Pharmacokinetic Interactions of Antiepileptic Drugs

Penny S. Albright and J. Bruni

ABSTRACT: The problem of antiepileptic drug interactions is significant in that many epileptic patients are treated with multiple drug therapy. Moreover, patients may also be receiving additional medication for other concurrent disorders. Most drug interactions are pharmacokinetic, involving changes in absorption, protein binding, metabolism, or excretion. As a result, plasma levels of the antiepileptic drug may decrease leading to exacerbation of seizures. Alternatively, plasma levels may rise resulting in toxic side effects. Similar changes may also occur with drugs given for other disorders. In this paper, possible mechanisms of drug interactions are discussed. This is followed by a description of clinically significant interactions involving phenytoin, carbamazepine, barbiturates, valproic acid, benzodiazepines, and succinimides. Potentially serious drug interactions may be minimized by using as few medications as possible and by regularly monitoring plasma levels of antiepileptic drugs.

RÉSUMÉ: Le problème des interactions entre les drogues anticonvulsivantes est important car beaucoup des patients épileptiques reçoivent le traitement des plusieurs médicaments. D'ailleurs quelques patients prennent aussi des drogues supplémentaires pour autres maladies. La plupart des interactions sont pharmacokinetiques, associées à des changements de l'absorption, de la fixation à des protéines, du métabolisme, ou de l'excrétion. En conséquence les niveaux plasmatiques de la drogue antépileptique peuvent diminuer et exacerber des convulsions. Au contraire, les teneurs plasmatiques peuvent augmenter avec le résultat de la toxicité. Des changements semblables pourraient avoir lieu avec des drogues ordonnées pour autres maladies. Dans cet article, les mécanismes possibles des interactions des drogues se discutent. Cela se suit de la description des interactions importantes cliniques en ce qui concerne la phénytoïne, la carbamazépine, les barbiturates, l'acide valproique, les benzodiazépines, et les succinimides. Les interactions entre les drogues les plus potentiellement graves peuvent être minimisées en employant le plus petit nombre des médicaments possible et en surveillant périodiquement les teneurs plasmatiques des drogues anticonvulsivantes.
Changes in Drug Metabolism

Most antiepileptic drugs undergo metabolism by enzymes located in the smooth endoplasmic reticulum of the liver. Metabolism consists of two phases which may occur separately or in series: phase 1 which involves oxidation-reduction reactions, and phase 2 which involves conjugation with glucuronide, sulphate, or acetic acid. This process converts the drugs into more polar, water-soluble metabolites which can then be easily eliminated by the kidney. The activity of hepatic drug metabolizing enzymes can be greatly affected by the administration of various therapeutic agents. Certain agents are known to increase or induce hepatic enzymes leading to increased biotransformation of various compounds. This effect would result in reduced efficacy except when active metabolites are produced. In that case, therapeutic effects may be enhanced depending on the relative activity of the metabolite and the parent drug.

Inhibition of drug metabolism is also known to occur when two compounds are co-administered. Inhibition may be competitive in that both drugs are competing for the same enzyme or it may be non-competitive in that one drug inactivates the enzyme responsible for metabolism of the other. Non-competitive inhibition is likely to cause a continuous increase in plasma levels of the primary drug whereas competitive inhibition would lead to stabilization at a new plateau (Kutt, 1975).

Changes in Protein Binding

If two highly protein-bound drugs are administered together, one drug may be displaced with a resultant increase in free levels. The ultimate effect of this change depends on whether elimination of the drug in question is restrictive or non-restrictive.

Restrictive elimination means that the extraction of the drug is less than the free fraction in blood. Displacement of such a drug from protein binding sites would initially lead to an elevation in free fraction with an increase in therapeutic effect or toxicity. However, this effect would be transient, since the increased free levels would lead to increased elimination. Free drug concentration would then return to the previous level although total plasma levels would be considerably reduced (Shand et al., 1975). Changes in total plasma levels can therefore be expected when two drugs are given which compete for acidic or basic compounds. Co-administration of two acidic drugs may retard tubular secretion of one of them. Secreted basic drugs may interact in a similar manner.

