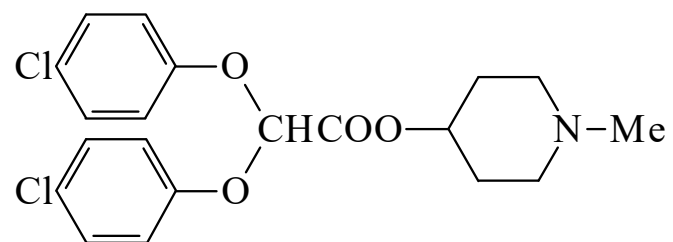




>>Entry Name: **Lifibrate**, INN, USAN

Synonym(s): 1-Methyl-4-piperidinyl bis(4-chlorophenoxy)acetate, 9Cl, 8Cl. MPCA. SaH 42-348



CAS Registry Number: 22204-91-7

Molecular Formula: $C_{20}H_{21}Cl_2NO_4$

Molecular Weight: 410.296

Accurate Mass: 409.084763

Percentage Composition: C 58.55%; H 5.16%; Cl 17.28%; N 3.41%; O 15.60%

Use/Importance: Antihyperlipidaemic agent

Physical Description: Cryst. (EtOH)

Development Status: Never marketed

Melting Point: Mp 93-95°

Calculated Log P Data: Log P 4.74 (calc)

>>Derivative: Hydrochloride

Physical Description: Cryst. (EtOH)

Melting Point: Mp 158-159°

>>Derivative: Fumarate

Physical Description: Cryst. (MeOH)

Melting Point: Mp 100-112 dec.°

References:

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