing them first through U. S. Sieve #10 and then through a #16. The granules passed through the #16 sieve and were retained on a #20 sieve.

D. Determination of the Pertinent Characteristics of the Granulations

Size Distribution Determination. A stereo microscope with a calibrated eyepiece and having magnification of twenty times and fixed focal length was used to determine size distribution of the granules. One thousand particles of granules were counted from each lot of granulation. The "mean linear intercept" (1) method was used for the measurement of the granules. In this method the mean length of a line intercept by the profile boundary which approximately bisects the area of the granule profile is taken as the diameter of the granule. The bisecting
line is taken parallel to a fixed direction, irrespective to the orientation of each particle. This has the effect of avoiding bias as to the direction in which the profiles observed are bisected. From this method of diameter count, reproducible graphs for average size distribution were obtained.

**Bulk Density Determinations.** For the determination of bulk density of granules, the method suggested by Butler, Ramsey, and Martin (13) was used. A sample of about 50cc. of the sized granules was carefully introduced into a 100cc graduated cylinder. The cylinder was then dropped onto a hard wood surface (table top) three times from a height of about one inch at approximately two second intervals. Bulk density was then determined by dividing the weight of the sample of granules in grams by the final volume in cc. of the sample contained in the cylinder.

**Granule Density Determinations.** A mercury displacement method similar to the method suggested by Strickland, Bussa and Higuchi (6) was used to determine granule density.

Figure I shows a diagram of apparatus and assembly used to determine density of the granules. The internal volume of the glass bulb (labeled "A" in diagram) was determined by calculations from the weighed quantity of merc-
Figure 1 Apparatus to determine Granule Density.
ury required to fill the bulb to the calibration mark. Granule density was determined by placing a known weight of granules in the empty bulb, pumping the air out of the bulb with a vacuum pump, allowing mercury to flow into the bulb to the calibration mark. The bulb containing the mercury and the granules was then removed from the assembly and weighed. From the known weight of the bulb and the granules and the calibrated volume of bulb, the volume displaced by granules was calculated. The granule density was then calculated by dividing the weight of granules by the volume displaced.

E. Compression of Granules

Equipment. A Colton Model 216 rotary tablet machine was used for all tablet compression.

Selection of operating parameters. From the preliminary experimentation some working parameters or conditions were set up for the study of the effects of granule and bulk density on tablet weight variation. In the first place, the No. 16 granulations were selected because the No. 16 granules were the only ones that could be prepared with sufficient yield from the formulas containing the larger proportions of bismuth subcarbonate.

The one-fourth inch dies and standard concave punch-
es were used for tablet compression. The weight of tablet was adjusted 100 mgms. with the lactose granulation while turning the machine by hand. The other 'standard' settings of the machine, which were kept constant during compression of all granulations, were: speed of machine or tableting rate (650 tab/min); overload pressure adjustments; and flow of granules through hopper.

Lubrication. All granulations were lubricated with 0.5% magnesium stearate; 5% cornstarch powder was added during lubricating as a disintegrating agent.

Determination of Tablet Characteristics

Weight. As the tablets were being produced from the machine five samples of twenty-five tablets each were collected in separate boxes, during the compression of the different granulations (i.e., granules containing 10%, 20%, 30% and 40% bismuth subcarbonate). Each tablet in a group was weighed individually and the average weight, mean deviation, and standard deviations were calculated.

Size. Thickness and diameter of each of fifty tablets in each group were measured and the average of thickness and diameter was determined.

Hardness. A Pfizer* hardness tester was used.

Twenty tablets selected at random from each lot of tablets produced, were tested for hardness, and the average hardness was determined for each lot of tablets.

**Disintegration Time.** The U.S.P. method* (basket-rack assembly) was used. One tablet of the sample from the lot of tablets was placed in each of the six tubes of the basket and the apparatus was operated, using distilled water at 37° as the immersion fluid. The time required for the complete disintegration of each tablet was noted and the average of six readings was determined.

