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1. <u>Purpose</u>:

To provide a reference of PDP required compounds, a listing of available marker pesticides and process controls, specification of PDP commodity groupings, requirements for method validation and continuing quality control (QC) for USDA/AMS Pesticide Data Program (PDP) samples.

2. <u>Scope</u>:

This standard operating procedure (SOP) shall be followed by all analytical laboratories conducting pesticide residue studies for PDP, including support laboratories conducting stability or other types of studies that may impact the program.

3. <u>Outline of Procedure</u>:

- 5.1 Required Compounds
- 5.2 Standards
- 5.3 Method Validation Background
- 5.4 General Method Validation Requirements
- 5.5 Method Validation Evaluation Guidelines
- 5.6 Method Validation Scenarios
- 5.7 Marker Pesticides
- 5.8 Process Control Compounds
- 5.9 PDP Commodity Groupings
- 5.10 Establishment of Limits of Detection (LODs) and Limits of Quantitation (LOQs)
- 5.11 Verification of LODs/LOQs
- 5.12 Changing LODs
- 5.13 Determination of Method Range
- 5.14 Precision and Accuracy Data Collection
- 5.15 Method Evaluation Reporting
- 5.16 Method Validation Evaluation by USDA/AMS
- 5.17 Blanks and Spikes Required per Set and Continuing QC
- 5.18 Criteria for Method Validation and Continuing QC
- 5.19 Proficiency Testing
- 5.20 Measurement Uncertainty

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Attachment 1 – Method Evaluation Flowchart

- Attachment 2 PDP Compound Groups, Pesticides Codes and Multi-residue Compound Groupings for Fruit and Vegetables
- Attachment 3 EPA, Codex, and Food and Drug Administration (FDA) Pesticide Analytical Manual (PAM) Commodity Groupings

Attachment 4 – FDA Information

Attachment 5 – Method Evaluation Reporting Forms [LOD Verification, Determination of Method Range, Precision and Accuracy Data Collection]

Attachment 6 – Process Control and Spike Recovery Acceptability Flowchart

4. <u>References</u>:

- de Kok et. al., *The Stability of Pesticide Standards and Solutions*, 5th European Pesticide Residue Workshop, Stockholm, Sweden, June 13-16, 2004
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- National Environmental Laboratory Accreditation Conference (NELAC), *Standards*, Appendix D, Section D.1.1.2.1, Laboratory Control Samples (LCS), June 5, 2003
- U.S. FDA, *Standard Operating Procedures for the Total Diet Study*, KCM TD G2, revision 0, Quality Assurance, January, 1993
- Association of Official Analytical Chemists (AOAC), *Quality Assurance Principles for Analytical Laboratories*, 1991, pp. 91-94
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- Federal Register, Rules and Regulations, Volume 49, Number 209, October, 1984
- Horwitz, W., Evaluation of Analytical Methods Used for Regulation of Foods and Drugs, Analytical Chemistry, Vol. 54, No. 1, pp. 67A-76A, 1982
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- U.S. EPA, Reagents and Solutions, 40 CFR 160.83
- U.S. EPA, Test, control and reference substance characterization, 40 CFR 160.105
- U.S. EPA, Test, control, and reference substance handling, 40 CFR 160.107

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- U.S. EPA, Pesticide Use Index Index of pesticide use sites: Corresponding Major Use Pattern(s) and Crop Group,

http://www.epa.gov/opp00001/regulating/usesite/terrestrial-food.pdf

- U.S. FDA, <u>Pesticide Analytical Manual Volume I (PAM) 3rd Edition</u>, Chapter 2, <u>http://www.fda.gov/downloads/Food/ScienceResearch/LaboratoryMethods/PesticideAnalysisManualPAM/ucm111500.pdf</u>
- Codex Alimentarius Commission, *Pesticide Residues in Food and Feed*, http://www.fao.org/fao-who-codexalimentarius/standards/pestres/commodities/en/

5. <u>Specific Procedures</u>:

This SOP represents minimum PDP requirements and is presented as a general guideline. Each laboratory shall have written procedures that provide specific details concerning how the procedure has been implemented in that laboratory.

5.1 Required Compounds

5.1.1 Refer to applicable commodity/compound-specific memoranda for commodity specific testing profiles.

5.1.2 Priority Levels

5.1.2.1 Each analyte of interest for each assigned commodity shall be designated with a priority level by the USDA/AMS. Priority levels for the individual compounds in the commodity-specific memoranda posted to the PDP Extranet are based on data needs identified by data users/stakeholders (e.g., U.S. Environmental Protection Agency, U.S. Food and Drug Administration, grower groups, industry, consumer/environmental groups), current tolerances and Action Levels (ALs), and national/international Maximum Residue Levels (MRLs). In addition, compounds that may not have tolerances in the U.S., but are known to be used in countries that export food to the U.S. are included; these compounds are comprised of compounds identified by EPA as having a high probability of consumption in selected imported products, and analytes identified by FDA or USDA Foreign Agricultural Service (FAS) as of interest in selected imported products, where applicable to a given commodity. It is recognized that not all

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compounds/metabolites on a given list are amenable to multiresidue testing and final screening lists will be determined based on method validation and ongoing testing results.

5.1.2.2 In the various commodity-specific memoranda (separate documents posted to the PDP Extranet), compounds identified as Priority 1 compounds are the most critical and those identified as Priority 4 are the least critical. The priority level is a combination of data needs and expected feasibility of current methods to recover a given compound. General priority levels are assigned according to the following protocol:

5.1.2.2.1 Priority 1 compounds are selected multiresidue-amenable pyrethroids, organophosphates, and carbamates and their associated metabolites. Priority 1 compounds are required for all commodities. These compounds are critical because they are scheduled for EPA Registration Review, as documented on the current EPA Office of Pesticide Programs Registration Review Schedule.

