Analysis of Drug Content and Weight Uniformity for Half-Tablets of 6 Commonly Split Medications

Shaynan W. Hill, PharmD; Andrew S. Varker, PharmD; Kelly Karlage, BS; and Paul B. Myrdal, PhD

BACKGROUND: Cost savings can be achieved with the practice of tablet splitting. Previous research has shown weight nonuniformity within tablet halves. However, limited research to date has found that the potential dose inaccuracy resulting from splitting tablets does not significantly affect clinical outcomes.

OBJECTIVE: To determine the drug content and weight in split-half tablets of 6 commonly split medications using drug assay analysis.

METHODS: This study was performed by 2 fourth-year pharmacy students using 30 randomly selected tablets of each of the following 6 medications: warfarin sodium 5 milligrams (mg), simvastatin 80 mg, metoprolol succinate 200 mg, metoprolol tartrate 25 mg, cilostazol 40 mg, and lisinopril 40 mg. A randomly selected half of the tablets were split by a single pharmacy student using a tablet cutter, and the remaining tablets were kept whole. Drug content was analyzed for 15 whole tablets and 30 half-tablets for each of the 6 drugs using high performance liquid chromatography, an analytical tool used to identify and quantify substances in solution. Drug content uniformity was assessed by comparing drug content within half-tablets with one-half of the drug content mean found for all whole tablets in the sample. Weight uniformity was assessed by comparing half-tablet weights, as determined by a Mettler analytical balance, with one-half of the mean weight for whole tablets in the sample. The percentages by which each whole tablet’s or half-tablet’s drug content and weight differed from sample mean values were compared with proxy United States Pharmacopeia (USP) specification ranges for drug content (95%-105% for warfarin sodium and 90%-110% for the other 5 drugs). Additionally, these outcomes were compared for nonscored versus scored tablets. The percent relative standard deviation (%RSD, ratio of the standard deviation to the mean), a commonly used measure of the repeatability and precision of assays used to analyze drug content, was also calculated in order to determine whether the drugs met proxy USP specification for %RSD (less than 6% for all drugs studied).

RESULTS: A total of 43 of 180 half-tablets (23.9%) differed from sample mean values by a percentage that fell outside of proxy USP specification for drug content; warfarin sodium (11 of 30 half-tablets, 36.7%), simvastatin (3 of 30 half-tablets, 10.0%), metoprolol succinate (10 of 30 half-tablets, 33.3%), metoprolol tartrate (4 of 30 half-tablets, 13.3%), cilostazol (5 of 30 half-tablets, 16.7%), and lisinopril (10 of 30 half-tablets, 33.3%). Half-tablets outside of proxy USP specification for weight included warfarin sodium (10 of 30 half-tablets, 33.3%), metoprolol succinate (6 of 30 half-tablets, 20%), and lisinopril (7 of 30 half-tablets, 23.3%). The %RSDs for drug content and weight fell outside of the proxy USP specification for %RSD for metoprolol succinate (drug content = 8.98%, weight = 7.70%) and lisinopril (drug content = 10.41%, weight = 8.13%). Mean percent weight loss after splitting was less than 1% for all drugs except lisinopril, which had an average weight loss of 1.25%. The total numbers of scored (nonscored) tablet halves that fell outside of proxy USP specification were 20 (23) for drug content and 10 (13) for weight. When measuring drug content, the numbers of out-of-range half-tablets for scored (nonscored) were 36 (44) at 95%-105%, 9 (23) at 90%-110%, 0 (10) at 85%-115%, and 0 (1) at 75%-125%. When measuring weight, the numbers of out-of-range half-tablets for scored (nonscored) drugs were 28 (38) at 95%-105%, 0 (14) at 90%-110%, 0 (3) at 85%-115%, and 0 (0) at 75%-125%.

CONCLUSION: Dose variation exceeded a proxy USP specification for more than one-third of sampled half-tablets of warfarin sodium, metoprolol succinate, and lisinopril and appeared to be greater for nonscored tablets as compared with scored tablets. Drug content variation in half-tablets appeared to be attributable primarily to weight variation occurring when tablets powder or fragment during the splitting process. Therefore, equal daily doses will be determined by the ability of patients to split tablets perfectly in half.

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What is already known about this subject

• Manufacturers of FDA-approved medications are required to adhere to the United States Pharmacopeia (USP) established ranges for drug content of whole tablets. The USP has created guidelines to compare drug content for whole tablets; however, no guidelines have been established to assess the drug content of half-tablets.

