

## Preparations

**USP 31:** Maprotiline Hydrochloride Tablets.

### Proprietary Preparations (details are given in Part 3)

**Austria:** Ludiomil; **Belg.:** Ludiomil; **Braz.:** Ludiomil; **Canad.:** Ludiomil; **Chile:** Mapromil; **Cz.:** Ludiomil; Maprotibene; **Denm.:** Ludiomil; Maludil; **Fr.:** Ludiomil; **Ger.:** Depriplep; Ludiomil; Maprolu; **Gr.:** Aprotilin; Ludiomil; **Hong Kong:** Ludiomil; **Hung.:** Ludiomil; Maprolu; **Indon.:** Ludiomil; **Lu.:** Sandepin; Tilsan; **Israel:** Melodil; **Ital.:** Ludiomil; **Malaysia:** Ludiomil; **Mex.:** Ludiomil; **Neth.:** Ludiomil; **NZ:** Ludiomil; **Pol.:** Ludiomil; **Port.:** Ludiomil; **Rus.:** Ludiomil (Людиомил); **S.Afr.:** Ludiomil; **Singapore:** Ludiomil; **Spain:** Ludiomil; **Swed.:** Ludiomil; **Switz.:** Ludiomil; **Thai:** Ludiomil; **Turk.:** Ludiomil; Maproti; **UK:** Ludiomil; **Venez.:** Ludiomil.

## Melitracen Hydrochloride (USAN, rINNM)

Hydrocloruro de melitraceno; Mélitracène, Chlorhydrate de; Melitraceni Hydrochloridum; N-7001; U-24973A. 3-(9,10-Dihydro-10,10-dimethyl-9-anthrylidene)propyldimethylamine hydrochloride.

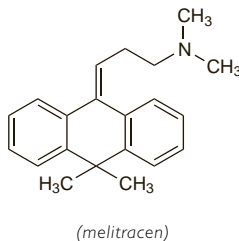
Мелитрацена Гидрохлорида

$C_{21}H_{25}N.HCl = 327.9$ .

CAS — 5118-29-6 (melitracen); 10563-70-9 (melitracen hydrochloride).

ATC — N06AA14.

ATC Vet — QN06AA14.



## Profile

Melitracen is a tricyclic antidepressant (see Amitriptyline, p.376).

In the treatment of depression (p.373) melitracen is given orally as the hydrochloride although doses are expressed in terms of the base; melitracen hydrochloride 28.1 mg is equivalent to about 25 mg of melitracen. The recommended initial dose is the equivalent of 25 mg two or three times daily gradually increased to a total of 225 mg daily if necessary. Elderly patients should generally be given reduced doses of 25 or 30 mg daily initially. Melitracen may also be given in lower doses with flupentixol (p.997) in the management of depression with anxiety. A combination of melitracen 10 mg and flupentixol 500 micrograms is given orally in the morning and at midday. In severe cases the morning dose may be doubled. The total daily dose should not exceed melitracen 40 mg and flupentixol 2 mg. Elderly patients may be given melitracen 10 mg with flupentixol 500 micrograms in the morning; in severe cases this dose may be given in the morning and at midday.

Melitracen should be withdrawn gradually to reduce the risk of withdrawal symptoms.

## Preparations

### Proprietary Preparations (details are given in Part 3)

**Austria:** Dixeran; **Belg.:** Dixeran†.

**Multi-ingredient:** **Austria:** Deanaxit; **Belg.:** Deanaxit; **Hong Kong:** An-free; Deanaxit; **Ital.:** Deanaxit; **Singapore:** Deanaxit; **Spain:** Deanaxit; **Switz.:** Deanaxit; **Thai:** Deanaxit.