IV
RESULTS AND DISCUSSION
PRELIMINARY EXPERIMENTS

Granule Characteristics

The lactose granulations obtained in the first part of the experiment were pure white. As mentioned before, in this series of preliminary experiments, granule size was controlled by passage through U. S. Standard Sieves Nos. 16, 20 and 40. Detailed studies on these granules with regard to size distribution, bulk density, and granule density were not carried out.

Effect of Granule Size

From Table II, which shows the influence of three different sizes (i.e. Nos. 16, 20 and 40) of granules on tablet weight and weight variation, it can be seen that the tablet weight increased as granule size decreased, and that the weight variation decreased with decreasing granule size. These results correspond to those obtained by Arambulo and Deardorff (5) who studied the effects of granule size from sieve Nos. 4 to 170. Arambulo and Deardorff attributed the increasing weight with decreasing granule size to a decrease in void space. They reasoned that large granules gave large proportions of void space
# TABLE II

**Tablet Weight and Coefficient of Variation Obtained from 3 Different Sizes of Granulations**

<table>
<thead>
<tr>
<th>Tablet Weight in Mg. from Granulation</th>
<th>16 Mesh</th>
<th>20 Mesh</th>
<th>40 Mesh</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>424</td>
<td>482</td>
<td>508</td>
</tr>
<tr>
<td></td>
<td>444</td>
<td>485</td>
<td>507</td>
</tr>
<tr>
<td></td>
<td>448</td>
<td>478</td>
<td>509</td>
</tr>
<tr>
<td></td>
<td>439</td>
<td>503</td>
<td>512</td>
</tr>
<tr>
<td></td>
<td>450</td>
<td>508</td>
<td>520</td>
</tr>
<tr>
<td></td>
<td>426</td>
<td>497</td>
<td>524</td>
</tr>
<tr>
<td></td>
<td>449</td>
<td>490</td>
<td>519</td>
</tr>
<tr>
<td></td>
<td>461</td>
<td>499</td>
<td>521</td>
</tr>
<tr>
<td></td>
<td>421</td>
<td>493</td>
<td>518</td>
</tr>
<tr>
<td></td>
<td>431</td>
<td>482</td>
<td>522</td>
</tr>
<tr>
<td><strong>Average of tab.</strong></td>
<td>439.3</td>
<td>491.7</td>
<td>516</td>
</tr>
<tr>
<td><strong>Max. Deviation</strong></td>
<td>21.7</td>
<td>16.3</td>
<td>9</td>
</tr>
<tr>
<td><strong>Coef. of Variation</strong></td>
<td>0.08</td>
<td>0.05</td>
<td>0.03</td>
</tr>
</tbody>
</table>
in the die before compression, resulting in relatively light weight tablets. As the granule size was reduced, void space was reduced and the average tablet weight increased.

**Effect of Tableting Rate**

As shown in Table III and Figure 2, the average tablet weight decreased as the tableting rate was increased. This trend followed in both sizes of granules used. As the tableting rate was increased, there was less time for granules to flow into the die cavity from the feed shoe. In addition, there was a tendency for the granules to be thrown outward at higher rotational speeds. In the case of the No. 16 granulations, at the maximum rate of tableting a 22.4% drop in the average weight of tablets was observed, indicating either that the die cavities were incompletely filled by the granules or that granules were being thrown out of the dies by centrifugal force. The latter factor would seem to be less important than the former in view of the later experiments which showed that, as die size was decreased the weight variation of tablets was increased. In general, as the speed of compression was increased the weight of the tablets produced was decreased, the variation in weight from the average tablet weight was also found to increase as the
TABLE III

