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Porphyrin-inducing Activity of Alfaxolone and Alfadolone Acetate in Chick Embryo Liver Cells

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The two steroid components of Alfashein®, alfaxolone and alfadolone acetate, have been tested for porphyrin-inducing activity in chick embryo liver cell culture and for hepatic ALA-synthetase-inducing activity in the 17-day-old chick embryo. In cell culture alfaxolone was shown to have potency comparable to that of thiopental, while alfadolone acetate had low potency.

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In the 17-day-old chick embryo alfaxolone has a third the potency of thiopental; alfadolone acetate showed low potency. The authors conclude that an induction dose of Alfashein® would be less likely than a comparable dose of thiopental to increase ALA-synthetase activity in a patient with hereditary hepatic porphyria. (Key words: Anesthetics, intravenous: steriod; thiopental; porphyria.)

The need to choose a suitable drug for induction of anesthesia for patients who have hereditary hepatic porphyria sometimes confronts the anesthetist. It was thus of interest to study the porphyrin-inducing activity of the new steroidal anesthetic Alfashein® in two well-known screening procedures, the chick embryo liver

![Graph showing porphyrin accumulation in cells and medium 24 hours after administration of increasing concentrations of A1A, alfadoxolone, and alfadolone acetate. The points represent the means of at least four determinations ± standard error.](http://anesthesiology.pubs.asahq.org/pdfaccess.ashx?url=/data/journals/jasa/931484/)
cell culture⁴ and the 17-day-old chick embryo.³ Alfathein consists of two steroids, alfaxyolone and alfaxyolone acetate, in a 3:1 ratio, dissolved in polyoxyethylated castor oil (Cremophor EL⁵).² Alfaxyolone is the principal anesthetic agent, and alfaxyolone acetate, which has half the anesthetic potency of alfaxyolone, is added to enhance the solubility of alfaxyolone. Each steroid component was tested separately.

**Methods**

The method used for investigating drug-induced porphyrin accumulation in chick embryo liver cells⁵ is a modification of the procedure of Granick.⁵ The steroids (1.5–100 µg) were dissolved in 10 µl of 95 per cent ethanol for addition to the cell culture medium (5 ml). Porphyrin accumulation was determined 24 hr after the addition of steroids by the procedure of Granick,² and was expressed as ng porphyrin formed per mg cellular protein. The procedure used for determining hepatic δ-aminolevulinic acid (ALA) synthetase activity in 17-day-old chick embryos was that of Racz and
Steroids (0.3–7 mg) were dissolved in 0.1 ml of dimethylsulfoxide, and injected through the chorioallantoic fluid surrounding the embryo; hepatic ALA-synthetase activity was determined six hours later. This activity was expressed as nmol ALA formed per hour per 100 mg liver protein. Liver and cellular proteins were determined by the method of Lowry et al. Neither the 95 per cent ethanol nor the dimethylsulfoxide had any effect on porphyrin biosynthesis or on ALA-synthetase activity.

Results and Discussion

Alfadalone acetate has low potency as a porphyrin-inducing agent in chick embryo liver cells when compared with a standard porphyrin-inducing chemical, allylisopropylacetamide (AlA), while alfadalone has a potency comparable to that of AIA (fig. 1). The hepatic ALA-synthetase-inducing activities of alfadalone, alfadalone acetate, and AIA in the 17-day-old chick embryo differ (fig. 2). Alfadalone acetate has very low potency, while alfadalone has a potency intermediate between those of AIA and alfadalone acetate. Parikh and Moore injected Alfathesin, 12 mg/kg, intraperitoneally daily into rats for four consecutive days and demonstrated a 2.5-fold increase in levels of hepatic ALA-synthetase. It was unclear from their study which of the steroids was responsible for the activity. Our study indicates that it is alfadalone. Until recently it was believed that 5β-steroids are potent porphyrin-inducing agents while 5α-steroids have a much lower potency. Recent studies have failed to substantiate this presumed difference. That alfadalone, a steroid with a 5α-configuration, is a potent porphyrin-inducing drug is in agreement with these recent findings.

From a clinical point of view it is important to compare the effects of alfadalone on the heme biosynthetic pathway with those of thiopental. This follows from the fact that these two drugs are used for comparable anesthetic purposes and thiopental is known to precipitate attacks of hepatic porphyria in patients who have the latent genetic disease. Thiopental, 0.3 mg/egg, injected into the 18-day-old chick embryo produces an increase in ALA-synthetase activity comparable to that produced by alfadalone, 1 mg/egg (fig. 3). Similarly, thiopental, 1 mg/egg, produces an effect comparable to that produced by alfadalone, 3 mg/egg. On this basis, alfadalone can be judged to have approximately a third the potency of thiopental. Alfadalone has a potency comparable to that of thiopental in chick embryo liver cell culture (fig. 4). Since the dosage of alfadalone used as an induction anesthetic in man (47.3 mg/70 kg) is considerably less than that of thiopental (350 mg/70 kg), the data indicate that alfadalone in therapeutic doses is less likely than comparable anesthetic doses of thiopental to increase ALA-synthetase activity in a patient who has hereditary hepatic porphyria.

An important question that remains to be answered concerns the relevance of the chick embryo data to patients who have hereditary hepatic porphyria. In a recent study, Marks compared the chick embryo data obtained with 29 drugs with clinical experiences in these drugs in hereditary hepatic porphyria. The results of the chick embryo data were in accord with clinical experience for 29 drugs; for the remaining six drugs the results were not definitive. It was concluded that tests in the chick embryo have considerable predictive value for hereditary hepatic porphyria.

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References