• Teng et al. (2002) found that only 3 of 11 medications passed an adjusted USP uniformity test when half-tablet drug content uniformity was estimated from half-tablet weight after splitting of tablets by hand or razor blade.

• To date, 4 published studies have shown that the potential inaccuracy of dose resulting from splitting tablets does not significantly affect clinical and humanistic outcomes.

What this study adds

• This is the first study to determine drug content uniformity within half-tablets using drug assay; target drug content was defined as one-half of the per-tablet mean drug content for all whole tablets in a sample of 15 whole tablets each of 6 commonly split medications.

• More than 30% of measured half-tablets (n=30 each drug) of warfarin sodium, metoprolol succinate, and lisinopril differed from the target drug content by a percentage that fell outside of the proxy USP specification (95%-105% for warfarin sodium and 90%-110% for the other 5 drugs). For all other medications studied (simvastatin, metoprolol tartrate, and cilostazol), 10%-17% of half-tablets fell outside of proxy USP specification for drug content.

• Only 5 of 180 (2.8%) half-tablets in a weight-adjusted analysis, as compared with 43 of 180 (23.9%) half-tablets in an analysis that was not weight-adjusted, fell outside of the proxy USP specification for drug content. Thus, drug content variation in half-tablets appears to be attributable primarily to weight variation occurring when tablets powder or fragment during the splitting process.
Tablet splitting has become increasingly common, especially within the geriatric and psychiatric communities, as a means of reducing medication dose and/or cost. Physicians frequently write prescriptions for half-tablets in order to achieve doses less than the smallest available manufactured strength. Prescribers also write for half- and quarter-tablet doses of higher-strength tablets in order to reduce costs because parity pricing (the use of flat rates for medications independent of dose strength) is common. Cohen and Cohen (2000) showed that an annual savings of $1.45 billion could be achieved when tablet splitting was performed for 12 specific psychotropic medications, while annual savings of $725 million and $325 million could be achieved from splitting one-half and one-fourth of prescriptions written, respectively. Cohen and Cohen later (2002) estimated potential cost savings of $1.7 billion nationally if tablet splitting was performed for 7 antidepressant medications. Miller et al. (2007) found that tablet splitting contributed $342,239 (about $1.30 per member per month [PMPM]) or 17.3% of total annualized savings of $1,983,153 attributed to 4 managed care interventions that included (a) moving certain brand name drugs in 6 drug classes to nonpreferred status, (b) removing low-sedating $1.30 PMPM.8

Although cost savings may be accomplished, problems may arise with tablet splitting such as poor cognitive function or memory, the inability of patients to effectively split tablets, and the fear of inaccurate dose. The “Uniformity of Dosage Units” section in the U.S. Pharmacopeia (USP) manual states that each unit within a single lot of a given medication should have drug substance content that is within a narrow range around the labeled claim. Several studies have looked at weight variation of split tablets as a means of estimating drug content uniformity. Teng et al. (2002) evaluated the weight uniformity of 11 commonly split medications through an analysis of half-tablet weights, using a uniformity test adapted from the USP specifications. Eight medications were split using a single-edged razor blade, and 3 were split by hand (i.e., using only tensile strength of the fingers or hands). Results revealed that only 3 of 11 medications passed their adapted USP specifications, and there were no obvious tablet features (e.g., scoring) that determined whether a tablet would pass or fail the uniformity test. This study also found that tablets split by hand showed less uniformity than did tablets split using a razor blade, even though splitting tablets by hand produced cleaner splits with less tablet crumbling.

A similar study performed by Polli et al. assessed content uniformity through the analysis of half-tablet weights using the same adapted USP methods as used by Teng et al.9 In contrast to the results found by Teng et al., this study, performed at a Department of Veterans Affairs (VA) center, found that 8 out of 12 medications split with a tablet-splitting device passed the adapted uniformity test. Another study analyzed the drug weight uniformity of cyclobenzaprine tablets split in half using either a tablet splitter or a kitchen knife. The results showed that both methods resulted in a wide variation in fragment weight between 49.9% to 149.5% of the theoretical weight (defined as one-half of the mean weight of the intact tablet) using a kitchen knife and 69.4% to 130.2% using the tablet splitter. Thus, both methods failed the quality standards for dosage uniformity of manufactured drugs as outlined by Teng et al. and Rosenberg et al. (2002) evaluated content uniformity of discontinued pharmacist-dispensed split-tablet samples taken from 4 long-term care facilities using a total of 560 fragments. These authors found that 2 of 22 medications had significantly different fragment weights as compared with the theoretical weight of the half-tablets. The researchers also found that 30 of the 560 fragments deviated by more than 15% of the sample mean fragment weight, and 32 of the 560 fragments deviated by more than 15% of the theoretical weight. Lastly, 15 of the 22 medications were found to have relative standard deviations for weight expressed as a percentage (%RSDs) in excess of 6%, the upper limit of the USP specification.