## Mianserin Hydrochloride (BANM, USAN, rINNM)

Hydrocloruro de mianserina; Mianserinihydrokloridi; Mianserin Hidroklorür; Miansérine, chlorhydrate de; Mianserin-hydrochlorid; Mianserinhydroklorid; Mianserini hydrochloridum; Mianserinihydrochloridas; Mianseriny chlorowodorek; Mianserinihydroklorid; Org-GB-94. 1,2,3,4,10,14b-Hexahydro-2-methyldibenzo[c,f]pyrazino[1,2-a]zepine hydrochloride.

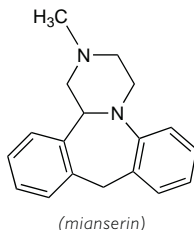
Миансерина Гидрохлорида

$C_{18}H_{20}N_2.HCl = 300.8$ .

CAS — 24219-97-4 (mianserin); 21535-47-7 (mianserin hydrochloride).

ATC — N06AX03.

ATC Vet — QN06AX03.



**Pharmacopoeias.** In *Eur.* (see p.vii).

**Ph. Eur. 6.2** (Mianserin Hydrochloride). A white or almost white crystalline powder or crystals. Sparingly soluble in water; slightly soluble in alcohol; soluble in dichloromethane. A 1% solution in water has a pH of 4.0 to 5.5. Protect from light.

## Adverse Effects

Antimuscarinic and cardiac adverse effects are fewer and milder with mianserin, a tetracyclic antidepressant, than with tricyclic antidepressants but effects are otherwise broadly similar (see Amitriptyline, p.376); mianserin may be associated with a lower risk of cardiotoxicity in overdose.

The most common adverse effect associated with mianserin is drowsiness. Mianserin also causes bone-marrow depression usually presenting as leucopenia, granulocytopenia, or agranulocytosis; aplastic anaemia has been reported. These adverse haematological reactions generally occur during the first few weeks of therapy and especially in the elderly.

Other adverse effects reported include disturbances of liver function and jaundice, breast disorders (gynaecomastia, nipple tenderness, and non-puerperal lactation), and polyarthralgia.

**Effects on the blood.** Between 1976 and the end of 1988 the UK CSM had received 239 reports of adverse haematological reactions associated with mianserin use.<sup>1</sup> The reports included 68 of agranulocytosis and 84 of granulocytopenia or leucopenia where mianserin was considered to be the probable or possible cause; there had been 17 fatalities. Allowing for the pattern of prescribing there was a greater number of reports of white blood cell disorders in patients over 65 years of age but there was no sex difference. The data also indicated that the adverse reactions were most likely to develop during the first 3 months of therapy. By the end of 1992 the number of reports of mianserin-induced agranulocytosis or neutropenia received by the CSM<sup>2</sup> had risen to 79 and 105, respectively.

A case of fatal aplastic anaemia associated with mianserin use has also been reported.<sup>3</sup>

Proposed mechanisms of mianserin haematotoxicity have included a direct toxicity<sup>4</sup> and an immunologically-mediated mechanism.<sup>5</sup> There is evidence from studies *in vitro* of a significant correlation between the desmethyl metabolite and cytotoxicity. Mianserin is given as a racemic preparation and the formation of metabolites was greater with the *R*(-)-enantiomer than with the *S*(+)-enantiomer.<sup>6</sup>

1. CSM. Mianserin and white blood cell disorders in the elderly. *Current Problems* 25 1989. Also available at: [http://www.mhra.gov.uk/home/idcplg?IdcService=GET\\_FILE&dDocName=CON2024441&RevisionSelectionMethod=LatestReleased](http://www.mhra.gov.uk/home/idcplg?IdcService=GET_FILE&dDocName=CON2024441&RevisionSelectionMethod=LatestReleased) (accessed 05/08/08)
2. CSM/MCA. Drug-induced neutropenia and agranulocytosis. *Current Problems* 1993; 19: 10–11. Also available at: [http://www.mhra.gov.uk/home/idcplg?IdcService=GET\\_FILE&dDocName=CON2024456&RevisionSelectionMethod=LatestReleased](http://www.mhra.gov.uk/home/idcplg?IdcService=GET_FILE&dDocName=CON2024456&RevisionSelectionMethod=LatestReleased) (accessed 05/08/08)
3. Durrant S, Read D. Fatal aplastic anaemia associated with mianserin. *BMJ* 1982; 285: 437.
4. O'Donnell JL, et al. Possible mechanism for mianserin induced neutropenia associated with saturable elimination kinetics. *BMJ* 1985; 291: 1375–6.
5. Stricker BHC, et al. Thrombocytopenia and leucopenia with mianserin-dependent antibodies. *Br J Clin Pharmacol* 1985; 19: 102–4.
6. Riley RJ, et al. A stereochemical investigation of the cytotoxicity of mianserin metabolites *in vitro*. *Br J Clin Pharmacol* 1989; 27: 823–30.

