The practice and clinical implications of tablet splitting in international health

Ivo Elliott1,2, Mayfong Mayxay1,2,3, Sengchanh Yeuchaixong1, Sue J. Lee2,4 and Paul N. Newton1,2

1 Lao-Oxford-Mahosot Hospital-Wellcome Trust Research Unit, Microbiology Laboratory, Mahosot Hospital, Vientiane, Lao PDR
2 Centre for Tropical Medicine, Nuffield Department of Medicine, Churchill Hospital, University of Oxford, Oxford, UK
3 Faculty of Postgraduate Studies, University of Health Sciences, Vientiane, Lao PDR
4 Mahidol-Oxford Tropical Medicine Research Unit, Faculty of Tropical Medicine, Mahidol University, Bangkok, Thailand

Abstract

OBJECTIVE Tablet splitting is frequently performed to facilitate correct dosing, but the practice and implications in low-income settings have rarely been discussed.

METHODS We selected eight drugs, with narrow therapeutic indices or critical dosages, frequently divided in the Lao PDR (Laos). These were split, by common techniques used in Laos, by four nurses and four laypersons. The mean percentage deviation from the theoretical expected weight and weight loss of divided tablets/capsules were recorded.

RESULTS Five of eight study drugs failed, on splitting, to meet European Pharmacopoeia recommendations for tablet weight deviation from the expected weight of tablet/capsule halves with 10% deviating by more than 25%. There was a significant difference in splitting accuracy between nurses and laypersons ($P=0.027$). Coated and unscored tablets were less accurately split than uncoated ($P=0.03$ and 0.0019 for each half) and scored (0.0001 for both halves) tablets.

CONCLUSION These findings have potential clinical implications on treatment outcome and the development of antimicrobial resistance. Investment by drug companies in a wider range of dosage units, particularly for narrow therapeutic index and critical dosage medicines, is strongly recommended.

keywords tablet splitting, therapeutic range, Laos, dose, essential medicines

Introduction

Tablet splitting is widely practised in many areas of healthcare. In the German primary care setting in 2006, an estimated one quarter of all drugs were split and in a large elderly care home in Canada 35% of all tablets were split (Fischbach et al. 2001; Quinzler et al. 2006). In low-income settings, such as the Lao PDR (Laos), tablet splitting is commonly performed, but the practice and implications have rarely been discussed.

The primary reason for tablet splitting is to increase dose flexibility, particularly for the elderly, children and those requiring titrating or tapering doses (Fischbach et al. 2001; Cohen & Cohen 2002; van Santen et al. 2002). Appropriate doses are often not manufactured or are unavailable. Drug costs per unit of active pharmaceutical ingredient (API) frequently decrease with increasing dose or flat charges may exist, independently of dose. Tablet splitting can therefore have an economic incentive, benefitting both the individual patient and the healthcare provider. Estimates of cost saving for splitting statins from innovative pharmaceutical companies are as high as 40–50% (Fawell et al. 1999; Duncan et al. 2002; Gee et al. 2002). This cost saving is, however, limited to relatively few drugs. Finally, a more pragmatic reason for tablet splitting is to aid swallowing.

Splitting tends to be performed by a variety of people including pharmacists, nurses and patients or their relatives. Tablets, particularly those with score lines, are usually split by hand. Those without score lines may require the use of a razor, knife or scissors. Commercially available splitting devices can also be employed where available (e.g. www.medimax.co.uk). Occasionally, capsules are split by emptying the powder and dividing it into equal portions.

The accuracy of tablet splitting has been demonstrated to vary considerably (Rosenberg et al. 2002; Teng et al. 2002; Polli et al. 2003; Verrue et al. 2011). Quality standards for uniformity of weight and content of manufactured drugs are outlined in the British (BP), United States (USP) and European Pharmacopoeias (EP). These require both the weight and content of the API of whole tablets to be within 85–115% of the intended dose with a relative standard deviation (RSD) less than or equal to 6%.
(United States Pharmacopeial Convention 1999). BP and USP standards apply only to whole dosage units with no guidance on split tablets. The revised 2008 EP standards for the division of scored tablets allow for no more than 1 in a set of 30 tablets to be outside the 85–115% range (European Pharmacopoeia Supplement 2008). If one tablet falls outside this range, it must fall within 75–125% of the expected mass/content. Although a homogeneous distribution of active drug is generally assumed in unscored tablets, some variation is expected and US Food and Drug Administration (FDA) bioequivalence standards permit variance of ±20%. The FDA, American Medical Society and American Pharmacists Association advise against splitting modified or sustained release, co-formulated, unscored, film-coated, friable or dose-critical tablets (American Pharmacists Association 2003).