Although studies of weight differences among split tablets have been performed, the more important analysis of drug content has yet to be explored. Studies to date have assessed drug content uniformity only as variation in half-tablet weights. These studies adapted the USP manual section entitled “Uniformity of Dosage Units”—criteria developed to ensure that actual drug content is equivalent to manufacturer-labeled drug content—and indirectly measured half-tablet drug content by measuring half-tablet weight. Although USP guidelines enforce strict adherence to drug content per dosage unit for whole tablets, guidelines for the drug content of split tablets have yet to be established.

In the present study, we defined the target drug content and weight of a half-tablet as equal to one-half of the mean drug content and weight, respectively, for all whole tablets in a sample of 6 commonly split medications. To assess the acceptability of variation in the half-tablets, defined as the percentage by which each individual whole tablet and half-tablet differed from the sample mean values, we adapted USP specifications for drug content and weight of whole tablets (proxy USP specification). We hypothesized that the drug content and weight of half-tablets would deviate from these proxy specifications.

Methods
Six drugs were studied: warfarin sodium 5 milligrams (mg), simvastatin 80 mg, metoprolol succinate 200 mg, metoprolol tartrate 25 mg, citalopram 40 mg, and lisinopril 40 mg (Table 1). These drugs were chosen because they were observed to be commonly split within a single VA health care network. A total of 30 whole tablets were randomly selected from each medication lot for each of the 6 drugs. All 30 whole tablets were weighed using a Mettler
Toledo AG204 (Mettler Toledo, Inc., Columbus, Ohio) analytical balance that is accurate to 0.1 mg. Fifteen of the 30 randomly selected tablets were split in half by a single pharmacy student (SH), using a Locking Tablet Cutter (Apothecary Products, Inc.), and weighed with the Mettler Toledo analytical balance.

The 15 whole tablets and 30 half-tablets for each of the 6 drugs were then dissolved separately using a combination of manual agitation and sonication techniques in an appropriate diluent adapted from respective USP official monographs. All tablets were assayed in accordance with USP methodology for determining content uniformity for whole tablets. Assay parameters for each drug were taken directly from USP monographs,\(^5\) customary changes were made to allow for column optimization (Appendix).\(^3\) After the tablets were completely dissolved, samples of each solution were assayed for drug concentration via a Waters Alliance High Pressure Chromatography system, consisting of a 2695 Separations Module coupled with a 2487 Dual Wavelength ultraviolet (UV) detector (Waters Corporation, Milford, MA). A standard curve was created for each drug, using pure drug powder (obtained from LKT Laboratories, St. Paul, MN, or Sigma-Aldrich, St. Louis, MO) diluted to 5 known concentrations. These standard curves were established to verify accurate analysis of the drug, as opposed to any inactive tablet constituents, and to quantify drug content by calculating concentration from area under the curve data obtained through high performance liquid chromatography (HPLC) analysis of whole- and half-tablet samples.

The following parameters were assessed for each of the 6 medications (Table 2):
1. Measured drug content:
   a. Each whole tablet’s drug content (n=15) was compared with the target drug content for whole-tablets, defined as one-half of the mean measured drug content for all whole tablets in the sample.
   b. Each half-tablet’s drug content (n=30) was compared with the target drug content for half-tablets, defined as one-half of the mean measured drug content for all whole tablets in the sample.
   c. Each nonscored tablet’s drug content (n=90) was compared with the target drug content for whole tablets, defined as one-half of the mean measured drug content for all whole tablets in the sample.

2. Weight-adjusted target drug content: To account for tablet powdering/fragmenting and the inability to split tablets into perfectly equal halves, each half-tablet’s target drug content (n=30) was adjusted for the weight of the fragment. The adjustment formula assumed that within a single half-tablet of known weight, the half-tablet’s proportion of the whole-tablet drug content should equal the half-tablet’s proportion of the whole-tablet weight (e.g., if a half-tablet was 51% of the whole-tablet weight, it should equal 51% of the whole-tablet target drug content).

3. Measured weight:
   a. Each whole tablet’s weight (n=15) was compared with the target weight for whole tablets, defined as the mean measured weight for all whole tablets in the sample.
   b. Each half-tablet’s weight (n=30) was compared with one-half of the target weight for whole tablets.