AVERAGE WEIGHT OF TABLETS FROM NO. 16 AND NO. 20 SIEVE GRANULES OBTAINED AT DIFFERENT SPE DS

<table>
<thead>
<tr>
<th>Tab/min</th>
<th>Av. Wt. mg</th>
<th>Coef. of Variation</th>
<th>Tab/min</th>
<th>Av. Wt. mg</th>
<th>Coef. of Variation</th>
</tr>
</thead>
<tbody>
<tr>
<td>392</td>
<td>376.6</td>
<td>0.16</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>508</td>
<td>375</td>
<td>0.19</td>
<td>580</td>
<td>438.0</td>
<td>0.11</td>
</tr>
<tr>
<td>632</td>
<td>374.2</td>
<td>0.20</td>
<td>724</td>
<td>428.0</td>
<td>0.12</td>
</tr>
<tr>
<td>768</td>
<td>370</td>
<td>0.25</td>
<td>912</td>
<td>409.3</td>
<td>0.14</td>
</tr>
<tr>
<td>932</td>
<td>345</td>
<td>0.38</td>
<td>1020</td>
<td>373.0</td>
<td>0.21</td>
</tr>
<tr>
<td>1180</td>
<td>290</td>
<td>0.46</td>
<td>1132</td>
<td>344.8</td>
<td>0.29</td>
</tr>
</tbody>
</table>

*hand turned weight 400 mg.  **hand turned weight 450 mg.
tabletting rate was increased. These results are shown in Table III and Figures 2 and 3. As a result of incomplete fill at higher speeds of compression a drop in the hardness of tablets was also observed. These results do not parallel those of Hasagawa who studied weight variation at two rates of tableting - 42 and 76 tab/min, and concluded that significant differences in tablet weight variation did not result at 76 and 42 tab/min tableting rates.

The difference between his results and the results reported here can probably be explained by the fact that Hasagawa used tableting rates of 76 and 42 tab/min which are very low compared to the 92 - 1180 tab/min rates used in these experiments. In addition Hasagawa used a single punch tablet machine (Kimura Model K.T. 2) whereas results reported here are from a Colton Model 216 rotary tablet machine.

Effect of Die Size (Diameter)

The effect of two different die sizes on weight variation at different speeds of compression is shown in Table IV. In the case of the smaller die diameter the coefficient of variation is higher and in the case of the larger die diameter, the coefficient of variation is less than former. The effect of die size on weight variation of the tablets is similar to data reported by Hasagawa. The reason for this type of effect is that larger die
Figure 2  Effect of tableting rate on coefficient of variance.

- Tablets from No. 20 sieve granules.
- Tablets from No. 16 sieve granules.
Figure 3  Effect of tableting rate on average weight of tablets.

- Tablets from No. 20 sieve granules.
- Tablets from No. 16 sieve granules.
cavities are easier to fill and fill evenly during the run of the tablet machine than smaller die cavities.

INTERRELATIONSHIP OF GRANULLE DENSITY, BULK DENSITY, TABLET WEIGHT AND WEIGHT VARIATION

Characteristics of Granulations

Size Distribution. In these studies the granule size was controlled within a narrow range and size distribution data were obtained to show this distribution. Tables V and VI show the size distribution of the granulations. In Table V only the No. 16 plain lactose granulation was considered, and four counts each of 1,000 granules were made to obtain the plain lactose size distribution plots. The average diameter of the granules in m.m. and the standard deviations in each count show that the results are very close. The reproducibility of these results is also shown by the standard error which is 1.35 ± 0.01. Table VI shows the size distribution of lactose granulations containing various concentrations (10%, 20%, 30%, 40%) of bismuth subcarbonate, compared with plain lactose granulations. It can be seen from these data that all the granulations had similar distributions, but the granulations containing 30% and 40% bismuth subcarbonate apparently contained slightly higher proportions
<table>
<thead>
<tr>
<th>Tab/min</th>
<th>Av. Wt. Mg</th>
<th>Coef. of Variance</th>
<th>Tab/min</th>
<th>Av. Wt. Mg</th>
<th>Coef. of Variance</th>
</tr>
</thead>
<tbody>
<tr>
<td>980</td>
<td>417.0</td>
<td>0.05</td>
<td>912</td>
<td>153</td>
<td>0.061</td>
</tr>
<tr>
<td>1130</td>
<td>396.0</td>
<td>0.10</td>
<td>1020</td>
<td>140.3</td>
<td>0.092</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>1132</td>
<td>127.0</td>
<td>0.16</td>
</tr>
</tbody>
</table>
TABLE V

