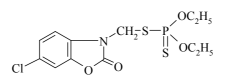
# **CIPAC STATUS REPORT**

### 28/06/2005



### 0109 Phosalone

	Allocated to F.
	CIPAC methods published in :
	CIPAC D, p. 141 (GLC)
CIPAC	10th meeting, June 1966 in France
	Dr. Prat stated that the method had only as yet been received in principle: it based on GLC. He was awaiting details from the firm.
CIPAC	11th meeting, June 1967 in London
	No collaborative work has been carried out.
CIPAC	13th meeting, June 1969 in Oeiras
	Dr. Povlsen reported that the analysis can probably be carried out similar to the analysis of malathion using a corresponding yellowcoloured cupric complex. However the potassium salt, in this case, seems to be difficult to prepare.
CIPAC	16th meeting, June 1972 in Stockholm
	Mr. Laurent presented a paper at the CIPACSymposium in Stockholm, on the methods of analysis for phosalone. Work to be initiated by F.
CIPAC	17th meeting, June 1973 in Wageningen
	To be considered by the French Committee.
CIPAC	18th meeting, June 1974 in London
	Work in progress (GLC method).
CIPAC	19th meeting, June 1975 in Oeiras
	GLC method ready for testing collaboratively, accepted as draft method.
CIPAC	20th meeting, June 1976 in Wädenswil
	The preliminary studies of the GLC method by the French Committee may be finished for the end of the year.
CIPAC	21st meeting, June 1977 in Braunschweig
	Study with GLC method will start soon.

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**CIPAC** 22nd meeting, June 1978 in Versailles

Work in progress.

**CIPAC** 23rd meeting, June 1979 in Baltimore

The GC column and the internal standard have been changed. Coll. study can start in the near future.

**CIPAC** 24th meeting, May 1980 in Salobrena

A small scale study using a HPLC method is under way (CIPAC/2816/R). A collaborative study on an international scale can probably start in November.

**CIPAC** 25th meeting, June 1981 in Gembloux

Mr Laurent reported (CIPAC/2967) that after a preliminary study phosalone was being collaboratively studied on an international level. The new method was better described than the one used in the first study.

**CIPAC** 26th meeting, May 1982 in Rome

Mr Laurent reported that the collaborative study would start in June 1982.

CIPAC 28th meeting, October 1984 in Baltimore

A collaborative study was planned against the end of the year (CIPAC/3197).

**CIPAC** 29th meeting, September 1985 in Copenhagen

The manufacturing process was reported to have been changed so that the impurities pattern was different. Mr Declercq said that additional work was needed before a collaborative study could be held.

**CIPAC** 30th meeting, June 1986 in Vienna

Mr Declercq reported that probably in September a decision would be taken concerning a possible collaborative study.

CIPAC 31st meeting, June 1987 in Cascais

Dr Declercq presented the report of a collaborative study (3416). He explained that a short GC column had been used to shorten the residence time of phosalone. Mr Declercq would send copies of the method to all participants. <u>Decision</u> The GLC method for phosalone technical and formulations, CIPAC/3418, was accepted as provisional CIPAC method.

CIPAC 32nd meeting, June 1988 in Geneva

<u>Decision</u> The provisional CIPAC method, CIPAC /3418 for phosalone technical and formulations was accepted as <u>full</u> CIPAC method.