The accuracy of tablet splitting is influenced by tablet size, shape, hardness, splitting method and human ability. Small, round or unusual-shaped tablets give rise to the greatest deviations and harder tablets are most likely to fragment or powder, leading to drug loss (McDevitt et al. 1998; Polli et al. 2003). Dividing tablets into quarters results in even greater ranges of weight differences (Biron et al. 1999; Kayumba 2006). Tablets with score lines, especially if deep, tend to split more uniformly (Gupta & Gupta 1988; Kayumba 2006). Hand-split tablets were less uniform than those split by razor, and knife splitting was less accurate than a tablet splitter (Teng et al. 2002; Cook et al. 2003). Weight loss due to fragmentation and powdering has been reported to vary between minimal (Stimpel et al. 1984; McDevitt et al. 1998) and 14% for tablet halves (Biron et al. 1999).

The implications of tablet splitting are wide ranging, but large deviations from the ideal weight after splitting are likely to be especially important (McDevitt et al. 1998; Rosenberg et al. 2002; Teng et al. 2002; Polli et al. 2003). This may result in incorrect dosing affecting clinical outcome, especially for medicines with narrow therapeutic indices. Under dosing may be an important factor in the development of resistance in diseases such as malaria (White et al. 2009).

We investigated the practice of tablet splitting in the low-income setting of Laos. We performed a study to replicate the usual circumstances and methods by which tablets are split and report the frequency with which split study drugs failed to meet EP guidelines on weight (dose) uniformity and identify factors that may influence the accuracy of tablet splitting.

**Methods**

**Participants**

Eight Lao volunteers were recruited to perform the tablet splitting at Mahosot Hospital, Vientiane. Four paediatric nurses at Mahosot Hospital and four laypersons (representing patients and relatives) with no healthcare-related experience were selected. These are the two major groups responsible for tablet splitting in Laos (pharmacists are less frequently involved).

**Study drugs**

Eight medicines frequently split in Laos, with narrow therapeutic indices or critical dosages, were selected (Table 1 and Table S1). Participants were requested to divide all drugs into halves with the exception of phenobarbitone, where one set was divided into halves and another into thirds, as is commonly required in pediatrics. For each study drug, 80 tablets (160 for phenobarbitone) were purchased from a local pharmacy, of the same dose, manufacturer and lot number per medicine. Doxycycline capsules are also frequently divided in Laos for paediatric dosing and were included in the eight study

| Drug          | Dose and formulation | Shape | Flat? | Score line? | Coated? | Split into? | Whole tablet weight (g) (SD) |
|---------------|----------------------|-------|-------|-------------|---------|------------|-------------------------------|
| Chloroquine   | 250 mg T             | Round | No    | No          | Yes     | 2          | 0.804 (0.019)                |
| Doxycycline   | 100 mg C             | Oblong| N/A   | N/A         | N/A     | 2          | 0.328 (0.011)                |
| Ofloxacin     | 200 mg T             | Oblong| Yes   | Yes         | No      | 2          | 0.466 (0.011)                |
| Enalapril     | 20 mg T              | Hexagonal| No | Yes         | No      | 2          | 0.201 (0.002)                |
| Atenolol      | 100 mg T             | Round | No    | No          | Yes     | 2          | 0.434 (0.005)                |
| Digoxin       | 0.25 mg T            | Round | Yes   | Yes         | No      | 2          | 0.155 (0.006)                |
| Glibenclamide | 5 mg T               | Oblong| Yes   | Yes         | No      | 2          | 0.160 (0.001)                |
| Phenobarbitone| 60 mg T              | Round | No    | No          | No      | 2 or 3     | 0.133 (0.001)                |

T, tablet; C, capsule; SD, standard deviation. Manufacturer and excipients are listed in supplementary material.
I. Elliott et al. Practice and implications of tablet splitting

drugs. All splitting was performed before drug expiry date.