PLAIN LACTOSE GRANULATION #16 MESH DIAMETER COUNT

| Size in m.m. | Count No. | 0.9 | 1.1 | 1.2 | 1.3 | 1.4 | 1.5 | 1.6 | 1.7 | 1.8 | 1.9 | 2.0 | 2.1 | 2.2 | 2.3 | Total Particles | Average Diameter | Standard Deviation |
|--------------|-----------|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|-----|----------------|-----------------|------------------|
| 1            | 1         | 14  | 42  | 107 | 133 | 226 | 148 | 137 | 80  | 48  | 30  | 14  | 10  | 6   | 3   | 2   | 1,000          | 1.37            | 0.228            |
| 2            | 1         | 14  | 41  | 121 | 136 | 228 | 149 | 130 | 74  | 46  | 29  | 13  | 10  | 4   | 2   | 0   | 1,000          | 1.36            | 0.222            |
| 3            | 1         | 12  | 60  | 105 | 143 | 208 | 141 | 131 | 65  | 43  | 27  | 14  | 10  | 7   | 5   | 3   | 1,000          | 1.36            | 0.229            |
| 4            | 1         | 16  | 59  | 121 | 157 | 230 | 150 | 112 | 62  | 41  | 30  | 10  | 6   | 4   | 2   | 0   | 1,000          | 1.34            | 0.222            |

\[ SE = 1.35 \pm .01 \]
of granules in the 1.0 to 1.2 m.m. range, resulting in a slightly lower average granule size. Plots of the size distributions of the various granulations are shown in Figures 4 and 5.

Granule Density. Table VII shows that granule density of the five different granulations was increased in proportion to the amount of Bismuth subcarbonate added. The granule density ranged from 1.488 for plain lactose granulations to 2.025 for the granules containing 40% Bismuth subcarbonate.

Bulk Density. The bulk density of the granulation is related to granule density, and when the density of the basic lactose granulation was increased by the addition of Bismuth subcarbonate, the bulk density of the granules was found almost proportionally increased (size distribution was controlled). Table VII shows the increase of bulk density as the concentration of Bismuth subcarbonate was increased. If the size distribution of granules had not been controlled, the bulk density of granules might have shown different a trend, since bulk density is also influenced by shape and size of granules.

Miscellaneous. The granulations prepared from plain lactose and those prepared by adding various con-
Figure 4 Granule size distribution of No. 16 sieve plain lactose granules.
Figure 5  Granule size distribution of No.16 sieve lactose granules containing various concentrations of Bismuth Subcarbonate.
TABLE VI

GRANULATION OF LACTOSE AND BISMUTH SUBCARBONATE

PARTICLE DIAMETER COUNT

<table>
<thead>
<tr>
<th>Particle Size in m.m.</th>
<th>.9</th>
<th>1</th>
<th>1.1</th>
<th>1.2</th>
<th>1.3</th>
<th>1.4</th>
<th>1.5</th>
<th>1.6</th>
<th>1.7</th>
<th>1.8</th>
<th>1.9</th>
<th>2</th>
<th>2.1</th>
<th>2.2</th>
<th>2.3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plain Lactose</td>
<td>14</td>
<td>42</td>
<td>107</td>
<td>133</td>
<td>226</td>
<td>148</td>
<td>137</td>
<td>80</td>
<td>48</td>
<td>30</td>
<td>14</td>
<td>10</td>
<td>6</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>Lactose + 10% Bismuth Subcarbonate</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Lactose + 20% Bismuth Subcarbonate</td>
<td>7</td>
<td>75</td>
<td>90</td>
<td>116</td>
<td>208</td>
<td>156</td>
<td>156</td>
<td>84</td>
<td>45</td>
<td>31</td>
<td>21</td>
<td>12</td>
<td>4</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Lactose + 30% Bismuth Subcarbonate</td>
<td>15</td>
<td>77</td>
<td>145</td>
<td>155</td>
<td>229</td>
<td>157</td>
<td>120</td>
<td>39</td>
<td>30</td>
<td>19</td>
<td>5</td>
<td>4</td>
<td>2</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Lactose + 40% Bismuth Subcarbonate</td>
<td>7</td>
<td>63</td>
<td>120</td>
<td>130</td>
<td>234</td>
<td>172</td>
<td>120</td>
<td>63</td>
<td>38</td>
<td>25</td>
<td>7</td>
<td>5</td>
<td>3</td>
<td>2</td>
<td>1</td>
</tr>
</tbody>
</table>