**Effects on the cardiovascular system.** Although mianserin is considered to be less cardiotoxic than the tricyclic antidepressants adverse effects have been noted in individual patients. Two elderly patients developed signs of disturbed cardiac function (cardiac failure, atrial and ventricular fibrillation, bradycardia, and frequent ventricular ectopic beats) which resolved after the drug was stopped.<sup>1</sup> One of the patients also developed hypokalaemia which was possibly caused by mianserin. It was suggested that persons most likely to experience problems were the elderly with a history of cardiovascular disorders. Further reports of mianserin-induced cardiac effects include recurrent ventricular fibrillation in a 61-year-old man after an overdose of mianserin<sup>2</sup> and bradycardia in a 50-year-old woman after a therapeutic dose.<sup>3</sup>

1. Whiteford H, et al. Disturbed cardiac function possibly associated with mianserin therapy. *Med J Aust* 1984; 140: 166–7.
2. Haefeli WE, et al. Recurrent ventricular fibrillation in mianserin intoxication. *BMJ* 1991; 302: 415–16.
3. Carcone B, et al. Symptomatic bradycardia caused by mianserin at therapeutic doses. *Hum Exp Toxicol* 1991; 10: 383–4.

**Effects on the liver.** By March 1985 the UK CSM had received 57 reports of hepatic reactions associated with mianserin use from a total of 5 million prescriptions. Reactions had included jaundice and other abnormalities of liver function, but no fatalities had been reported.<sup>1</sup>

Case reports have also been published concerning jaundice;<sup>2–5</sup> liver function returned to normal after stopping mianserin or lowering the dose.

1. CSM. Dangers of newer antidepressants. *Current Problems* 15 1985. [http://www.mhra.gov.uk/home/idcplg?IdcService=GET\\_FILE&dDocName=CON2024422&RevisionSelectionMethod=LatestReleased](http://www.mhra.gov.uk/home/idcplg?IdcService=GET_FILE&dDocName=CON2024422&RevisionSelectionMethod=LatestReleased) (accessed 14/08/08)
2. Adverse Drug Reactions Advisory Committee. Mianserin: a possible cause of neutropenia and agranulocytosis. *Med J Aust* 1980; 2: 673–4.
3. Goldstraw PW, et al. Mianserin and jaundice. *N Z Med J* 1983; 96: 985.
4. Zarski J-P, et al. Toxicité hépatique des nouveaux anti-dépresseurs: a propos d'une observation. *Gastroenterol Clin Biol* 1983; 7: 220–1.
5. Otani K, et al. Hepatic injury caused by mianserin. *BMJ* 1989; 299: 519.

**Effects on the musculoskeletal system.** A patient developed an acute polyarthritides affecting the hands and feet 6 days after starting therapy with mianserin;<sup>1</sup> at that time the UK CSM had received 19 reports of arthritis and arthralgia associated with mianserin. For the suggestion that effects at serotonin receptors may be involved in such adverse effects see under Mirtazapine, p.410.