Tablet splitting

Participants worked separately and were provided with 10 tablets or capsules of each study drug (total 90 tablets or capsules). They were asked to divide these, using their preferred method, into halves (both halves and thirds for phenobarbitone) and return the pieces or powder into separately labelled zipped plastic bags. Halves are referred to as A and B and thirds as A, B and C.

Weighing

Each individual whole tablet or capsule was weighed the day before splitting using a Sartorius BL210S (Göttingen, Germany) analytical balance. Weight was recorded to 0.001 g, and whole, half and tablet thirds were weighed using the same machine. Zipped bags containing doxycycline powder were weighed before and after washing out the powder with tap water (followed by thorough drying) as well as the capsule itself to accurately determine the weight of each half of powder. The same individual, blinded to the initial whole tablet weight, weighed the divided tablets/capsules. Interobserver variability was assessed using 50 randomly selected pairs of tablets and two observers.

Data analysis

Mean whole tablet weight [range and standard deviation (SD)] was calculated for each study drug. The median percentage deviation [interquartile range (IQR)] from the theoretical expected weight of split formulations and the maximum percentage deviation were calculated. A Wilcoxon matched-pairs signed-ranks test was used to compare the weight of each study drug half (or third) and its theoretical weight. Median weight loss, IQR, and maximum weight loss for each study drug were determined. The frequencies (%) of split study drugs falling outside the USP/EP recommended ranges of >85% and <115%, >75% and <125% and >6% relative SDs are reported.

The percentage deviation from the expected weight and the percentage weight loss were compared between coated and uncoated tablets, tablets with and without a scoreline and nurses and laypersons using a Mann–Whitney U test. Methods used to split tablets were analysed using the Kruskal–Wallis test. Fifty randomly selected pairs of half tablets from one randomly selected drug were re-weighed to check for interobserver variability

using the Bland–Altman method (Bland and Altman 1995).

Results

Eight drugs (with phenobarbitone used twice) were included in the study with a broad range of tablet characteristics of size, shape, coating and presence of a scoreline plus one powder-filled capsule (doxycycline) formulation. The percentage RSD of whole tablet weight ranged from 0.63% for glibenclamide to 2.63% for chloroquine, well within the recommended maximum of 6% (Table 1).

By Bland–Altman analysis for interobserver variability, the mean differences were normally distributed and no significant difference was seen between the first weights and re-weighing (mean difference 0.000185 (95% CI 0.000151–0.000217) g for half A and 0.000137 (0.000107–0.000167) g for half B.

A statistically significant difference (P < 0.01) was seen for median percentage deviation from theoretical for split drugs for four of nine sets of tablets/capsules (Table 2). Digoxin tablets (P = 0.0006) and the uncoated chloroquine (P = 0.01) were both inaccurately split. Phenobarbitone was the most inaccurately split (P < 0.0001) when divided into 3 (not when divided into 2). Dividing the powder content of doxycycline capsules also resulted in significant deviation from expected weight (P < 0.01). Median weight loss of tablets for each study drug ranged from 0.22% to 3.75% (Table 2). Maximum weight loss varied greatly from 1.85% (glibenclamide) to 23.48% (phenobarbital halves).

Table 3 shows the frequency of tablet halves (or thirds) falling outside the USP/EP recommendations. In total, 336 (25%) of the divided tablets were outside 85–115% of the theoretical weight and 140 (10%) deviated by more than 25% from the theoretical weight. Six of nine sets of tablets failed to meet the EP requirements following splitting, although one of these (glibenclamide) only failed on the basis of a single tablet divided inaccurately (27.5 and 30% deviation from theoretical for each half). When phenobarbitone was divided into 3, half the pieces deviated by >50% from the mean theoretical weight.

Comparing nurses with laypersons, splitting by nurses resulted in less deviation from theoretical weight of tablet halves (P = 0.0273), but not for weight loss. The greatest difference was seen for the division of doxycycline capsules (P = 0.0009). There was a significantly greater deviation from theoretical weight and weight loss for unscored (chloroquine and phenobarbitone) tablets than scored tablets (P = 0.0001 for both half weights and weight loss). A similar result was seen for coated (chloroquine and atenolol) versus uncoated tablets (P = 0.030
Wilcoxon matched-pairs signed-ranks test. All drugs were divided into halves, labelled half A and half B, or thirds, labelled A, B and C.

†expected weight (i.e. whole tablet weight divided by 2 or 3).