5.1.2.2.2 Priority 2 compounds include other multiresidue-amenable compounds with a current tolerance for the given commodity that are highly important because they also have upcoming reviews scheduled or have been identified by a stakeholder as a highly important data need. Cyphenothrin, imiprothrin, and tetramethrin are also included as priority level 2 compounds for all commodities. Additionally, chemicals used in other countries may be included as Priority 2 compounds, dependent upon their anticipated method behavior.

5.1.2.2.3 Priority 3 compounds include other analytes with tolerances (including food handing establishment tolerances) or ALs (e.g., environmental contaminants/extraneous residues – aldrin, BHC, chlordane, DDD, DDE, DDT, dieldrin, endrin, heptachlor, and heptachlor epoxide) for the given commodity and are routinely analyzed by multiresidue methods. Priority 3 compounds may also include chemicals used in other countries, dependent upon their anticipated method behavior.

5.1.2.2.4 Priority 4 compounds include pesticides that have current tolerances, but likely require single analyte methods (e.g., glyphosate/AMPA, paraquat/diquat, EBDCs). Priority 4 compounds may also include chemicals used in other countries, dependent upon their anticipated method behavior.

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5.1.2.3 Laboratories should include all Priority 1 compounds, as many Priority 2 compounds as possible, and as many Priority 3 compounds as feasible.

5.1.2.4 In some cases, PDP will authorize the development of new methods to detect certain compounds (e.g., triazole metabolites, phenoxies, formetanate hydrochloride).

5.2 Standards

5.2.1 Ordering Analytical Standards

5.2.1.1 Standards may be obtained from the EPA Repository, registrants, or commercial vendors. When requesting standards from the Repository, identify your laboratory as a PDP laboratory in the comment section of the order form so that the Repository staff will know that the order takes precedence. If the request is urgent, note that in the Comment section of the order form as well.

The EPA repository is located at:

EPA National Pesticide Standard Repository Environmental Science Center 701 Mapes Road Fort Meade, MD 20755-5350 Phone: (410)305-2931 FAX: (410) 305-2999 http://www.epa.gov/pesticides/labs/standards_repository.html

5.2.1.2 Procurement of standards from all sources must meet the following minimum requirements:

Availability of a current and valid "Certificate of Analysis" (CoA) (as a minimum requirement the certification shall identify the substance, its purity, and the production lot), traceability, and current expiration date.

An exemption for CoA and current expiration date is allowed for extraneous environmental contaminants that are covered by FDA Action Levels and compounds that

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have been revoked and no longer have existing U.S. registrations. Extraneous environmental contaminants include aldrin, BHC, chlordane, DDT (and metabolites), dieldrin, endrin, heptachlor (and metabolite), and lindane. Examples of revoked compounds that no longer have existing U.S. registrations include parathion ethyl, chlorfenvinphos, and fenchlorphos.

For all other analytical standards, in some cases, a current and valid CoA may not accompany the analytical standard. In this case, the laboratory shall contact the vendor to determine if one is available; if one is not available, the laboratory is exempt from the requirement to maintain a current and valid CoA for that standard.

5.2.2 Receipt of Analytical Standards

Custody of a standard begins when the standard is received in the laboratory. Each standard shall be given a code that uniquely identifies the standard from neat material to final dilutions. Receipt of standards shall be documented and each standard shall be traceable. Records shall include name, unique code, purity, lot number, date received, and expiration date (see 5.2.1.2 for exemption).

5.2.3 Storage of Analytical Standards

5.2.3.1 Neat standards shall be kept in a separate standards freezer, preferably at approximately -20° C or lower unless degradation occurs at such temperatures. In these cases, neat standards shall be stored at the recommended temperature.

5.2.3.2 Stock standards and dilutions including mixed standards shall be kept in refrigerators or freezers separate from those used for samples. Stock standards and dilutions shall be stored in teflon-lined, screw-capped, glass bottles or sealed glass ampules.

5.2.3.3 Access to the freezers and refrigerators shall be controlled and standards usage documented through the use of appropriate records (e.g., log books). These records shall contain at a minimum: standard name and/or unique code, date and time removed, initials of person removing standard, date and time returned, initials of person returning standard.

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5.2.3.4 Refrigerator and freezer temperatures shall be checked either by taking readings each working day, or by automatic temperature recording devices.

5.2.3.5 When a neat standard is removed from freezer storage, the standard should be stored in a desiccator while it is brought to room temperature to minimize the potential for hydrolysis.

5.2.4 Preparation of Stock Standard Solutions

Stock standard solutions shall be prepared in a separate standard preparation area to avoid contamination of samples with pesticide standards. Each stock standard shall be given a unique identifying code and shall be labeled with a minimum of: pesticide name, concentration, solvent, date of preparation, initials of preparer, and expiration date of solution. Written SOPs for stock standard preparation shall include the method for preparing standards, calculations used in standard preparation, documentation that provides for standard traceability and safety guidelines.

5.2.5 **Preparation of Intermediate Dilutions**

Intermediate dilutions, including mixed standards, shall be prepared in a separate standard preparation area. Each standard shall be given a unique identifying code and shall be labeled with pesticide name, concentration, solvent, date of preparation, initials of preparer, and expiration date of solutions. Written SOPs shall include the method for standard preparation and documentation that provides for standard traceability.

