Grapefruit juice and clopidogrel

Bailey and colleagues⁠¹ are to be commended for highlighting the potential of many medications to interact with grapefruit.¹ However, one medication on their list (Table 1¹) deserves closer scrutiny.

At first glance, clopidogrel seems to fit the criteria of a medication that is a known substrate of CYP3A4 with low total bioavailability, but on closer examination the issue is less clear.

First, clopidogrel is a prodrug that needs to be metabolized to its active form.² Therefore, it is the only medication listed in Table 1¹ whose interaction with grapefruit via CYP3A4 may lead to a loss of efficacy instead of toxicity (this was correctly stated in Table 1¹). However, the biotransformation of clopidogrel to its active form is a two-step process involving multiple CYP isoenzymes. Only in the second step in the process is CYP3A4 involved.³ Whether intestinal CYP3A4 (as opposed to hepatic CYP3A4) is even involved in this second step is unclear.

Second, the enzyme that is primarily responsible for the low bioavailability of clopidogrel is not a cytochrome P450 enzyme at all. Instead, 85% of clopidogrel is metabolized to an inactive compound through esterase enzymes.² Interestingly, some suggest that grapefruit juice may inhibit intestinal esterase enzymes.⁴ If this were the case, grapefruit may increase the amount of parent drug available to be converted to its active form, which could lead to increased antiplatelet effect. Therefore, given the complex metabolic pathway of clopidogrel, what, if any, effect grapefruit would have on pharmacokinetics or pharmacodynamics of this agent is not clear. A randomized study examining the effect of grapefruit juice on the antiplatelet activity of clopidogrel will hopefully shed more light on this issue (ClinicalTrials.gov: NCT00817999).

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References


The author responds

Friesen¹ has provided thoughtful comments about the “loss of efficacy” of clopidogrel from grapefruit interactions that was reported in our article.² Because clopidogrel is commonly prescribed in Canada and is used to prevent common and serious thrombotic-mediated cardiovascular events, reduction in the effectiveness of clopidogrel has potentially broad and significant clinical consequences.³

Friesen correctly notes that clopidogrel requires metabolic conversion before it can exert an antiplatelet effect and that this entails multiple sequential bioactivation steps involving cytochrome P450 enzymes (CYPs). One of these CYPs is CYP3A4, which grapefruit is known to inactivate. Because erythromycin and troleandomycin (CYP3A4 inhibitors) decrease and rifampicin (CYP3A4 inducer) increases the clinical antiplatelet activity of clopidogrel, one could conclude that this enzyme plays a key role in the therapeutics of this drug.⁴ These drugs most likely modulate the activity of CYP3A4 in both the small intestine and liver. Friesen¹ rightly remarks that grapefruit primarily inactivates only intestinal CYP3A4. Moreover, he also puts forth the possibility that the low oral bioavailability of clopidogrel may not be from the action of CYP3A4, which is a fundamental characteristic for drugs that interact with grapefruit.

Friesen¹ mainly contends that whether grapefruit can attenuate the antiplatelet action of clopidogrel clinically is not currently clear. He cites the study “The Impact of Grapefruit Juice on the Response to Clopidogrel” found on ClinicalTrials.gov.¹ Friesen¹ believes that results from this study will be instrumental in deciding the relevance of this interaction.

The results of the completed ClinicalTrials.gov trial revealed that in the loading-dose phase, clopidogrel 300 mg with grapefruit juice versus water produced platelet inhibition of 23% compared with 41%, respectively. In the maintenance phase, clopidogrel 75 mg with grapefruit juice versus water for 7 days resulted in inhibition of 25% compared with 59%, respectively. Thus, our concern regarding “loss of efficacy” of clopidogrel from grapefruit interactions appears currently justified.

This information is not mentioned in the prescribing information for clopidogrel possibly because the previous version, dated April 2012, of the prescribing information may have preceded completion of the trial in August 2012.

Of greater concern is that there is no indication of the effects of inhibition and induction of CYP3A4 caused by drugs or herbals — particularly St. John’s wort — which have been known for several years.⁵

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References

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