Total Particles: 1,000
Average: 1.37
Standard Deviation: .228
TABLE VII

GRANULE AND BULK DENSITY OF
DIFFERENT GRANULATIONS

<table>
<thead>
<tr>
<th>Granulations</th>
<th>Granule Density</th>
<th>Bulk Density</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plain Lactose</td>
<td>1.488</td>
<td>0.432</td>
</tr>
<tr>
<td>Lactose + 10% Bismuth subcarbonate</td>
<td>1.564</td>
<td>0.483</td>
</tr>
<tr>
<td>Lactose + 20% Bismuth subcarbonate</td>
<td>1.653</td>
<td>0.497</td>
</tr>
<tr>
<td>Lactose + 30% Bismuth subcarbonate</td>
<td>1.81</td>
<td>0.582</td>
</tr>
<tr>
<td>Lactose + 40% Bismuth subcarbonate</td>
<td>2.02</td>
<td>0.683</td>
</tr>
</tbody>
</table>
centrations of bismuth subcarbonate showed little change in gross appearance except with respect to color. The plain lactose granulations were pure white in color, but as the increasing proportions of bismuth subcarbonate (i.e. 10%, 20%, 30% and 40%) were added the white color was changed gradually to off white (or pale tan) color. The granulations containing bismuth subcarbonate were harder, possibly due to the larger amounts of binder solution required to prepare the granules. When magnified, it was observed that the bismuth subcarbonate granules were somewhat less regular in shape than those of the plain lactose granulations.

RELATIONSHIP OF BULK DENSITY TO GRANULE DENSITY

When the size and size distribution of granular particles are controlled within narrow limits, variations in other physical characteristics such as porosity and shape are reflected by changes in the granule density and bulk density.

Granule density as determined by the mercury displacement method, reflects changes in intraparticle porosity (due to totally enclosed air pockets, and micro-
pores too small for the mercury to enter at ordinary pressure) and changes in the average true density of the materials used to make up the granulation. Bulk density reflects the changes in granule density and interparticle void spaces. Since the interparticle void space is dependent upon the packing characteristic of the granulations, shape differences in the granulation will be reflected by the changes in the bulk density. The mathematical relationship as shown below:

In the mercury displacement method granule density

\[ P_g = \frac{\text{weight of granules}}{\text{vol. of mercury displaced}} \]

is given by the bulk volume, \( V_b \) - granule volume, \( V_g \).

Now, in the case of bulk density, the mass of the powder is divided by Bulk volume \( \frac{W_g}{V_b} \). Bulk volume includes Void Porosity which is defined as shown below:

\[
\text{Void Porosity} = \frac{V_b}{V_g} = 1 - \frac{\text{weight/granule density}}{\text{weight/bulk density}} = 1 - \frac{\text{Bulk density}}{\text{granule density}}
\]

From this derivation it can be seen that the bulk volume of granules and therefore bulk density is related to granule density. Now, the void porosity or inter
spaces can be controlled by controlling the size and shape of the granules. And, now if the size of the granules is controlled and density of the granules is increased, then, the bulk density should show changes parallel to the granule densities. Figure 9 shows that this parallel was obtained experimentally when granule size and size distribution were very carefully controlled. A nearly straight line relationship was obtained. As size distribution data (Figures 5 and 6) shows, all five granulations had very similar size distributions. It seems probable that, for a given type of granulation passing through a No. 16 sieve, the granules would have approximately the same void spaces, and therefore the bulk density would be increased as the granule density increased. The small deviation from linearity that was observed may have been due either to small differences in granule shape and consequent changes in packing characteristics or to the small changes in size distributions seen in Figures 5 and 6.