5.2.6 Standard Checking

5.2.6.1 Stock solutions of neat pesticide standards not previously prepared or not currently in use in the laboratory shall be prepared in duplicate and the two standards compared to each other. Responses for standards of comparable concentrations must match within 15% relative percent difference (RPD):

$$RPD = \frac{|RF_1 - RF_2|}{\left[\frac{RF_1 + RF_2}{2}\right]} \times 100$$

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where RF_1 is the response factor¹ of the first analytical standard and RF_2 is the response factor of the second standard. If standards do not match, a third standard shall be made and compared. This process shall be continued until two matching standards are prepared.

5.2.6.2 New stock solutions that are prepared from neat pesticides currently used in the laboratory shall be compared to the old stock solution. The two standards must match within 15% RPD. If the two standards do not match, the problem must be identified and solved before the standard is used for quantitation. A suggested approach is to make new dilutions of both the old and new standards to check for dilution errors. If no dilution errors are found, a second stock dilution should be made to determine whether an error was made in the original preparation from neat material. If these two stocks match, then the standard may be used. If they do not match, a third stock solution should be made. Whenever possible, duplicate injections shall be used.

5.2.6.3 Documentation of the standard checking process shall be kept through appropriate records (i.e. logs). Chromatograms of all standards shall be kept indicating the standard comparisons of old and new standards and the calculated difference.

5.2.7 Expired Standard Verification

If a laboratory has an expired neat analytical standard and cannot obtain a replacement with a valid expiration date from an approved PDP vendor or the EPA National Pesticide Standard Repository, with a deviation from USDA/AMS on file, the laboratory may proceed with validation and analysis of samples using the expired standard under the following conditions:

5.2.7.1 If the standard is recertified by the vendor and new documentation is obtained, it shall be recorded in the laboratory's standard records.

5.2.7.2 If the standard is not recertified, it shall be compared to an unexpired neat when one is available to verify its integrity.

5.2.7.2.1 If the two standards' response factors are within 15%, the expired standard being used shall be considered fit for purpose and this data shall be recorded in the laboratory's records.

¹ Area or height of each standard divided by the concentration of that standard.

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5.2.7.2.2 If the two standards' response factors are not within 15%, USDA/AMS shall be contacted.

5.2.7.2.2.1 If there are residues, USDA/AMS and the laboratory's Technical Program Manager (TPM) and Quality Assurance Officer (QAO) shall develop an agreement on how to proceed with samples containing residues (e.g., re-extract and analyze with unexpired standard, code data as estimates, change to "unable to analyze," etc.). The agreement shall be documented and recorded in the laboratory's records. USDA/AMS will update any transmitted data in the USDA/AMS database.

5.2.7.2.2.2 If there are non-detects and the expired standard produces a response less than the response of the unexpired standard, the LOD shall be raised (consult with USDA/AMS to determine the level) and this information shall be recorded in the laboratory's records. USDA/AMS will update any transmitted data in the USDA/AMS database. The expired standard being used shall be considered fit for purpose for qualitative analysis only and this declaration shall be recorded in the laboratory's records.

5.2.8 Working Dilutions/Mixed Standards

5.2.8.1 Working dilutions and mixed standards shall be checked to ensure integrity of the solutions. These solutions should be made as frequently as necessary to ensure that concentrations do not change and/or individual pesticides do not degrade. Each laboratory shall determine the frequency of remaking dilutions/mixed standards. Documentation supporting this decision shall be maintained. A suggested guideline is six months for stock mixed standards and one month for working dilutions. Some pesticides may require more frequent dilution from the stock.

5.2.8.2 An archive file of all old mixed standards shall be kept and the dates the standards were used shall be indicated. The archive file shall be maintained a minimum of five years.

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5.2.8.3 All working/mixed standards shall be identified by a unique and traceable code. Working/mixed standard records shall contain a minimum of pesticide name, solvent, date of preparation, expiration date, and preparer.

5.2.9 Detector Profiles

Standard retention time and response shall be characterized by analysis on the detectors used in each laboratory. These include but are not limited to: GC-ECD, GC-FPD, GC-ELCD, GC-XSD, GC-MSD, GC-ITD, LC-MS, and tandem MS. Libraries of all standards shall be developed for confirmatory instruments (GC-MS and LC-MS systems).

5.2.10 Disposal of Analytical Standards

Each laboratory shall establish the proper procedures for disposal (e.g., disposal by a licensed contractor) of expired analytical standards (both neat standards and dilutions). Disposal shall be in accordance with the laboratory's Chemical Hygiene Plan and shall be documented.

5.3 Method Validation Background

5.3.1 Marker compounds and commodity groups were created to facilitate the validation and ongoing QC of the enormous number of combinations of pesticides and commodities included in PDP. Each concept seeks to group pesticides or commodities by common properties and exploits these common properties to reduce the possible combinations to a manageable number.

5.3.2 This method evaluation framework makes the following assumptions:

5.3.2.1 Commodities are grouped in such a way that assessment of method performance in one commodity in the group can be extended to apply to all commodities in the group.

5.3.2.2 Marker pesticides are chosen to be representative of a broad range of similar pesticides. The assessment of method performance for these pesticides can be extended to apply to similar pesticides.

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5.3.2.3 LOD is specific to a <u>pesticide-commodity pair</u> and must be evaluated for every pesticide-commodity pair.

5.3.2.4 Although a method may be extended to other commodities and pesticides, a minimum amount of LOD verification and recovery data must be obtained to confirm this assumption.