*Split into thirds.

Table 2 Median deviation from theoretical weight for each divided study drug and median and maximum weight loss

| Drug            | Median% deviation from theoretical of half A (IQR) | Median% deviation from theoretical of half B (IQR) | Median% weight loss (IQR) |
|-----------------|-----------------------------------------------|-----------------------------------------------|----------------------------|
| Chloroquine     | 6.63 (3.00–13.96)                            | 8.31 (3.80–15.66)                            | 48.97 (1.24–2.25)         |
| Doxycycline     | 11.76 (4.57–16.13)                           | 7.24 (3.87–15.17)                            | 43.97 (3.75–2.58)         |
| Ofloxacin       | 2.39 (1.03–4.88)                             | 2.27 (1.27–5.66)                            | 18.71 (0.22–0.43)         |
| Enalapril       | 5.03 (2.46–7.49)                             | 4.95 (2.00–8.37)                            | 20.20 (0.50–1.90)         |
| Atenolol        | 6.59 (2.41–11.88)                            | 7.30 (2.31–13.97)                            | 45.37 (1.14–1.45)         |
| Digoxin         | 7.19 (3.80–14.38)                            | 8.92 (3.59–16.03)                            | 56.69 (1.95–2.84)         |
| Glibenclamide   | 4.67 (2.51–4.97)                             | 4.38 (2.50–6.69)                            | 30.00 (0.62–0.63)         |
| Phenobarbitone  | 9.43 (3.77–19.47)                            | 9.85 (3.03–19.77)                            | 80.45 (1.50–2.79)         |
| Phenobarbitone† | 15.47 (8.27–23.16)                           | 0.001                                       | 100                        |
|                | 12.83 (5.32–27.84)                           | <0.0001                                    |                            |
|                | 16.54 (9.09–23.88)                           | <0.0001                                    |                            |

All drugs were divided into halves, labelled half A and half B, or thirds, labelled A, B and C.

*Wilcoxon matched-pairs signed-ranks test. P value = actual weight of divided halves (or thirds) compared with the theoretical expected weight (i.e. whole tablet weight divided by 2 or 3).

†Split into thirds.

Table 3 Frequencies of tablet halves or thirds deviating by more than 15% or 25% and study drugs with a >6% RSD

| Drug            | No. of halves (or thirds) deviating by >15% (%) | No. of halves (or thirds) deviating by >25% (%) | >6% RSD |
|-----------------|-----------------------------------------------|-----------------------------------------------|--------|
| Chloroquine     | 40/160 (25)                                  | 19/160 (12)                                  | Yes    |
| Doxycycline     | 44/160 (0)                                   | 18/160 (0)                                   | Yes    |
| Ofloxacin       | 4/160 (2.5)                                  | 0/160 (0)                                    | No     |
| Enalapril       | 7/160 (4.4)                                  | 0/160 (0)                                    | No     |
| Atenolol        | 29/160 (18)                                  | 10/160 (6.3)                                 | Yes    |
| Digoxin         | 39/160 (24)                                  | 19/160 (12)                                 | Yes    |
| Glibenclamide   | 1/160 (0.6)                                  | 1/160 (0.6)                                 | No     |
| Phenobarbitone  | 53/160 (33)                                  | 21/160 (13)                                 | Yes    |
| Phenobarbitone* | 119/240 (50)                                 | 52/240 (22)                                 | Yes    |
| Total           | 336 (25)                                      | 140 (10)                                    |        |

*Split into thirds.

and 0.0019 for each half). Weight loss was also significantly higher for coated tablets (P = 0.0001).

Significant differences were seen between the different methods used to divide the tablets or capsules for both weight deviation from theoretical weight and weight loss (all P ≤ 0.0001). The greatest differences were seen when dividing the powder in doxycycline capsules [median weight deviation of half A 11.74% (4.57–16.13) and half B 7.24% (3.87–15.17)], followed by knife [half A = 11.03% (2.38–22.02) and half B = 9.43% (3.76–21.80)] and scissors [half A = 6.92% (3.14–13.64) and half B = 7.58% (3.03–13.73)]. Dividing tablets by hand was most accurate [half A = 3.86% (1.88–6.80) and half B = 3.28% (1.69–6.25)].