1. Hughes A, Coote J. Arthropathy associated with treatment with mianserin. *BMJ* 1986; 292: 1050.

**Effects on the skin.** Reports of adverse dermatological reactions in individual patients related to mianserin therapy have included toxic epidermal necrolysis<sup>1</sup> and erythema multiforme.<sup>2,3</sup>

1. Randell P. Tolvon and toxic epidermal necrolysis. *Med J Aust* 1979; 2: 653.
2. Quraishi E. Erythema multiforme during treatment with mianserin—a case report. *Br J Dermatol* 1981; 104: 481.
3. Cox NH. Erythema multiforme due to mianserin—a case against generic prescribing. *Br J Clin Pract* 1985; 39: 293–4.

**Effects on the tongue.** Glossitis associated with mianserin therapy was reported in 2 patients.<sup>1</sup> Additionally, glossitis accompanied by severe facial oedema has been noted in another patient.<sup>2</sup> In all cases symptoms resolved after withdrawal of mianserin.

1. de la Fuente JR, Berlanga C. Glossitis associated with mianserin. *Lancet* 1984; i: 233.
2. Leibovitch G, et al. Severe facial oedema and glossitis associated with mianserin. *Lancet* 1989; ii: 871–2.

**Epileptogenic effect.** By March 1985 the UK CSM had received 64 reports of convulsions associated with mianserin use from a total of 5 million prescriptions.<sup>1</sup> In a previous review<sup>2</sup> concerning 40 of these cases it was considered that a causal connection could be established only in a minority. It was suggested that mianserin is no more epileptogenic than tricyclic antidepressants, an opinion that was also shared by other reviewers.<sup>3</sup>

1. CSM. Dangers of newer antidepressants. *Current Problems* 15 1985. Also available at: [http://www.mhra.gov.uk/home/idcplg?IdcService=GET\\_FILE&dDocName=CON2024422&RevisionSelectionMethod=LatestReleased](http://www.mhra.gov.uk/home/idcplg?IdcService=GET_FILE&dDocName=CON2024422&RevisionSelectionMethod=LatestReleased) (accessed 05/08/08)
2. Edwards JG, Glen-Bott M. Mianserin and convulsive seizures. *Br J Clin Pharmacol* 1983; 15: 299S–311S.
3. Richens A, et al. Antidepressant drugs, convulsions and epilepsy. *Br J Clin Pharmacol* 1983; 15: 295S–298S.

**Overdose.** Experience with 100 consecutive cases of intoxication with mianserin<sup>1</sup> revealed that when it was the only drug ingested symptoms were mild and neither deep coma nor convulsions occurred. More serious symptoms and 2 fatalities were seen in patients who had taken multiple drug overdoses. The results suggested that after an acute overdose mianserin is less toxic than the tricyclic antidepressants. This conclusion was also supported by a large follow-up study<sup>2</sup> comparing the outcome of suicide attempts among patients who had taken mianserin in overdose with those who had taken amitriptyline.

1. Chand S, et al. One hundred cases of acute intoxication with mianserin hydrochloride. *Pharmakopsychiatrie* 1981; 14: 15–17.
2. Inman WHW. Blood disorders and suicide in patients taking mianserin or amitriptyline. *Lancet* 1988; ii: 90–2.

## Precautions

As for tricyclic antidepressants in general (see Amitriptyline, p.378). Although mianserin is less cardiotoxic than the tricyclic antidepressants, it still should be used with caution in patients with cardiovascular disorders, such as heart block, or after recent myocardial infarction. Similarly, patients with angle-closure glaucoma or prostatic hyperplasia should be monitored even though antimuscarinic effects are rare. Mianserin should be used with caution in patients with diabetes mellitus, epilepsy, and hepatic or renal impairment; it should be avoided in severe hepatic disease.

Patients should be carefully monitored during early antidepressant therapy until significant improvement in depression is observed because suicide is an inherent risk in depressed patients. For further details, see under Depression, p.373. Suicidal thoughts and behaviour may also develop during early treatment with antidepressants for other disorders; the same precautions observed when treating patients with depression should therefore be observed when treating patients with other disorders.