4. For each tablet, the percentage weight loss due to fragmenting and/or powdering was calculated as ([measured weight of whole tablet – measured weight of both half-tablets] / measured weight of whole tablet) x 100.

5. Nonscored tablet tablets (n=90; simvastatin, metoprolol succinate, and lisinopril) were compared with scored drug tablets (n=90; warfarin sodium, citalopram, and metoprolol tartrate) on 2 outcome measures, half-tablet drug content and half-tablet weight.

To assess the amount and acceptability of variations in drug content and weight, several measures were calculated. The measured drug content expressed as a percent of target drug content was calculated for both whole and half-tablets using the following equation: ([target drug content – measured drug content] / target drug content) x 100. Individual values for whole tablets should
TABLE 2 Definitions of Terms

<table>
<thead>
<tr>
<th>Definition</th>
<th>Description</th>
</tr>
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<tbody>
<tr>
<td>United States Pharmacopeia (USP)</td>
<td>A publication that contains legally recognized standards of identity, strength, quality, purity, packaging, and labeling for drug substances, dosage forms, and other therapeutic products, including nutritionals and dietary supplements.</td>
</tr>
<tr>
<td>Percent (%) RSD</td>
<td>Ratio of the standard deviation to the mean, standard deviation for measured variable x 100 measured variable mean</td>
</tr>
<tr>
<td>5-point standard curve</td>
<td>A curve consisting of 5 known concentrations of the drug created using pure drug—used to determine that the drug was isolated by HPLC as opposed to other tablet constituents.</td>
</tr>
<tr>
<td>High Performance (Pressure) Liquid Chromatography (HPLC)</td>
<td>A form of liquid chromatography used to separate compounds that are dissolved in solution.</td>
</tr>
<tr>
<td>Measured drug content</td>
<td>The amount of drug (mg) determined to be within the whole or half-tablet analyzed using HPLC.</td>
</tr>
<tr>
<td>Measured weight</td>
<td>The weight (mg) of the whole or half-tablet as measured using a Mettler analytical balance.</td>
</tr>
</tbody>
</table>
| Target drug content for individual tablet or half-tablet (measured drug content mean per tablet for sample) | Whole tablets: Σ drug content for whole tablets number of whole tablets  
Half-tablets: Σ drug content for half-tablets number of half-tablets  
| Target weight for individual tablet or half-tablet (measured weight mean per tablet for sample) | Whole tablets: Σ weight for whole tablets number of whole tablets  
Half-tablets: Σ weight for half-tablets number of half-tablets  
| Weight-adjusted target drug content                                      | The amount expected to be found within a single half-tablet of known weight, assuming that the half-tablet's proportion of whole tablet drug content equals the half-tablet's proportion of whole tablet weight. measured half-tablet weight x target drug content for whole tablets measured whole tablet weight  
| Percent of weight-adjusted drug content                                  | Measured half-tablet drug content as a percent of weight-adjusted target drug content. measured drug content for half-tablet x 100 weight-adjusted target drug content  
| Percent of target drug content                                           | For each tablet or half-tablet, expresses measured drug content as a percentage of target drug content.  
Whole tablets: Measured drug content for whole tablet x 100 Target drug content for whole tablets  
Half-tablets: Measured drug content for half-tablet x 100 Target drug content for half-tablets  
| Percent of target weight                                                 | For each tablet or half-tablet, expresses measured weight as a percentage of target weight.  
Whole tablets: Measured weight for whole tablet x 100 Target weight for whole tablets  
Half-tablets: Measured weight for half-tablet x 100 Target weight for half-tablets  
| Mean percent weight loss                                                 | The amount of drug loss caused by the splitting process. weight of whole tablet - weight of half-tablet #1 - weight of half-tablet #2 x 100 weight of whole tablet  
| Proxy USP specification for drug content                                 | Measured drug content of whole or half-tablets within 95%-105% of target drug content for half-tablets for warfarin sodium and within 90%-110% of target drug content for half-tablets for simvastatin, metoprolol succinate, metoprolol tartrate, citalopram, and lisinopril.  
| Proxy USP specification for weight                                       | Measured weight of whole or half-tablets within 95%-105% of target weight for half-tablets for warfarin sodium and within 90%-110% of target weight for half-tablets for simvastatin, metoprolol succinate, metoprolol tartrate, citalopram, and lisinopril.  
| Proxy USP specification for %RSD                                         | %RSD for whole or half-tables less than 6%.  