5.3.3 This SOP details various scenarios and their corresponding method validation requirements.

5.3.4 When problems occur, such as instrument reproducibility and/or linearity, an investigation of causes shall be conducted. A flow diagram is attached (*see Attachment 1 – Method Evaluation Flowchart*) which further clarifies these concepts.

5.4 General Method Validation Requirements

5.4.1 Methods selected for use by PDP laboratories, and significant changes to approved methods, are subject to prior approval by USDA/AMS.

5.4.2 The laboratory shall complete all required method validation modules, with the exception of precision and accuracy data collection (extracted, analyzed, and reviewed) prior to the extraction of any routine analytical sample sets.

5.4.3 An extraction/detection system includes the whole method: extraction, clean-up, chromatography, and analytical technique.

5.5 Method Validation Evaluation Guidelines

5.5.1 The following scenarios shall be followed for validation of new methods or changes/additions to existing methods. The following scenarios of changes/additions are possible:

5.5.1.1 Implementing a new method (5.6.1)

5.5.1.2 Changing an analytical method

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- 5.5.1.2.1 Extraction (5.6.2.1)
 5.5.1.2.2 Post-extraction/pre-instrumentation (5.6.2.2)
 5.5.1.2.3 Instrumentation new Limit of Detection (LOD) (5.6.2.3)
- **5.5.1.2.4** Minor Modifications (5.6.2.4)

5.5.1.3 Adding a new commodity grouping (5.6.3)

5.5.1.4 Adding a raw agricultural commodity or a processed commodity to an existing commodity group (5.6.4)

5.5.1.5 Adding pesticides related to marker pesticide groups to an existing commodity group (5.6.5). (see Attachment 2 – PDP Compound Groups, Pesticides Codes and Multi-residue Compound Groupings for Fruit and Vegetable).

5.5.1.6 Adding a new pesticide that is not related to marker pesticide groups to an existing commodity group. (5.6.6)

5.5.2 Evaluation takes place through the performance of method evaluation modules. These modules are chosen to meet the requirements of each scenario. The modules are:

- Establishment of LODs and Limits of Quantitation (LOQs) (5.10)
- Verification of LODs/LOQs (5.11)
- Determination of Method Range (from 1xLOQ to 10xLOQ) (5.13)
- Precision and Accuracy Data Collection at 2xLOQ (5.14)
- Method Evaluation Reporting (5.15)

5.5.3 Section 5.6 of this SOP lists each scenario and the modules that must be performed in that scenario. Sections 5.10 through 5.15 outline the detailed procedures to be followed for each module.

5.6 Method Validation Scenarios

The TPM and QAO will determine which scenario described in the following subsections applies for the analytes/commodities/methods pairings (see Attachment 2 - PDP Compound Groups,

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Pesticides Codes and Multi-residue Compound Groupings for Fruit and Vegetable). If local agreement cannot be reached, the PDP Technical Director shall be contacted to determine which modules should be performed.

5.6.1 New method implementation – Proceed with:

- Establishment of LODs and (LOQs) (5.10)
- Verification of LODs/LOQs for all compounds (5.11)
- Determination of Method Range for marker compounds (5.13)
- Precision and Accuracy Data Collection for all compounds (5.14)
- Method Evaluation Reporting (5.15)

5.6.2 Method changes

5.6.2.1 Major Extraction Change - Examples would be using a different solvent, solid phase extraction (SPE) sorbent bed, or a new technique. Proceed with:

- Establishment of LODs and (LOQs) (5.10)
- Verification of LODs/LOQs for all compounds (5.11)
- Determination of Method Range for marker compounds (5.13)
- Precision and Accuracy Data Collection for all compounds (5.14)
- Method Evaluation Reporting (5.15)

5.6.2.2 Major changes in post-extraction/pre-instrumentation procedures (cleanup) - Proceed with:

- Verification of LODs/LOQs for all compounds (5.11)
- Determination of Method Range for marker compounds (5.13)
- Precision and Accuracy Data Collection for all compounds (5.14)
- Method Evaluation Reporting (5.15)

5.6.2.3 Instrumentation Changes - The TPM and QAO will determine if the instrument change warrants completion of the following sections. *This is dependent upon the extent of modification. If local agreement cannot be reached, the PDP Technical Director shall be contacted for further resolution.*

For new LOD - Proceed with:

• Establishment of LODs/LOQs for all compounds (5.10)

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- Verification of LODs/LOQs for all compounds (5.11)
- Method Evaluation Reporting (5.15)

The laboratory shall use best professional judgment to determine if Precision and Accuracy Data Collection (subsection 5.14) is necessary.

5.6.2.4 Minor modifications of existing method - The TPM and QAO will determine which portions of the following sections will be completed. *This is dependent upon the extent of modification*. *If local agreement cannot be reached, the PDP Technical Director shall be contacted to determine which sections should be performed.*

- Establishment of LODs and LOQs of affected analytes (5.10)
- Verification of LODs/LOQs of affected analytes (5.11)
- Determination of Method Range of affected markers (5.13)
- Precision and Accuracy Data Collection of affected analytes (5.14)
- Method Evaluation Reporting (5.15)

5.6.3 Adding a new commodity group - Proceed with:

- Verification of established LODs/LOQs for all required pesticides in the new commodity (5.11)
- Determination of Method Range for the marker pesticides (5.13)
- Precision and Accuracy Data Collection for all required analytes (5.14)
- Method Evaluation Reporting (5.15)

5.6.4 Adding a raw agricultural commodity or processed commodity (i.e., canned/frozen/dried/ juice) to an existing commodity group. Proceed with:

- Verification of established LODs/LOQs for all required pesticides (5.11)
- Precision and Accuracy Data Collection (2 points) for all required pesticides (5.14)
- Method Evaluation Reporting (5.15)

The laboratory shall use best professional judgment to determine if additional validation is necessary based on matrix behavior.