Discussion

Table splitting is widely practised throughout the world, but has rarely been examined in low-/middle-income countries. In this study in Laos, six of the nine sets of split tablets/capsules failed to meet EP guidelines on weight uniformity of divided fragments, with 25% deviating by >15% and 10% by >25%. Previous studies report seven of 22 split drugs (Rosenberg et al. 2002), eight of 11 (Teng et al. 2002), four of 12 (Polli et al. 2003) and seven of eight (Verrue et al. 2011) failing to meet USP/EP guidelines. These findings are broadly similar to our own, despite the use of different tablets and methods of splitting.

Three study drugs, glibenclamide, ofloxacin and enalapril, were accurately divided and suffered the least weight loss (median <0.62%), and all participants reported these to be simple to divide either by hand or using scissors. All three formulations were scored and lacked a coating. Both these factors were significantly associated with more accurate tablet division. Atenolol and digoxin halves failed to meet EP guidelines. Digoxin had a statistically significant difference between weight of halves and theoretical weight; however, atenolol did not quite reach statistical significance (P = 0.0556). This was despite the presence of a scoreline in both tablets. Atenolol was coated, rounded and hard making it very difficult to split by hand and easy to fragment into multiple pieces when using knife or scissors. Digoxin tablets were very small, soft and crumbled easily when divided, reflecting a marked median weight loss of drug (1.95%). Phenobarbitone was a very small, round and hard tablet.
Division into two pieces resulted in failure to meet the EP guidelines, but as for atenolol, this did not reach statistical significance for the weights of the halves compared with the theoretical weight ($P = 0.109$). Division into three pieces proved the most difficult with very large differences between the weight of thirds and theoretical weight ($P < 0.0001$), median weight loss of 2.27% and as much as a 100% deviation from theoretical weight. Dividing the powder from doxycycline capsules in half also proved difficult and resulted in significant differences in weight and the greatest median weight loss (3.75%). There was a statistically significant difference in results seen between nurses and laypersons, with nurses performing more accurate division. We selected paediatric nurses working at Mahosot Hospital, who regularly divided drugs as part of their practice. This experience is likely to have resulted in greater accuracy of division.

Median tablet/capsule weight losses due to splitting in this study were 0.22–3.75%, consistent with the few other studies to report this (Gupta & Gupta 1988; Teng et al. 2002). In our study, a maximum weight loss of 23.5% for a phenobarbitone tablet was seen. One study reported a maximum of 27% weight loss when dividing oral anticoagulants into quarters (Biron et al. 1999).

The clinical impact of these often-large variations in weight between fragments has been investigated primarily for drugs with wide therapeutic indices. The impact of RSDs of more than 10% of halved statins had no clinical impact on patient LDL cholesterol or total cholesterol (Duncan et al. 2002; Gee et al. 2002), and similar variations of weight of lisinopril fragments did not impact blood pressure over a prolonged follow-up period (Rindone 2000). One study examined content uniformity of the narrow therapeutic index levothyroxine after splitting and suggested that sub- or super potency could prove clinically detrimental; however, no clinical studies were performed (Shah et al. 2010).

No specific recommendations were provided by the manufacturers of any of the study drugs, scored or unscored, specifically advising on whether tablet or capsule splitting could be performed. The uniformity of distribution of API in the study drugs was also not recorded. Scored tablets facilitate splitting, and the importance of manufacturing quality assurance standards for API distribution in whole tablets has recently been highlighted (Anonymous 2014). In the absence of explicit documentation by the manufacturer allowing splitting of unscored or coated tablets, we recommend education of healthcare staff and patients against dividing these tablets. Best practices guidelines for tablet splitting are available (US Food & Drug Administration). However, where alternative dosage units do not exist, manufacture of these is thus urgently needed.

Inadequate antimalarial treatment doses, particularly in patients with hyperparasitaemia, may be an important source of de novo resistance and recrudescence (Simpson et al. 2000; White et al. 2009). One study performed in Africa found that 13% of quinine sulphate tablets deviated in weight by more than 35% from the theoretical (Kayumba 2006). Our study demonstrated difficulty in accurately splitting chloroquine tablets, which may be an important factor in developing drug resistance. Subtherapeutic doxycycline dosing in severe malaria may also be a factor in emerging resistance for partner artemisinin drugs (Newton et al. 2005; Dondorp et al. 2009) and could result in resistance developing in bacteria such the rickettsiae for which doxycycline is frequently used in Laos and elsewhere in Asia. Tablet splitting is contraindicated for co-formulated drugs including the widely used amoxicillin/clavulanic acid (American Pharmacists Association 2003; Anonymous 2014). Although not investigated in this study, division of this antibiotic may also act as a driver for resistance.