HPLC = high performance liquid chromatography; mg = milligrams; RSD = relative standard deviation.

fall within 95%-105% for warfarin sodium and 90%-110% for the other 5 drugs studied (proxy USP specification for drug content) according to the individual USP drug monographs. Because no USP criteria for drug content uniformity of half-tablets have yet been established, this study applied the proxy USP specification for whole tablets to half-tablets. It should be noted that the proxy USP specification ranges chosen for this study are the more stringent of ranges described within the individual USP monographs.

The tighter range is typically applied to samples of 20 or greater. However, individual tablets are typically subjected to a specification range of 85%-115%.

Relative standard deviation expressed as a percentage (%RSD), which is a ratio of the standard deviation to the mean of the variable being analyzed, was calculated for whole tablets (drug content and weight) and for half-tablets (drug content, weight-adjusted drug content, and weight). The %RSD is widely used to
assess the repeatability and precision of the assays used to analyze drug content. The %RSD for drug content for all drugs studied was calculated using the following equation: (standard deviation for measured drug content / measured drug content mean) x 100. The %RSD for weight for all drugs studied was calculated using the following equation: (standard deviation for measured weight / measured weight mean) x 100. Individual medication lots for whole tablets are targeted to have a %RSD less than 6% (proxy USP specification for %RSD).3

The percentage by which weight-adjusted drug content differed from target drug content was calculated using the following equation: (measured drug content for half-tablet / weight-adjusted target drug content for half-tablet) x 100.

### Results

#### Drug Content

For all whole tablets studied, measured drug content expressed as a percent of target drug content was found to fall within the proxy USP specification percentage range (Table 3). All whole tablets also met the proxy USP specification for %RSD. The measured drug content expressed as a percent of target drug content for half-tablets fell outside of the proxy USP specification for drug content for at least 3 half-tablets of each drug studied. A total of 43 of 180 half-tablets (23.9%) fell outside of the proxy USP specification for drug content; warfarin sodium (11 of 30 half-tablets, 36.7%), simvastatin (3 of 30 half-tablets, 10.0%) metoprolol succinate (10 of 30 half-tablets, 33.3%), metoprolol tartrate (4 of 30 half-tablets, 13.3%), citalopram (5 of 30 half-tablets, 16.7%), and lisinopril (10 of 30 half-tablets, 33.3%). The measured drug content variations expressed as a percent of target drug content for half-tablets were, from smallest to the largest, warfarin sodium (90.01-109.40%), simvastatin (95.21%-111.35%), metoprolol succinate (82.77%-115.92%), metoprolol tartrate (94.83%-112.37%), citalopram (96.50-111.93%), and lisinopril (81.15%-125.72%). Metoprolol succinate and lisinopril were the only agents analyzed that fell outside of the proxy USP specification for %RSD, with %RSD values of 8.98% and 10.41%, respectively.

Weight-adjusted drug content expressed as a percent of target drug content for half-tablets fell outside of the proxy USP specification for drug content for at least 1 half-tablet of 3 drugs—warfarin sodium, citalopram, and lisinopril (Table 3). A total of 5 of 180 half-tablets (2.78%) fell outside of the proxy USP specification for drug content for at least 1 half-tablet of each drug studied. A total of 43 of 180 half-tablets (23.9%) fell outside of the proxy USP specification for drug content; warfarin sodium (11 of 30 half-tablets, 36.7%), simvastatin (3 of 30 half-tablets, 10.0%) metoprolol succinate (10 of 30 half-tablets, 33.3%), metoprolol tartrate (4 of 30 half-tablets, 13.3%), citalopram (5 of 30 half-tablets, 16.7%), and lisinopril (10 of 30 half-tablets, 33.3%). The measured drug content variations expressed as a percent of target drug content for half-tablets were, from smallest to the largest, warfarin sodium (90.01-109.40%), simvastatin (95.21%-111.35%), metoprolol succinate (82.77%-115.92%), metoprolol tartrate (94.83%-112.37%), citalopram (96.50-111.93%), and lisinopril (81.15%-125.72%). Metoprolol succinate and lisinopril were the only agents analyzed that fell outside of the proxy USP specification for %RSD, with %RSD values of 8.98% and 10.41%, respectively.
USP specification for drug content after weight adjustment; these included warfarin sodium (3 of 30 half-tablets, 10%), citalopram (1 of 30 half-tablets, 3.3%), and lisinopril (1 of 30 half-tablets, 3.3%). Metoprolol succinate and lisinopril were the only agents analyzed that fell outside of the proxy USP specification for %RSD after weight adjustment, with %RSD values of 7.67% and 8.07%, respectively.

