MEMORANDUM

SUBJECT: Post-Phase II Registration Standard Support Team Meeting For Terbufos FRSTR

TO: Addressees

The Office of Pesticide Programs' (OPP) procedures for development of Registration Standards includes two major meetings of the OPP Registration Standard Project Support Team. The first of these meetings (the Pre-Phase II Meeting) is an informational meeting intended to provide an overview (regulatory history) for the chemical and put it in regulatory and scientific perspective for the support team. This meeting occurs at the very beginning of the data review and evaluation phase (Phase II) of the Registration Standard process. The presence of external program offices at this meeting will provide those offices the opportunity for up-front involvement in Registration Standard decision-making. The second of these meetings (the Post-Phase II Meeting) occurs at the conclusion of Phase II of the Registration Standard process and is intended to establish agreement on the science conclusions, data requirements, and regulatory position/rationale which flow from the science findings.

The Pre-Phase II Meeting for Terbufos FRSTR was held on September 17, 1987. Phase II of the Registration Standard process for terbufos was completed on April 14, 1988.

In keeping with OPP's procedures for development of Registration Standards, the Post-Phase II Meeting for terbufos has been scheduled for June 17, 1988 at 9:30 AM in Room 813, in Crystal Mall Building 2.

As stated above, the purpose of the Post-Phase II Meeting is to establish agreement on the science conclusions, data requirements, and regulatory position/rationale which flow from the science findings.
The attachments provide the information each participant will need for evaluating the pesticide and its uses in developing the Agency's regulatory position and rationale with regard to reregistration. These attachments are listed and described below:

A. Draft Registration Standard: This document includes the proposed regulatory position and rationale, unique label language, science summary, data tables and tolerance reassessment.

B. Preliminary Quantitative Use Analysis: This document, prepared early in Phase I of the registration standard process by the Benefits and Use Division, provides a brief description of the use(s) for the pesticide under review.

C. Science Chapters: These chapters, prepared by the scientific reviewers in Hazard Evaluation Division, document the scientific reviewer's evaluation of the data to support the reregistration of the pesticide and its uses. These chapters include:

   1) Product and Residue Chemistry Chapter
   2) Toxicology Chapter
   3) Environmental Chapter
   4) Exposure Assessment Chapter

D. Scientific Summary Memo: This memo, prepared by the Science Integration Staff of the Hazard Evaluation Division, provides the "bottom line" finding from the science chapters.

E. Summary of Registered Uses: This summary, prepared by the Benefit Use Division, as a table of contents for the larger "Index of Currently Acceptable Uses," provides a listing of the major use categories and the crops under each use.

If after reviewing the attached material you find that you are not interested in taking part in the discussion on this Registration Standard, please notify me of your decision by completing the attached "Notice of Non-Participation" form.
and return it and the material to the PM at room number 211, Crystal Mall Building 2.

If, however, you find that you are interested in taking part in the discussion on this Registration Standard you should:

- Review the attachments referred to above and come to the meeting ready to discuss the items on the attached agenda.

- If you have specific questions, let me know them ahead of time so I or other members of the team can be prepared to respond to them at the meeting.

- If you would like to revise the language of the document, or add a section, bring copies of the proposed language to the meeting for the team's consideration.

- If you only have a general interest come to the meeting and participate in the discussion.

Participants at this meeting, and decision meetings that will follow, should be authorized to state a position on behalf of their office, so that decisions made are the official position of the organizations represented.

The Registration Standard, as revised at the meeting, will be circulated to all offices after the meeting for formal concurrence. If you have any further questions, you may contact Marilyn Mautz at 557-2785.

John Tice
Acting Product Manager (16)
Insecticide-Rodenticide Branch
Registration Division (TS-767C)
Addressees:

External Program Offices

J. Reinert (OPPE)
P. Flaherty (OCM)
K. Lee (OGC)
J. MacDonald (C&T)

OPP Offices

H. Manning (HED/EAB)
H. Craven (HED/EEB)
W. Hazel (HED/RCB)
K. Barbehenn (HED/SIS)
R. Levy (HED/TB)
P. Johnson (BUD/SSB)Index)
W. Gross (BUD/SSB)(QUA)
R. Esworthy (BUD/EAB)(PQUA)
C. Grimes (PMSD/ISB)
V. Prunier (RD/PCS)
C. Bashor (PMSD/PFSS)

Attachments: Meeting Agenda
Preliminary Quantitative Use Analysis
Science Chapters
Science Summary Memo
Data Tables
Summary of Uses
Notification of Non-Participation Form
TERBUFOS

Post-Phase II Meeting Agenda

1) INTRODUCTION
   * Introduction of attendees
   * Explanation of the purpose and scope of the meeting
   * Brief description of document

2) AGREEMENT ON SCIENCE CONCLUSIONS INCLUDING THE DATA TABLES
   * Are data requirements adequately covered?
   * Are all data gaps identified?
   * Are special studies needed?
   * At what point should the data generated under the Standard be reviewed?
   * Resolution of issues.

3) DEVELOPMENT OF REGULATORY POSITION AND RATIONALE
   * Worker reentry risks:
   * Wildlife risks:
   * Resolution of issues.

4) SUMMARIZATION OF DECISIONS
NOTICE OF NON-PARTICIPATION FORM FOR TERBUFOS

TO: John Tice
Acting Product Manager (16)
Insecticide-Rodenticide Branch
Registration Division (TS-767C)
Room No. 211
Crystal Mall Building No. 2

We will not be taking part in the discussions and development of the registration standard for this chemical for the following reason(s):

__________________________________________
(name)

__________________________________________
(title)

__________________________________________
(office)

__________________________________________
(date)