

Ciprofibrate

(RS)-2-[4-(2,2-dichlorocyclopropyl)phenoxy]-2-methylpropanoic acid

Ciprofibrate is a hypolipidemic drug used to lower plasma lipid levels. It modulates lipid metabolism through activation of peroxisome proliferator-activated receptor-alpha (PPAR α).

Ciprofibrate passed the animal data screen, underwent a preliminary toxicological evaluation, and is being brought to the Carcinogen Identification Committee for consultation. This is a compilation of the relevant studies identified during the preliminary toxicological evaluation.

Epidemiological data

No cancer epidemiology studies were identified.

Animal carcinogenicity data

- 60-week dietary studies
 - Male F344 rats: Rao *et al.* (1984)
 - *Increase in liver tumors (by pairwise comparison)*
 - Male F344 rats: Rao and Subbarao (1997)
 - *Increase in liver tumors (by pairwise comparison)*
 - Male F344 rats: Rao and Subbarao (1999)
 - *Increase in liver tumors (by pairwise comparison)*
- 70-week dietary studies
 - Male F344 rats: Rao *et al.* (1986)
 - *Increase in liver tumors (by pairwise comparison)*
- Two-year dietary studies
 - Male and female F344 rats: Spencer *et al.* (1989)
 - *Increase in rare carcinoid tumors of the fundus of the stomach (by pairwise comparison and trend)*
 - *One rare carcinoid tumor of the fundus of the stomach occurred in females*
- 14- to 22-month dietary studies
 - Male F344 rats: Milano *et al.* (1987)
 - *Increases in liver tumors and metastasis to the lung (by pairwise comparison)*

- 22-month dietary studies
 - Male F344 rats: Rao and Subbarao (1995)
 - *No findings of treatment-related pancreatic and testicular tumors. Other sites not examined.*
- 18- and 21-month dietary studies
 - Male C57BL/6N mice: Rao *et al.* (1988)
 - *Increase in liver tumors (by pairwise comparison and trend)*
- Two-year dietary studies
 - Male B6C3F₁ mice: Hegi *et al.* (1993)
 - *Increase in liver tumors (by pairwise comparison)*
- 155-week gavage studies
 - Male and female marmosets: Graham *et al.* (1994)
 - *No findings of treatment related tumors in three animals. (Comment: small numbers and study duration was < 25% of marmoset lifespan)*

Other relevant data

- Genotoxicity
 - Rat hepatocyte sister chromatid exchanges, chromosome aberrations, and micronuclei (*positive*): Reisenbichler and Eckl (1993)
 - DNA alterations and 8-hydroxydeoxyguanosine adduct formation *in vivo* in rat liver (*positive*): Kasai *et al.* (1989), Randerath *et al.* (1991), Huang *et al.* (1994)
 - DNA synthesis *in vivo* in rat liver (*positive*): Yeldandi *et al.* (1989)
- Mechanistic considerations
 - H- and K-*ras* gene mutations in ciprofibrate mouse liver tumors (*differ from mutations in spontaneous tumors*): Hegi *et al.* (1993)
 - Gene expression profiling in rat liver slices (*expression profile of ciprofibrate is similar to that of other peroxisome proliferators*): Werle-Schneider *et al.* (2006)
 - Transcription factor NF- κ B activity in rat liver *in vivo* (*increased*): Li *et al.* (1996)
 - NF- κ B and AP-1 activities in mouse liver *in vivo* (*increased*): Nilakantan *et al.* (1998)
 - Chromatin sphingomyelin alterations in rat liver *in vivo* (*induced*): Albi *et al.* (2003)
 - Reviews of PPAR α agonists and human relevance of rodent liver tumors: IARC (1994), Guyton *et al.* (2009)

- Structure activity considerations
 - Structurally similar to two other fibrate compounds that induce liver tumors in rodents and are Proposition 65 carcinogens: clofibrate and gemfibrozil

References¹

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¹ Excerpts or the complete publication have been provided to members of the Carcinogen Identification Committee, in the order in which they are discussed in this document.

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