<table>
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<tr>
<th>Table 1: Mechanisms of Pharmacokinetic Drug Interactions</th>
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<tr>
<td>1. Absorption</td>
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<td>2. Protein Binding</td>
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<th>Table 2: Drug Interactions Involving Phenytoin</th>
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<td>Drugs Increasing Phenytoin Levels</td>
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<tr>
<td>Disulfiram</td>
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<tr>
<td>Sulthiame</td>
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<td>Isoniazid</td>
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<td>Chloramphenicol</td>
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<td>Bishydroxyxoumarin</td>
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<td>Propoxyphene</td>
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<td>Methsuximide</td>
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| Drugs Decreasing Phenytoin Levels               |
| Ethanol                                        |
| Carbamazepine                                  |
| Valproic Acid (transiently)                    |
| Antacids                                       |
| Phenobarbital                                  |

| Other Drugs Affected by Phenytoin              |
| (Plasma Levels are decreased)                  |
| Oral Contraceptives                            |
| Bishydroxyxoumarin                             |
| Dexamethasone                                  |
| Metapyrone                                     |
| Quinidine                                      |

Changes in Renal Excretion

Drug interactions at the renal level typically involve changes in tubular secretion or reabsorption. Theoretically they also may involve changes in glomerular filtration and renal blood flow although these are rarely observed (Shand et al., 1975). Changes in tubular reabsorption can be caused by alterations in urinary pH. For example, weak acids are eliminated more rapidly in alkaline urine whereas weak bases are excreted more effectively when the pH is low. Tubular secretion can be affected when two drugs are given which compete for acidic or basic compounds. Co-administration of two acidic drugs may retard tubular secretion of one of them. Secreted basic drugs may interact in a similar manner.

Antiepileptic Drug Interactions

Phenytoin

Drugs elevating plasma phenytoin levels. A number of drugs are reported to cause increases in plasma phenytoin levels (Table 2). With respect to other antiepileptic drugs, only a few are thought to raise phenytoin concentrations. For example, the metabolite of methsuximide competes with phenytoin for metabolic enzymes and, when both are administered, phenytoin toxicity may occur (Rambeck, 1979). Competitive inhibition between phenytoin and phenobarbital also has been reported (Patsalos and Lascelles, 1977). However, the ability of phenobarbital to induce the enzymes involved in phenytoin biotransformation can counteract this effect.

With respect to other types of medications, disulfiram (Olesen, 1967) and sulthiame (Houghton and Richens, 1974) both inhibit phenytoin metabolism leading to increases in plasma concentrations. Isoniazid, an antituberculosis agent, increases phenytoin levels in about 10% of epileptic patients (Kutt et al., 1966). Slow acetylators are more likely to exhibit this effect (Brennan et al., 1970). Although chloramphenicol can increase plasma phenytoin levels, 2) chelation of drugs with metals or ions; and 3) stimulation or inhibition of gut enzymes. Changes in gastric motility may affect the rate of absorption but rarely alter the total amount of drug absorbed.

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Drugs subject to non-restrictive elimination are highly extracted by the liver and the extraction ratio therefore exceeds the free fraction of the drug. Displacement of such drugs from protein binding sites results in an increase in volume of distribution and half-life (Shand et al., 1975). Free drug levels may rise since little increase in elimination would occur. In fact, however, protein binding interactions involving these drugs are seldom clinically important (Shand et al., 1975).
levels (Koup et al., 1978), antimicrobial agents are rarely given for long enough periods to induce severe clinical complications. Bishydroxycoumarin (Hansen et al., 1966) and propoxyphene (Dam et al., 1980) also are reported to elevate phenytoin concentrations.

**Drugs decreasing phenytoin levels.** Phenytoin concentration in plasma is decreased by the drugs shown in Table 2. The antiepileptic drugs, phenobarbital and carbamazepine, can induce phenytoin metabolism leading to reduced levels (Kutt, 1975). Valproic acid causes a transient reduction of total phenytoin levels in the majority of patients (Bruni et al., 1979). This interaction is caused by valproic acid displacing phenytoin from protein binding sites, leading to elevated free phenytoin levels. The increase in free concentration results in a compensatory increase in metabolism, and total phenytoin concentration declines. After a number of weeks, valproic acid inhibits phenytoin metabolism and phenytoin levels rise again. (Bruni et al., 1980a).

Of the non-antiepileptic drugs, chronic ethanol consumption can reduce phenytoin levels by inducing metabolic enzymes (Kater et al., 1969). However, drug metabolism may be inhibited during acute ethanol intake (Kutt, 1975). Antacids have been reported to reduce absorption of phenytoin from the G.I. tract (Gdnett et al., 1979).