5.6.5 Adding pesticides related to the marker pesticide groups to an existing commodity group – Proceed with:

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- Establishment of LODs and LOQs for each pesticide added (5.10)
- Verification of LODs/LOQs for each pesticide added (5.11)
- Precision and Accuracy Data Collection for each pesticide added (5.14)
- Method Evaluation Reporting (5.15)

5.6.6 Adding pesticides that are not related to the marker pesticide groups to an existing commodity group: (For example, the addition of imidacloprid analyzed by the same multiresidue procedure. The new pesticide may then become a marker pesticide for similar pesticides that are later added.) - Proceed with:

- Establishment of LODs and LOQs for each pesticide added (5.10)
- Verification of LODs/LOQs for each pesticide added (5.11)
- Determination of Method Range for compound(s) that are to become marker(s) (5.13)
- Precision and Accuracy Data Collection for each pesticide added (5.14)
- Method Evaluation Reporting (5.15)

5.7 Marker Pesticides

5.7.1 Assigning Compounds to Marker Groups

5.7.1.1 Compounds are placed into marker groups based on a combination of analyte chemistry and method performance behavior. Initial compound designations are made by the Technical Advisory Group (TAG), with applicable analytical laboratory input based on known method behavior, if those data are available. For new compounds, behavior data may not be available.

5.7.1.2 Final marker group assignment, and any marker group assignment changes, are based on laboratory experience. USDA/AMS maintains an "Effective Date" field that tracks initial group assignment as well as any changes in that initial assignment.

5.7.2 Multi-residue Screening

5.7.2.1 A laboratory may choose to use marker groups, rotate spike mixtures between analytical sets, or spike all compounds analyzed, as long as each extraction/detection system is adequately represented within each set.

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5.7.2.2 For laboratories using marker groups, each laboratory shall select at least one compound from each applicable group (*see Attachment 2 – PDP Compound Groups, Pesticides Codes and Multi-residue Compound Groupings for Fruit and Vegetable*) to serve as a marker pesticide. Applicable groups are those that contain at least one compound analyzed by that laboratory for that commodity. For each applicable group, a marker pesticide shall be included for each extraction/detection system used to analyze that group.

5.7.2.3 For laboratories rotating spike mixtures between analytical sets, each laboratory shall ensure that each extraction/detection system is adequately represented within each set.

5.7.2.4 For laboratories analyzing multiple commodities, a single list of marker compounds may be specified to represent all commodities. The lists of required compounds for commodities analyzed should be combined and at least one compound from each applicable group chosen to serve as a marker compound.²

5.7.3 Selected/single analyte residue studies utilize the selected analyte as the marker pesticide.

5.7.4 "Marginal Performing Analytes" are analytes that do not meet linearity, calibration integrity, ion ratio, recovery (individual or mean), or precision and accuracy criteria during method validation or continuing quality control (QC) as specified in Section 5.18. Marginal performing analytes are determined in conjunction with USDA/AMS.

5.8 Process Control Compounds

Samples analyzed by each extraction/detection system shall include the analysis of a process control compound. More than one process control may be required. The laboratory shall make every effort to choose a compound that is not expected to be an incurred residue.

 $^{^2}$ For laboratories analyzing multiple commodities, compounds in single groupings only need apply to that required commodity.

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5.9 PDP Commodity Groupings

<u>Fruits and Vegetables:</u> Apples (AP), Apple Juice (AJ), Applesauce (AC), Asparagus (AS), Avocado (AV), Baby Foods (see *Attachment* 3 for corresponding codes), Bananas (BN), Blueberries (BB), Broccoli (BR), Cabbage (CG), Canned Beans (BC), Canned Garbanzo Beans (ZB), Canned beets (BT), Cantaloupe (CN), Carrots (CR), Cauliflower (CF), Celery (CE), Cherry Tomatoes (CT), Cilantro (CL), Cranberries (CA), Cucumbers (CU), Eggplant (EP), Grapefruit (GF), Grapes (GR), Grape Juice (GJ), Green Beans (GB), Green Onions (GO), Greens (GS), Honeydew Melons (HD), Hot Peppers (HP), Kale (GK), Lettuce (LT), Mangoes (MA), Mushrooms (MU), Nectarines (NE), Canned Olives (OL), Onions (ON), Oranges (OG), Orange Juice (OJ), Papaya (YA), Canned Peaches (CC), Peaches (PC), Pears (PE), Pear Juice (PJ), Peas (PS), Canned Pineapples (NC), Pineapples (PN), Dried Plums/Prunes (PD), Plums (PU), Potatoes (PO), Raspberries (RS), Snap Peas (SN), Spinach (SP), Strawberries (ST), Summer Squash (SS), Sweet Bell Peppers (PP), Sweet Cherries (CH), Sweet Corn (CS), Sweet Potatoes (SW), Tangerines (TA), Canned Tomatoes (TC), Tomatoes (TO), Watermelon (WM), Winter Squash (WS)

<u>Cereal Grains (Low Oil)</u>: Barley (BY), Corn Grain (CO), Oats (OA), Rice (RI), Wheat (WH), Wheat Flour (WF)

Cereal Grains (High Oil): Almonds (AL), Peanut Butter (PB), Soybeans (SY),

<u>Animal Tissue/High Protein:</u> Beef (adipose – BA, liver – BL, muscle – BM), Catfish (FC), Eggs (EG), Pork (adipose – KA, muscle – KM), Poultry (adipose – PA, liver – PL, muscle – PM, breast – PR, thigh – PT), Salmon (FS)

Dairy Products: Butter (BU), Heavy Cream (CM), Milk (MK)

<u>Water:</u> Untreated Drinking Water (WU), Treated Drinking Water (WR), Bottled Water (WB), Groundwater (WG)

<u>Single Commodities:</u> For example, Corn Syrup (CY), Dairy-based Infant Formula (DF), Raisins (RA), Soy-based Infant Formula (YF), Tomato Paste (TP), Honey (HY).