### Weight

For all whole tablets studied, measured weight expressed as a percent of target weight was found to fall within the proxy USP specification for weight (Table 4). All whole tablets also met the proxy USP specification for %RSD. Measured weight expressed as a percent of target weight for half-tablets fell outside of the proxy USP specification for weight for at least 6 half-tablets of warfarin sodium, metoprolol succinate, and lisinopril. A total of 23 of 180 half-tablets (12.8%) fell outside of the proxy USP specification for weight; these included warfarin sodium (10 of 30 half-tablets, 33.3%), metoprolol succinate (6 of 30 half-tablets, 20.0%), and lisinopril (7 of 30 half-tablets, 23.3%). Metoprolol succinate and lisinopril were the only agents analyzed that fell outside of the proxy USP specification for %RSD, with %RSD values of 7.70% and 8.13%, respectively.

Mean percent weight loss, after splitting, was less than 1% for all drugs with the exception of lisinopril: warfarin sodium (0.50%), simvastatin (0.08%), metoprolol succinate (0.17%), metoprolol tartrate (0.57%), citalopram (0.24%), and lisinopril (1.25%; Table 4).

### Scored Versus Nonscored Tablets

A total of 20 of 90 (22.2%) half-tablets of scored medications, and a total of 23 of 90 (25.6%) half-tablets of nonscored medications fell outside of the proxy USP specification for drug content (Table 5). The numbers of half-tablets for scored (nonscored) drugs falling outside of range for drug content were 36 (44) for 95%-105%, 9 (23) for 90%-110%, 0 (10) for 85%-115%, and 0 (1) for 75%-125%.

A total of 10 of 90 (11.1%) half-tablets of scored medications, and a total of 13 of 90 (14.4%) half-tablets of nonscored medications fell outside of the proxy USP specification for weight (Table 6). The numbers of half-tablets for scored (nonscored) drugs falling outside of range for weight were 28 (38) for 95%-105%, 0 (14) for 90%-110%, 0 (3) for 85%-115%, and 0 (0) for 75%-125%.

### Discussion

When measured half-tablet drug content was compared against target drug content (one-half of the sample mean drug content) for each of 6 study medications, 43 of 180 half-tablets (23.9%) fell outside of a proxy USP specification percentage range. Warfarin sodium had the highest number of half-tablets falling out of its proxy specification range, most likely due to its narrower specification window of 95%-105%. Metoprolol succinate and lisinopril were found to have a relatively large number of half-tablets with drug content falling outside of the range of 90%-110%. Variation in half-tablet drug content was greatest with lisinopril, which had tablet halves ranging from 81.15% to 125.72% of the target drug content.
content for half-tablets. Thus, when tablet splitting is performed for this lot of lisinopril, patients may receive daily doses that vary by as much as 45%.

Several potential reasons could explain the observed variation in lisinopril half-tablet drug content. Inaccuracy during the tablet splitting process may have produced variability between tablet halves due to unequal half-tablet size. This argument is supported by the weight-adjusted data: Only a single lisinopril half-tablet fell outside of the range of 90%-110% when half-tablet drug content was adjusted for weight as compared with 10 half-tablets when the data were not adjusted for weight. The results for lisinopril may also have been affected by weight loss due to the powdering and fragmenting that occurred during tablet splitting. Although lisinopril half-tablets were shown to have a mean percent weight loss of 1.25%, the majority of lisinopril half-tablets did not fall outside the proxy USP specification for drug content until the weight loss was greater than 1.72%

Both metoprolol succinate and metoprolol tartrate may also have been affected by the inability of the tablet splitting device to accurately split medications into 2 equal halves. Additionally, for metoprolol succinate, a greater percent of drug content variation may be attributed to tablet formulation, specifically regarding the sustained release mechanism. This drug is found within nonsolvin pellets intended to provide a slow release of the drug. Without the ability to visualize the inside of individual drug pellets, complete dissolution could not be easily determined.

Metoprolol succinate and lisinopril were shown to have %RSD values greater than 6% with regard to drug content and weight. This finding indicates that tablets of these medications are not easily split into equal halves when using a tablet-splitting device. Lisinopril had the greatest degree of drug content variability, with a %RSD of greater than 10%, perhaps in part because lisinopril had the greatest amount of weight loss from splitting. The high level of variability for metoprolol succinate may be due to the extended release drug delivery system.