**Effects of phenytoin on other drugs**

Table 2 shows those drugs which are affected by co-administration of phenytoin. One of the most significant interactions is the increased metabolism of contraceptive steroids leading to contraceptive failure in women taking phenytoin (Hempel and Klinger, 1976). Other drugs which are reduced by phenytoin administration include bishydroxycoumarin (Hansen et al., 1971), dexamethasone (McClelland and Jack, 1978), metapyrone (Meikle et al., 1969) and quinidine (Data et al., 1976).

**Carbamazepine**

**Drugs elevating carbamazepine levels.** Administration of propoxyphene to patients taking carbamazepine produces increased plasma levels and signs of carbamazepine toxicity (Levy and Pitlick, 1982).

**Drugs decreasing carbamazepine levels.** With chronic use, carbamazepine induces its own metabolism (autoinduction) with a resultant decrease in half-life (Eichelbaum et al., 1976). Carbamazepine metabolism is also highly inducible by other antiepileptic agents. For example, carbamazepine levels are reduced by concomitant therapy with phenytoin, phenobarbital (Christiansen and Däm, 1973), and primidone (Schneider, 1975). This phenomenon does not necessarily reduce therapeutic efficacy since the concentration of carbamazepine epoxide, the active metabolite, is higher in these patients (Levy and Pitlick, 1982).

**Effects of carbamazepine on other drugs.** Carbamazepine's powerful enzyme-inducing properties can also affect the metabolism of other drugs. Carbamazepine has been shown to increase the clearance of ethosuximide (Warren et al., 1980), clonazepam (Lai et al., 1978), and valproic acid (Bowdle et al., 1979).

**Barbiturates (phenobarbital and primidone)**

**Drugs elevating barbiturate levels.** Table 3 summarizes drug interactions involving barbiturates. One of the most clinically significant interactions is observed when phenobarbital and valproic acid are co-administered. When valproic acid is initially given to patients receiving phenobarbital, signs of somnolence or even coma may appear within days or weeks. These symptoms are accompanied by an increase in plasma phenobarbital levels (Bruni et al., 1980b). The mechanism of this effect is thought to involve an inhibition of phenobarbital metabolism as indicated by a reduction in the excretion of metabolite (Bruni et al., 1980b; Kapetanovic et al., 1981). However, since a large proportion of phenobarbital is excreted unchanged, additional factors are thought to contribute to phenobarbital accumulation (Kutt and Paris-Kutt, 1982).

It is of interest that when primidone and valproic acid are combined phenobarbital levels do not increase as predictably (Flachs et al., 1977). It has been speculated that valproic acid also inhibits the conversion of primidone to phenobarbital (Bruni, 1981; Windorfer and Sauer, 1977). Co-administration of phenytoin may also elevate phenobarbital levels, although this is an infrequent occurrence. A competition for metabolic enzymes is thought to be responsible for this effect (Kutt and Paris-Kutt, 1982). Phenytoin is also thought to promote the conversion of primidone to phenobarbital (Fincham et al., 1974) although the clinical expression of this effect is questionable. Phenobarbital and primidone should not be given together since primidone is metabolized in part to phenobarbital. Most other antiepileptic drugs do not alter plasma phenobarbital levels.

**Drugs decreasing barbiturate levels.** Ammonium chloride can accelerate the elimination of phenobarbital and reduce plasma levels. Alkalization of the urine by this or other agents increases renal excretion of the drug (Kutt and Paris-Kutt, 1928).

**Effects of barbiturates on other drugs.** Phenobarbital decreases the absorption of phenytoin and griseofulvin. Because of its powerful enzyme-inducing properties, phenobarbital may decrease phenytoin levels in some patients. However, it also competes with phenytoin as a substrate for parahydroxylation and glucuronidation. The net effect of these opposing changes is little or no change in phenytoin levels (Kutt et al., 1969). Phenobarbital also may cause a reduction in carbamazepine levels although this effect also is variable (Dam et al., 1973).

