

US EPA ARCHIVE DOCUMENT

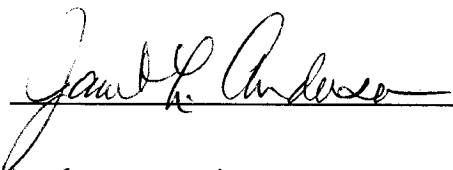


**US Environmental Protection Agency
Office of Pesticide Programs**

**Dried Blood Final Work Plan (FWP)
for Registration Review**

November 2007

Dried Blood Final Work Plan
Registration Review - Case 4030
November 2007

Approved by: 

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Biopesticides and Pollution Prevention Division

Date: November 27, 2007

Introduction:

The Agency is implementing the new Registration Review program and plans to review each registered pesticide every 15 years to determine whether it continues to meet the Federal Insecticide, Fungicide, and Rodenticide Act (FIFRA) standard for registration. Changes in science, public policy, and pesticide use practices will occur over time. The registration review program is intended to make sure that, as the ability to assess risk evolves and as policies and practices change, all registered pesticides continue to meet that statutory standard. The public phase of registration review begins when the initial docket is opened for each case. Information on this program is provided at: http://www.epa.gov/oppsrrd1/registration_review/.

This is EPA's *Final Work Plan* for the registration review of dried blood. The work plan includes the expected registration review time line. There has been a request for voluntary cancellation by the registrant of the only registered product containing the active ingredient (request from R.M. Horton to USEPA dated September 18, 2007). The Agency will inform the public of the registrant's intent to voluntarily cancel the product registration through an FR notice which is expected to be published early in 2008. If the Agency receives no comments from the public during the 30 day comment period, the registration will be cancelled. There is not a tolerance or an exemption from the requirement of a tolerance for this active ingredient.

Risk Assessment and Data Needs:

Due to the fact that the only registered pesticide containing dried blood as an active ingredient is intended to be cancelled, further risk assessments and submission of data are not necessary.

Next Steps:

The *Final Work Plan* for dried blood will be posted on the Agency's website (<http://www.epa.gov>) and a Federal Register Notice (FR-Notice) will be issued with respect to the cancellation of the pesticide containing the chemical an active ingredient.