INTERRELATIONSHIPS OF TABLET WEIGHT, WEIGHT VARIATION AND GRANULE AND BULK DENSITY

Table VIII shows the interrelationship between granule and bulk density and tablet weight, weight variation. Tablet weight was seen to increase linearly, in proportion
Figure 6  Effect of Bismuth Subcarbonate on Granule Density.
<table>
<thead>
<tr>
<th>Granulation</th>
<th>Granule Density</th>
<th>Bulk Density</th>
<th>Average Weight</th>
<th>Mean Deviation</th>
<th>Standard Deviation</th>
<th>Coef. of Variance</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plain Lactose</td>
<td>1.488</td>
<td>.432</td>
<td>90.505</td>
<td>2.1154</td>
<td>2.568</td>
<td>0.0284</td>
</tr>
<tr>
<td>Lactose + 10% Bismuth subcarbonate</td>
<td>1.564</td>
<td>.483</td>
<td>98.637</td>
<td>2.127</td>
<td>2.627</td>
<td>0.026</td>
</tr>
<tr>
<td>Lactose + 20% Bismuth subcarbonate</td>
<td>1.653</td>
<td>.497</td>
<td>102.65</td>
<td>2.0032</td>
<td>2.580</td>
<td>0.025</td>
</tr>
<tr>
<td>Lactose + 30% Bismuth subcarbonate</td>
<td>1.61</td>
<td>.582</td>
<td>119.85</td>
<td>1.556</td>
<td>2.428</td>
<td>0.018</td>
</tr>
<tr>
<td>Lactose + 40% Bismuth subcarbonate</td>
<td>2.025</td>
<td>.6834</td>
<td>139.81</td>
<td>2.836</td>
<td>3.606</td>
<td>0.026</td>
</tr>
</tbody>
</table>
to the bulk density change. The relationship of tablet weight to granule density was somewhat more variable (as might be expected) but with the granulations used, a fairly close relationship was seen. This relationship would be expected to vary with particle shape, and with particle size and size distribution. The bulk density of plain lactose granules was 0.432 and the average weight of tablets prepared from these granules was 90.5 mg., compared to a bulk density 0.683 of lactose + 40% bismuth subcarbonate and an average weight of tablets from these granules of 139.8 mg. The coefficient of variation for plain lactose tablets was 0.028, and standard deviation was 2.568, compared to coefficient of variation 0.026 and standard deviation 3.606 that of lactose + 40% bismuth subcarbonate tablets. These figures show that bulk density of granules affected the average weight and weight variation of the tablets, and that the variation in weight within a granulation on a relative basis was practically identical. These data are shown in Figure 7 in which the tablet average weight is plotted against granule density and in Figure 8 in which tablet average weight is plotted against bulk density. The bulk density of granules containing 20% bismuth subcarbonate was found slightly lower than that expected. Although the size of the granules was controlled to within a narrow range, the size
Figure 7  Relationship between granule density and average weight of tablets.
Figure 8 Relationship between bulk density and average weight of tablets.
distribution data (Figures 4 and 5) show a slightly higher proportion of granules in the 1 mm - 1.3 mm region. This variation in distribution could have affected the bulk density of those granules containing 20% bismuth subcarbonate. Another factor affecting the bulk density, could have been shape of the particles and their packing characteristics. When the shapes of different granules were compared under the microscope, the plain lactose granules were found to be smooth and round, but those granules containing bismuth subcarbonate were a little more irregular in shape. The granules containing 20% bismuth subcarbonate had the least smooth surfaces. Both the size distribution and the shape differences could have caused bulk density lower than expected in this granulation.