Following the splitting process, tablet-weight measurements revealed unequal splitting; this finding is likely a result of tablet powdering and limitations of the tablet splitter, person, and device. When half-tablet drug content was adjusted for weight, a large reduction in drug content variation was found. Thus, half-tablet weight appears to be directly correlated with drug content. When compared with the target drug content of a perfectly split tablet half, 43 of 180 half-tablets (23.9%), but only 5 of 180 weight-adjusted half-tablets (2.8%), fell outside of proxy USP specification for drug content. Warfarin sodium accounted for the majority of weight-adjusted half-tablets falling outside of proxy

### TABLE 5
Comparison of Scored and Nonscored Half-Tablets: Drug Content

<table>
<thead>
<tr>
<th>Tablet type</th>
<th>n</th>
<th>Percent of Target Drug Content - Range</th>
<th>Outside of Proxy USP Specification</th>
<th>Out of Range (95% - 105%)</th>
<th>Out of Range (90% - 110%)</th>
<th>Out of Range (85% - 115%)</th>
<th>Out of Range (75% - 125%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scored</td>
<td>90</td>
<td>89.85 – 112.37</td>
<td>20 (22.2%)</td>
<td>36 (40.0%)</td>
<td>9 (10.0%)</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Nonscored</td>
<td>90</td>
<td>81.15 – 125.72</td>
<td>23 (25.6%)</td>
<td>44 (48.9%)</td>
<td>23 (25.9%)</td>
<td>10 (11.1%)</td>
<td>1 (1.1%)</td>
</tr>
</tbody>
</table>

A range (smallest to largest) representing measured drug content for whole or half-tablets expressed as a percent of target drug content.

B Number of half-tablets with measured drug content not within 95%-105% of target drug content for half-tablets for warfarin sodium or 90%-110% of target drug content for half-tables for simvastatin, metoprolol succinate, metoprolol tartrate, citalopram, and lisinopril.

C Determined by HPLC. The number (%) of half-tablets that fell outside of the range listed for drug content expressed as a percentage of target half-tablet drug content.

D Warfarin sodium, metoprolol tartrate, and citalopram tablets were scored.

E Simvastatin, metoprolol succinate, and lisinopril tablets were not scored.

HPLC = high performance liquid chromatography; USP = United States Pharmacopeia.

### TABLE 6
Comparison of Scored and Nonscored Half-Tablets: Weight

<table>
<thead>
<tr>
<th>Tablet type</th>
<th>n</th>
<th>Percent of Mean Weight - Range</th>
<th>Outside of proxy USP Specification</th>
<th>Out of Range (95% - 105%)</th>
<th>Out of Range (90% - 110%)</th>
<th>Out of Range (85% - 115%)</th>
<th>Out of Range (75% - 125%)</th>
</tr>
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<tbody>
<tr>
<td>Scored</td>
<td>90</td>
<td>90.69 – 109.05</td>
<td>10 (11.1%)</td>
<td>28 (31.1%)</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Nonscored</td>
<td>90</td>
<td>82.16 – 113.27</td>
<td>13 (14.4%)</td>
<td>38 (42.2%)</td>
<td>14 (15.6%)</td>
<td>3 (3.3%)</td>
<td>0</td>
</tr>
</tbody>
</table>

A range (smallest to largest) representing measured weight for whole or half-tablets expressed as a percent of target weight.

B Number (%) of half-tablets with measured weight not within the 95%-105% specification range for warfarin sodium or 90%-110% for the other 5 drugs.

C Determined by Mettler analytical balance. The number (%) of half-tablets that fell outside of the range listed for drug weight expressed as a percentage of target half-tablet drug weight.

D Warfarin sodium, metoprolol tartrate, and citalopram tablets were scored.

E Simvastatin, metoprolol succinate, and lisinopril tablets were not scored.

USP = United States Pharmacopeia.
USP specification for drug content (3 of the 5 half-tablets). It was also observed that the %RSD for weight-adjusted drug content for all medications was reduced in comparison with non-weight-adjusted drug content. These results would appear to indicate that the drug is uniformly dispersed within single whole tablets.

The data suggest greater variability in half-tablet drug content and weight for nonscored medications than for scored medications. Although nonscored and scored tablets produced roughly an equivalent number of half-tablets falling outside of proxy USP specification for drug content and weight, it was found that greater variability existed with the nonscored drug tablets. More nonscored drug half-tablets were found to have drug content and weight within the ranges of 85%-115% and 75%-125%. This finding suggests that when a tablet-splitting device is used, dose administration may be more accurate and consistent for scored than nonscored medications; however, a larger sample of scored and nonscored tablets is needed to determine if there is a significant difference between scored and nonscored tablets.