Our findings are likely to underestimate the effect of tablet splitting. Divided tablets may fragment further when kept in a container after splitting, hygroscopic absorption from high humidity and the transfer of skin oils onto the tablets may have overestimated fragment weights. We were unable to determine the uniformity of distribution of API in whole tablets. Uneven distribution of API in divided tablets may lead to even greater risks for the emergence of antimicrobial resistance. We did not perform a direct comparison of splitting techniques to determine an optimal method, but division by hand appeared to be most accurate, followed by scissors or knife. Least accurate was the division of powdered drug from capsules. The accuracy of splitting by hand may reflect the choice available to participants, as tablets that were easy to divide, for example ofloxacin was divided so by hand, but the most difficult to split, for example atenolol, could only be divided using scissors or a knife. Verrue et al. (2011) showed that a splitting device was significantly more accurate than division with scissors or by hand; however, these devices are expensive and unavailable in low-income settings. An additional limitation to this study was that participants split a large number of tablets in a period of approximately 90 min. This does not reflect usual practice and may have influenced accuracy.

Further analysis should be performed to better understand the pharmacokinetic–pharmacodynamic consequences of tablet splitting for particular pathogens/disease states. That division of uncoated tablets with
scorelines resulted in the most accurate tablet division suggests that pharmaceutical manufacturers of medicines that are commonly split should consider such lines, if technically feasible, based on pharmaceutical specifications including the proof of uniform distribution of API. These data suggest that some key medicines such as doxycycline capsules, chloroquine and digoxin tablets should not be split and that phenobarbitone tablets should not be split, especially into thirds.

Conclusion

This study highlights the widespread practice and inaccuracy of tablet and capsule splitting of medicines. There is clear evidence that tablet design, with the lack of a coating and presence of a scoreline, allows significantly more accurate tablet splitting. The potential clinical implications of this are far reaching and may have a significant impact on successful outcome and the development of antimicrobial resistance. Vulnerable groups including paediatric and elderly patients are most at risk of the consequences of inaccurate tablet division. Investment by drug companies in the production of a wider range of dosage units or tablets better designed for splitting and better product information on the suitability of tablets for splitting, particularly for narrow therapeutic or critical dosage drugs, is strongly recommended.

Acknowledgements

We thank the Directors and staff of Mahosot Hospital, Laos, for their help. We thank Dr. Koukeo Phommasone for assistance with study logistics and Patricia Tabernero for comments on the manuscript. We are very grateful to the nurses and laypersons for their time in taking part in this study. The work was funded by the Wellcome Trust of Great Britain.