5.9.1 Based on their experience with a commodity, laboratories may request changes to the assigned commodity groupings from the PDP Technical Director.

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5.9.2 Environmental Protection Agency (EPA), Codex, and Food and Drug Administration (FDA) Pesticide Analytical Manual (PAM) commodity grouping information can be found in attachments 3 and 4 of this SOP.

5.10 Establishment of LODs and LOQs

5.10.1 Method Noise

5.10.1.1Method noise is the combination of instrument noise and the matrix noise contributions.

5.10.1.2 Method noise determination must be completed for all required PDP analytes.

5.10.1.3Method noise will be determined utilizing instruments and operating conditions, which are routinely used for the analysis of samples. Noise for the LOD and LOQ calculations will be determined by examining chromatograms of the blank commodity in the chromatographic time segment of the pesticides of interest.

5.10.2 Establishment of LOD

5.10.2.1 LOD may be estimated by whatever means the laboratory chooses to employ, but the response shall be at least 3x signal to noise.

For MS systems, ions used for quantitation and for qualitative analysis/confirmation shall meet the 3x signal to noise requirement.

For example: 1) take two equal portions from the same matrix blank extract; 2) spike one aliquot with a known amount of the analyte of interest; 3) inject both aliquots under the same conditions; 4) magnify the baseline of the unfortified blank at the analyte retention time window of interest to obtain the instrument response for the tallest (height) or the broadest (area) noise; and 5) convert the response into concentration (ppm, ppb, or ppt) from the known concentration of the spiked extract. Compare the two concentrations (blank vs. spiked) to estimate the LOD.

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5.10.2.2 LODs may be established at a level greater than 3x noise.

5.10.2.3 In addition to signal-to-noise considerations, LODs estimated for zero noise instruments (e.g. triple quadrupoles) may also include consideration of replication injection data (e.g. injecting an LOD standard 10x).

5.10.2.4 The reported LOD shall be the highest value obtained using the validated method. For instance, for dual column systems, the confirmatory column LOD must be AT LEAST that of the primary/quantitative column.

5.10.2.5 For multi-peak compounds, such as many of the pyrethroids, the laboratory may base the LOD on the largest peak if a mass spectrometry system is used for both quantitation and confirmation. If other systems are used for quantitation, the laboratory may base the LOD on the larger peak if the smaller peak is <20% of the total response.

5.10.2.6 LOD is method dependent and shall be experimentally verified in matrix as detailed in Section 5.10.1.

5.10.3 Establishment of LOQ

5.10.3.1 LOQ will be calculated/determined for each analyte in each commodity tested following the establishment of LOD.

5.10.3.2 For all detection systems other than mass spectrometry, LOQ will be established by multiplying the response of method noise level by at least ten and then converting the total response into concentration (i.e., ppm, ppb, or ppt), or by multiplying the LOD by no less than ten/thirds (10/3) if the LOD is established above 3x method noise.

5.10.3.3 For mass spectrometric systems, ions to be used for qualitative analysis/confirmation shall be at least 3x signal to method noise. Ions to be used for quantitation shall be at least 10x signal to method noise.

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5.10.3.3.1 In order to maximize the number of compounds screened by MS systems while maximizing the number of scans per second and dwell times, it may be desirable to perform the initial identification and quantitation using fewer than three ions for some or all of the compounds. Presumptive-positive samples shall be re-injected or data reprocessed to meet all MS confirmation criteria.

5.10.3.4 The reported LOQ shall be the highest value obtained using the validated method.

5.11 Verification of LODs/LOQs

5.11.1 During method validation, all calculated or established LODs must be verified by fortifying duplicate blank commodities at approximately the LOD level and subjecting them to the analytical method for each extraction/detection system used in the analysis of PDP samples. In the instance where the LOD=LOQ this verification suffices for the LOD and LOQ. If method range is performed (see subsection 5.13) for verification of LOQ then section 5.11 is not required.

5.11.2 Verification consists of the observation of detectable peaks in the chromatogram at 3x the current noise level (run within the last three months). Variability is expected to be high. Therefore, recoveries can be reported as present or not present. If detectable peaks are not observed, the LOD must be re-estimated and the verification repeated.

5.11.3 Prepare summary form(s) of the acquired data for all systems and all columns used for analysis and/or confirmation (*see Attachment 5 - Method Evaluation Reporting Forms*).

5.11.4 For water only, the LOD for each reported compound shall be verified, at least every two years, by extraction of a single LOD spike. Reporting these results to MP is optional If the LOD Verification Form in Attachment 5 is used, then recording only one LOD spike is required.