The pharmacokinetics and the mechanisms by which these medications act would appear to dictate that half-tablet regimens may or may not have a clinical impact on long-term patient outcomes. Metoprolol succinate, lisinopril, and citalopram are agents with long durations of action, in which minor dose variation should have no significant impact on steady state plasma concentrations. Additionally, citalopram efficacy is highly subjective; thus, daily efficacy measurements can be variable regardless of small dose variation. Statins, including simvastatin, are agents designed to prevent downstream medical problems such as acute coronary syndromes and stroke; thus, small changes in daily dose should have no significant impact on long-term clinical endpoints. Lastly, antihypertensives, including angiotensin-converting enzyme inhibitors and beta blockers, are used to prevent medical problems associated with an elevated blood pressure over an extended period of time, thus, daily fluctuations in dose would be expected to affect blood pressure measurements and side effects and have no effect on long-term clinical end points.

In contrast, caution should be taken when splitting warfarin sodium due to the potential for significant adverse events with minimal change in daily dose. However, daily variation of international normalized ratio (INR) values, the parameter used to monitor warfarin sodium efficacy, can result from food interactions, drug interactions, and variations in daily dose. For this reason, it cannot be stated that the minor differences in warfarin sodium half-tablet drug content will predict clinical outcomes.

Comparison With Previous Research and Clinical Significance
In order to determine true clinical significance of tablet splitting, studies looking at clinical outcomes must be examined. Four studies known to these authors have evaluated the clinical impact of half-tablet regimens: 3 assessing statins and 1 assessing lisinopril.10-13 All 4 studies have shown that the dose inaccuracy experienced from splitting tablets does not significantly affect primary outcomes. The study by Duncan et al. (2002) performed at a VA medical center examined triglycerides, low-density lipoprotein cholesterol (LDL-C), high-density lipoprotein cholesterol (HDL-C), and total cholesterol values for 109 patients enrolled in a statin (atorvastatin and simvastatin) tablet splitting program.10 The study concluded that there was a significant decrease in total cholesterol (187.6 mg per deciliter [dL] to 179.7 mg per dL; \( P = 0.005 \)) and LDL-C (111.6 mg per dL to 105.1 mg per dL; \( P = 0.004 \)) after at least 6 weeks following the initiation of the split tablet program. Duncan et al. concluded that there was no clinically significant difference comparing the time periods before and after the initiation of the tablet-splitting program.

A similar study performed at a different VA medical center assessed clinical endpoints before and after the initiation of a tablet-splitting program for 2,019 patients.11 This study by Gee et al. (2002) also found no clinically significant changes in serum lipid levels before and after implementation of the tablet-splitting program. A more recent retrospective chart review was performed across 6 VA medical centers, comparing 3,196 patients assigned to a split-tablet regimen with a whole-tablet regimen of varying simvastatin doses.12 Similar to the other previously mentioned studies, no statistically significant difference in LDL-C was found between patients in the split-tablet group and in the whole-tablet comparison group (\( P = 0.304 \)). A randomized crossover study performed at another VA medical center found no statistically significant differences in systolic or diastolic blood pressure for patients treated with whole- and half-tablet regimens for lisinopril.13 No studies to date have been performed that assess the clinical impact of half-tablet regimens for citalopram, metoprolol tartrate, metoprolol succinate, or warfarin sodium; thus, no conclusions about the clinical impact of half-tablet regimens for these agents can be made.

Limitations
First, the USP has not created a method for assessing half-tablet drug content uniformity; thus, previous studies assessing half-tablet drug content uniformity have used adapted USP methods for assessing weight variability as a means of estimating drug content uniformity. Second, all of the medications in this study are now available generically, and there is little financial value in splitting these particular drugs today. Third, the medications chosen for analysis were determined by prevalence of tablet splitting within a single health care network. The medications studied may not be representative of the most commonly split medications, and the purpose of the present research is not to suggest which drug classes may or may not be split. For these reasons, health care practitioners may not extrapolate the findings of this study to medications not studied. Fourth, the only tablet-splitting technique studied was the use of a tablet-splitting device. Splitting by hand or with sharp instruments including knives and razor blades are also commonly used techniques.
Analysis of Drug Content and Weight Uniformity for Half-Tablets of 6 Commonly Split Medications

DISCLOSURES

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Hill was primarily responsible for the study concept and design, with assistance from Karlage and Myrdal. The data were collected primarily by Hill and Varker, with assistance from Karlage. Hill and Myrdal interpreted the data, with assistance of the other 2 authors. Hill and Varker wrote the manuscript, and Varker was primarily responsible for revision of the manuscript, with assistance from the other 3 authors.

REFERENCES