With respect to the nonantiepileptic drugs, phenobarbital can reduce levels of bishydroxycoumarin and warfarin (McDonald and Robinson, 1968; Cucinelli et al., 1965). Induction of metabolism and possibly decreased absorption can account for these findings. Careful monitoring of prothrombin times should avoid serious consequences of this interaction. Enzyme induction can also
lead to decreased levels of chlordopromazine (Loga et al., 1975), desipramine (Kutt and Paris-Kutt, 1982), and oral contraceptives (Hempel and Klinger, 1976). The elimination of endogenous substances such as bilirubin (Thompson et al., 1969), cholesterol, bile salts, lipids (Linares et al., 1973), endogenous steroids (Burstein and Klaiber, 1965), and vitamin D (Hunter, 1976) is accelerated by phenobarbital administration.

**Valproic Acid**

*Drugs elevating valproic acid levels.* The only factor known to increase valproic acid levels is the discontinuation of enzyme-inducing antiepileptic drugs.

*Drugs decreasing valproic acid levels.* Concurrent use of other antiepileptic drugs can induce metabolic enzymes and reduce valproic acid concentrations (Reunanen et al., 1980). Removal of concomitant therapy may then cause a rebound increase in valproic acid levels (Johannessen and Henrikson, 1980). Administration of salicylate displaces valproic acid from protein binding sites (Fleitman et al., 1980) and leads to increased clearance and lower plasma concentrations (Schoeben et al., 1978, Viswanathan and Levy, 1981). Endogenous fatty acids can also elevate the free fraction of valproic acid (Zimmerman et al., 1981).

*Effects of valproic acid on other drugs.* The interactions between valproic acid and phenytoin, phenobarbital, and primidone have been described in previous sections. Valproic acid may also lead to increased levels of carbamazepine (Mattson et al., 1982), and ethosuximide (Mattson and Cramer, 1980). In addition, the combined use of valproic acid and clonazepam may produce signs of toxicity and even absence status (Jeavons and Clark, 1974). The mechanism of this effect is unknown. Table 4 summarizes the major interactions involving valproic acid.

**Benzodiazepines**

*Drugs elevating benzodiazepine levels.* The clearance of diazepam and its active metabolite, N-desmethyldiazepam is reduced by disulfiram (MacLeod et al., 1978) and cimetidine (Klotz et al., 1979).

*Drugs decreasing benzodiazepine levels.* The addition of other antiepileptic drugs may lower levels of clonazepam presumably by the induction of metabolic enzymes. Specific effects have been noted for carbamazepine (Lai et al., 1978), phenobarbital (Nanda et al., 1977), and phenytoin (Sjö et al., 1975).

*Effects of benzodiazepines on other drugs.* Pharmacodynamic interactions can occur between benzodiazepines and ethanol, barbiturates, antihistamines, phenothiazines, and tricyclic antidepressants. Other than these, the benzodiazepines do not produce prominent interactions with other drugs probably because they have little capacity to induce hepatic enzymes. Clonazepam has little effect on phenytoin and carbamazepine although diminished phenobarbital levels have been reported (Benetello et al., 1977).

**Succinimides**

*Drugs elevating succinimide levels.* No significant interactions seem to produce increased levels of the succinimides.

*Drugs decreasing succinimide levels.* Reduced levels of ethosuximide after addition of carbamazepine are probably due to enzyme induction (Warren et al., 1980).

**Effects of succinimides on other drugs.** The simultaneous administration of methsuximide with phenobarbital or phenytoin results in increased levels of the latter drugs. This effect is presumed to result from competitive inhibition of metabolic enzymes (Rambeck, 1979). Some increases in plasma phenytoin concentration occur after the addition of ethosuximide (Lander et al., 1975).

**CONCLUSIONS**

Important drug interactions may occur when two or more antiepileptic drugs are administered concurrently or if an antiepileptic drug is given together with another medication. Most drug interactions involve pharmacokinetic rather than pharmacodynamic changes. Such interactions can pose clinically important problems for the patient. In some cases the therapeutic efficacy of the antiepileptic medication is reduced leading to increased seizure frequency or, on the other hand, plasma concentrations may increase leading to toxic side effects. Similar effects have been observed with medications given for other disorders. Many potentially serious interactions can be avoided by careful monitoring of plasma levels of antiepileptic drugs. Although total concentrations are sufficiently informative for most purposes, free levels should be assessed in some situations (i.e. phenytoin-valproate interaction). However, maintenance of the epileptic patient on a single antiepileptic drug is the most effective way to avoid unwanted drug interactions.

**REFERENCES**