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5.12 Changing LODs

5.12.1 LODs may be raised for analytes in an individual sample set at the discretion of the TPM.

5.12.2 LODs may not be lowered without verification subject to the analytical method, TPM approval, and QAO review.

5.13 Determination of Method Range

5.13.1 During method validation, samples fortified with marker compounds (only marker compounds are required, however, other compounds may be used in addition to the markers, if desired) are to be run through the entire analytical method on the primary analytical system. If more than one type of chromatography system (e.g., GC versus LC) and/or detector system (e.g., FPD versus MSD) combinations are to be used for quantification, they must be likewise evaluated.

5.13.2 Fortify samples in triplicate at approximately 1xLOQ, 5xLOQ, and 10xLOQ for each marker or compound being validated. Process these fortified samples through the entire analytical method. A reagent and matrix blank shall be subjected to the analytical method along with the fortified samples.

5.13.3 For each data point, calculate the Percent Recovery compared to known standards to three significant figures if greater than 100% or to two significant figures if less than 100%.

5.13.4 Calculate the mean Percent Recovery (%R) and Coefficient of Variation (%CV) for each level. A definition of Horwitz expected intralaboratory and interlaboratory %CVs may be found in SOP PDP-Glossary. The appropriate values may be used as a guideline when evaluating data.

5.13.5 Prepare summary form(s) of the acquired data by analyte, level, and commodity group (*see Attachment 5 - Method Evaluation Reporting Forms*).

5.13.6 Method Range Extension

If more than 20 findings per life of the commodity for a particular analyte/commodity pair exceed the highest validated spiking level, then in order to verify the ability of the method to

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extract the analyte at the higher level, the laboratory shall fortify at least one spike at or above the level of the highest finding. Reagent and matrix blanks shall accompany these spikes. If the matrix spike recoveries do not meet QC criteria (per section 5.18) any affected findings shall be coded (or recoded) as estimates. Method range extension spikes may be reported via RDE as "other" spikes. Marker pesticide spikes may be used to represent other compounds in that group. Method range extension for a given commodity can represent another commodity in that group. Laboratories may perform the range extension at various times:

- preemptively during initial validation (based on intelligence or experience with the commodity),
- in subsequent batches following the high finding
- periodically (e.g. annually) to conserve resources, or
- internal blind check samples may be used for this purpose.

Method range extension results should be reported to USDA/AMS following QA review. USDA/AMS expects any **coding changes** for calendar year samples to be submitted by May 31st following the end of the calendar year. This does not remove the requirement to report all data sets for the calendar year by March 31st of the following calendar year.

5.14 Precision and Accuracy Data Collection

5.14.1 The precision and accuracy data collection shall be compiled from the commodity groupings as specified by USDA/AMS. Each marker, single analysis, new or other required PDP analyte shall be spiked at 2xLOQ and evaluated using a minimum of seven data points, with at least two points from each commodity in the group analyzed in a particular laboratory.

5.14.2 The required data points shall be obtained from:

• 2xLOQ data points completed after Determination of Method Range

and/or

• data points from matrix spikes analyzed concurrently with samples.

These two options provide slightly different data. The second option is preferable since it provides information about the repeatability of the method over time. The first option is

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permitted when running concurrent spikes would extend the data collection over more than six months and/or concurrent spikes would make the size of sample sets unmanageable.

5.14.3 For each data point, calculate the Percent Recovery compared to known standards to three significant figures if greater than 100% or to two significant figures if less than 100%.

5.14.4 Calculate the mean Percent Recovery (%R) and Coefficient of Variation (%CV) for each pesticide using the seven data points. A definition of Horwitz expected intralaboratory and interlaboratory %CVs may be found in SOP PDP-Glossary. The appropriate values may be used as a guideline when evaluating data and/or determining whether analytes should be considered a Marginal Performing Analyte. In addition, Marginal Performing Analytes may be determined based on linearity, calibration integrity, or individual recovery values.

5.14.5 Prepare summary form(s) of the acquired data (*see Attachment 5 - Method Evaluation Reporting Forms*). Refer to Sections 5.17 for PDP acceptance criteria.

5.15 Method Evaluation Reporting

5.15.1 The methodology, method evaluation records, summary form(s), chromatograms, and any other supporting data generated during method evaluation shall be maintained by the laboratory.

5.15.2 Local Approval

5.15.2.1 Any request for and written modification of an approved analytical method shall be reviewed and approved by the QAO and TPM.

5.15.2.2 All validation documentation shall be reviewed and approved by the QAO and TPM.

5.15.3 Letter of Intent

5.15.3.1 Once the Verification of LODs and LOQs and Determination of Method Range has been completed, reviewed, and approved by the QAO and TPM, a Letter of Intent shall be submitted to the PDP Technical Director with copies to the Method

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Validation Coordinator and the assigned liaison chemist stating that these modules have been completed, reviewed, and approved and will be submitted at a later date with the Precision and Accuracy Data.

5.15.3.2 This letter shall also include a list of commodity(ies) and analyte(s) with their LOD(s) that the laboratory intends to analyze and shall be submitted within 90 days of the applicable commodity entering the program.

5.15.3.3 The Letter of Intent is not required if all required method validation data will be/is submitted within 90 days of the commodity entering the program.

5.15.3.4 USDA/AMS will perform a brief preliminary review and upon laboratory request, will issue a provisional letter of concurrence allowing the laboratory to transmit data to their liaison chemist for review while the full method validation package undergoes a multi-level review by USDA/AMS. Data may be changed, in consultation with the lab, based on the results from the full method validation package review.