References

American Pharmacists Association (2003) Tablet splitting: evaluating appropriateness for patients. Journal of the American Pharmacists Association 2004, 324–325.
Anonymous (2014) Tablet splitting. JAMA 311, 521.
Biron C, Licznar P, Hansel S & Schved JF (1999) Oral anticoagulant drugs: do not cut tablets in quarters. Thrombosis and Haemostasis 82, 1201.
Bland JM & Altman DG (1995) Comparing methods of measurement: why plotting difference against standard method is misleading. Lancet 346, 1085–1087.
Cohen CI & Cohen SI (2002) Potential savings from splitting newer antidepressant medications. CNS Drugs 16, 353–358.
Cook TJ, Edwards S, Gyemah C, Shah M, Shah I & Fox T (2003) Variability in tablet fragment weights when splitting unscored cyclobenzaprine 10 mg tablets. Journal of the American Pharmacists Association 2004, 583–586.
Dondorp AM, Nosten F, Yi P et al. (2009) Artemisinin resistance in Plasmodium falciparum malaria. New England Journal of Medicine 361, 455–467.
Duncan MC, Castle SS & Streetman DS (2002) Effect of tablet splitting on serum cholesterol concentrations. Annals of Pharmacotherapy 36, 205–209.
Fawell NG, Cookson TL & Scranton SS (1999) Relationship between tablet splitting and compliance, drug acquisition cost, and patient acceptance. American Journal of Health System Pharmacy 56, 2542–2545.
Fischbach MS, Gold JL, Lee M, Dergal JM, Littner GM & Rochon PA (2001) Pill-splitting in a long-term care facility. CMAJ 164, 785–786.
Gee M, Hasson NK, Hahn T & Ryono R (2002) Effects of a tablet-splitting program in patients taking HMG-CoA reductase inhibitors: analysis of clinical effects, patient satisfaction, compliance, and cost avoidance. Journal of Managed Care Pharmacy 8, 453–458.
Gupta P & Gupta K (1988) Broken tablets: does the sum of the parts equal the whole? American Journal of Hospital Pharmacy 45, 1498.
Kayumba PC (2006) Impact du cassage de comprimes de sulfate de quinine sur l’exactitude de la dose administrée aux enfants. Le pharmacien d’Afrique 193, 11–16.
McDevitt JT, Gurst AH & Chen Y (1998) Accuracy of tablet splitting. Pharmacotherapy 18, 193–197.
Newton PN, Chaulet JF, Brockman A et al. (2005) Pharmacokinetics of oral doxycycline during combination treatment of severe falciparum malaria. Antimicrobial Agents and Chemotherapy 49, 1622–1625.
Polli JE, Kim S & Martin BR (2003) Weight uniformity of split tablets required by a Veterans Affairs policy. Journal of Managed Care Pharmacy 9, 401–407.
Quinzler R, Gasse C, Schneider A, Kaufmann-Kolle P, Szecsenyi J & Haeefeli WE (2006) The frequency of inappropriate tablet splitting in primary care. European Journal of Clinical Pharmacology 62, 1065–1073.
Rindone JP (2000) Evaluation of tablet-splitting in patients taking lisinopril for hypertension. Journal of Clinical Outcomes Management 7, 22–24.
Rosenberg JM, Nathan JP & Plakogiannis F (2002) Weight variability of pharmacist-dispensed split tablets. Journal of the American Pharmaceutical Association 42, 200–205.
van Santen E, Barends DM & Frijlink HW (2002) Breaking of scored tablets: a review. European Journal of Pharmaceutics and Biopharmaceutics 53, 139–143.
Shah RB, Collier JS, Sayeed VA, Bryant A, Habib MJ & Khan MA (2010) Tablet splitting of a narrow therapeutic index drug: a case with levethoxyxine sodium. American Association of Pharmaceutical Scientists 11, 1359–1367.
Simpson JA, Watkins ER, Price RN, Aarons L, Kyle DE & White NJ (2000) Mefloquine pharmacokinetic-pharmacody-
Practice and implications of tablet splitting

Veree C, Mehuys E, Boussery K, Remon JP & Petrovic M (2011) Tablet-splitting: a common yet not so innocent practice. *Journal of Advanced Nursing* 67, 26–32.

White NJ, Pongtavornpinyo W, Maude RJ *et al.* (2009a) Hyperparasitaemia and low dosing are an important source of antimalarial drug resistance. *Malaria Journal* 8, 253.

White NJ, Pongtavornpinyo W, Maude RJ *et al.* (2009b) Hyperparasitaemia and low dosing are an important source of antimalarial drug resistance. *Malaria Journal* 8, 253.

Supporting Information

Additional Supporting Information may be found in the online version of this article:

Table S1. Study drug tablet/capsule excipients.

I. Elliott *et al.* Practice and implications of tablet splitting

dynamic models: implications for dosing and resistance. *Antimicrobial Agents and Chemotherapy* 44, 3414–3424.

Stimpel M, Kuffer B, Groth H & Vetter W (1984) Breaking tablets in half. *Lancet* 1, 1299.

European Pharmacopoeia Supplement (2008) Tablets. Monograph 0478 (Suppl. 64).

Teng J, Song CK, Williams RL & Polli JE (2002) Lack of medication dose uniformity in commonly split tablets. *Journal of the American Pharmaceutical Association* 42, 195–199.

United States Pharmacopeial Convention (1999) Uniformity of Dosage Units. *United States Pharmacopeia* 24/National *Formulary* 19: Rockville, MD, United States Pharmacopeial Convention, pp. 2000–2003.

US Food and Drug Administration Best Practices for Tablet Splitting. www.fda.gov/Drugs/ResourcesForYou/Consumers/BuyingUsingmedicineSafely/EnsuringSafeUseofmedicine/ucm184666htm.