CHARACTERISTICS OF THE TABLETS PRODUCED FROM THE DIFFERENT GRANULATIONS

The tablets obtained from the plain lactose granulation were pure white in color, but the tablets containing bismuth subcarbonate were somewhat off white in color, the color darkening as the proportion of the bismuth subcarbonate was increased. The hardness of the tablets was found to increase gradually as the percentage of bismuth subcarbonate added; and therefore, the density of granules
Figure 9 Relationship between Granule Density and Bulk Density.
<table>
<thead>
<tr>
<th>Granulation</th>
<th>Granule Density</th>
<th>Bulk Density</th>
<th>Average Weight</th>
<th>D. Time Secs</th>
<th>Hardness Kgs.</th>
<th>Thickness Inch</th>
<th>Diameter Inch</th>
<th>T. Density</th>
<th>Moisture</th>
</tr>
</thead>
<tbody>
<tr>
<td>Plain Lactose</td>
<td>1.488</td>
<td>.432</td>
<td>90.505</td>
<td>34.8</td>
<td>2.24</td>
<td>.1200</td>
<td>.251</td>
<td>1.499</td>
<td>1.61</td>
</tr>
<tr>
<td>Lactose + 10% Bismuth sub-</td>
<td>1.564</td>
<td>.483</td>
<td>98.367</td>
<td>35.5</td>
<td>2.595</td>
<td>.1212</td>
<td>.251</td>
<td>1.576</td>
<td>1.621</td>
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<tr>
<td>Lactose + 20% Bismuth sub-</td>
<td>1.653</td>
<td>.497</td>
<td>102.65</td>
<td>102.5</td>
<td>2.695</td>
<td>.1205</td>
<td>.251</td>
<td>1.643</td>
<td>1.731</td>
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<tr>
<td>Lactose + 30% Bismuth sub-</td>
<td>1.81</td>
<td>.582</td>
<td>119.85</td>
<td>135.5</td>
<td>3.105</td>
<td>.122</td>
<td>.2515</td>
<td>1.850</td>
<td>1.681</td>
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</tr>
<tr>
<td>Lactose + 40% Bismuth sub-</td>
<td>2.025</td>
<td>.6834</td>
<td>139.81</td>
<td>15 min</td>
<td>4.90</td>
<td>.124</td>
<td>.2510</td>
<td>2.21</td>
<td>1.701</td>
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was increased. This is seen in Table IX where hardness of tablets containing only lactose was 2.24 Kg. and hardness of tablets containing 40% bismuth subcarbonate was found to be 4.90 Kg.

This table also compares the disintegration time of tablets from five different granulations. It can be seen that the disintegration time was gradually increased with the increasing concentrations of bismuth subcarbonate.

The average disintegration time of plain lactose tablets was 34.8 seconds and the average disintegration time of tablets containing 40% bismuth subcarbonate was more than 15 minutes, even though 5% corn starch was added to each granulation as a disintegrating agent. This gradual increase in disintegration time was probably due to the increase in hardness of the tablets resulting in tablets less readily penetrated by water.

The tablet diameter in all kinds of tablets was found to be identical, but the tablet thickness was found to be slightly increased (0.1200 to 0.1249 inch) as the density of the granulations increased. The tablet density was found to follow the same trend as granule density.
SUMMARY AND CONCLUSIONS

A study of several factors affecting tablet weight and weight variation was carried out. In a preliminary series of experiments, lactose granulations were prepared and effects of granule size, speed of compression, and die diameter were studied, compressing tablets on a Colton Model 216 rotary table press. Results of these experiments were found to be similar to those reported in the literature. From the data and techniques of these experiments, some optimum conditions, such as granule size, rate of tableting, die diameter, etc., were set up to study the effects of granule and bulk density on tablet weight and weight variation. The density of a basic lactose granulation was increased by the addition of various proportions of bismuth subcarbonate (true density 6.86). Bulk and granule density of these granules was determined. The size of these granulations was controlled to within a narrow range (No. 16 and No. 20 sieves) and the average size distribution was determined by an optical method. All granulations were compressed into tablets on fixed standard settings of a Colton Model 216 rotary press and the following conclusions were drawn:

(1) The density of a lactose granulation can be
successfully increased by the addition of various proportions of heavy substances such as bismuth subcarbonate.

(2) When granule size distribution is carefully controlled within narrow limits, bulk and granule density are linearly related.

(3) The increase in the average weight of the tablets was observed to be a linear function of the bulk density of the granulations.

(4) A similar relationship was found to exist between granule density and tablet weight, as might be expected.

(5) The increase in the granule density and in bulk density also affected other physical properties of tablets such as hardness, thickness and disintegration time.

(6) The coefficient of weight variation of tablets within a granulation was found to be practically identical for all the different granulations even though bulk density of the granulations was varied from 0.432 to 0.6834. The standard deviations varied with the average weight of the tablets produced.
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REFERENCES

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12. Nasegawa, J., Japan J. Ph. and Chem. 5 15 (1957)