5.15.4 Upon conclusion of the Precision and Accuracy Data Collection module, summary form(s) of validation documentation, and a brief narrative shall be sent to the PDP Technical Director with copies to the Method Validation Coordinator and the assigned liaison chemist with a cover memo detailing the submission (state which scenario(s) and module(s) that the submission is intended to represent). These may be sent by e-mail (preferred), hardcopy delivery (USDA/AMS PDP, 1400 Independence Ave, S.W. Washington DC 20250 or fax [(202) 619-1724].

5.15.5 A narrative accompanying the validation documentation shall include the following.

5.15.5.1 Description of the method.

5.15.5.2 Identification of any data that is only intended to be used for confirmation. Otherwise, USDA/AMS will evaluate the data as if quantitation will be performed on the instrument/analyte combination.

5.15.5.3 Requests for designation of any analytes as Marginal Performing Analytes - if USDA/AMS agrees to consider any analytes as Marginal Performing Analytes, that designation will be documented in the Letter of Concurrence.

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5.15.5.4 Identification of previous method validation data used. The laboratory shall be responsible for clearly identifying the data used and the rationale for their use. For example, if a previously validated commodity returns and the laboratory has not made any method changes and will be using the same instrumentation, the laboratory shall submit a letter to USDA/AMS explaining how the previous validation data will be used.

Example narrative for a data package:

Enclosed is the complete method validation summary of all compounds we are screening for in commodity "y" to support the addition of the commodity to the 2010 PDP program. The specific scenario used in validation was 5.6.1, New Method Implementation. Required modules included establishment and verification of LODs and LOQs, determination of method range, precision and accuracy data collection, method evaluation reporting for GC/MSD, GC/FPD, GC/XSD, and LC/MS/MS instrumentation. For compound "a", GC/FPD is the primary detection system and LC/MS/MS data is intended for confirmation purposes only. The following analytes were dropped during method development due to difficulty in analysis (e.g., solubility, poor chromatography, sensitivity, and/or loss in SPE cleanup): compound "b", compound "c", and compound "d". Due to problems with recovery, the following analytes should be considered Marginal Performing Analytes and if it is agreed, will be coded as such in reporting: compound "e", compound "f", and compound "g". If there are questions about this submission please contact: XXXXXX. All references to this submission should use QA# ###-####.

Example narrative for a previously validated returning commodity with no method, analyte, or instrumentation changes:

In 2011, commodity "y" returned to the 2011 PDP program. This commodity was previously validated in 2008 and there have been no changes to the method, target analytes, and instrumentation since then. Therefore, the 2008 validation data submitted on Month, Day, Year, is still applicable. If there are questions about this submission please contact: XXXXXX. All references to this submission should use QA# ###-####.

An example format for the submission follows:

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Title

Summary to include purpose, results, data anomalies.

Methods

Sample Preparation (example):

- 50g homogenized sample extracted with 100 ml ACN by gently mixing
- 5ml extract purified by a C-18 SPE cartridge, eluted with MeOH, and concentrated to 5ml
- 1 ml eluate further purified by florisil SPE and eluted with 5 ml 50:50 hexane/acetone
- Eluate dried down to 0.5 ml, re-suspended in acetone, and filtered
- Derivatizaton accomplished by reaction with dansyl chloride.

Analysis (example):

- Instrument GC/HPLC/detector
- Column (DB-)
- Post-column derivatization (where applicable).

5.16 Method Validation Evaluation by USDA/AMS

5.16.1 Letter of Intent

5.16.1.1 Letters of Intent shall be tracked and maintained in centralized files by the Method Validation Coordinator.

5.16.1.2 The USDA/AMS chemist assigned to that facility submitting a Letter of Intent shall review the letter and verify the submitted LOD/LOQ values against electronically submitted data (upon availability) and upon laboratory request issue a provisional letter of concurrence (see Section 5.15.3).

5.16.2 Method Validation Data Packages

5.16.2.1 After receipt by the PDP Technical Director, method validation data packages undergo a multi-tiered review by USDA/AMS. Details of this review process are specified in SOP PDP-ADMIN.

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5.16.2.2 The method validation package is reviewed to ascertain the physical presence and completeness of data submitted for method validation and to determine whether these data adhere to PDP criteria and are to be considered validated.

5.16.2.3 For data that do not meet PDP criteria for linearity, calibration integrity, ion ratios, individual or mean recovery (50-150%) or reproducibility (%CV values within the expected Horwitz intralaboratory values) USDA/AMS and the laboratory shall use scientific judgment to determine whether the compound shall be considered validated, designated as a Marginal Performing Analyte or designated as unvalidated for that pesticide/commodity pair.

Note: *The Horwitz values are used as <u>guidelines</u> only and do not preclude a compound from being considered validated.*

5.16.2.4 Once the USDA/AMS review of the method validation package has been completed, the laboratory TPM and QAO will receive a Letter of Concurrence that identifies the status of the instrument/detector results for the commodity/analyte pairing (e.g., validated, not validated, Marginal Performing Analyte, incomplete). If the data are deemed incomplete by USDA/AMS, the Letter of Concurrence will identify the deficiency and include a request for the remaining data e.g., monitoring of daily matrix fortifications or addition of a spike compound with the same functional group to the fortification profile).

5.16.2.5 Once a compound is designated as a Marginal Performing Analyte, that designation shall not be changed unless approved by the Technical Director.

5.17 Blanks and Spikes Required Per Set and Continuing QC

5.17.1 Sample set

A sample set is a group of samples, which are spiked individually with the designated process control(s), extracted with the required QC samples, and analyzed with the applicable required QC samples. Each set shall not exceed 35 samples. Required QC samples per set consist of a reagent blank, matrix blank, and matrix spike(s).

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Each laboratory is given the option of combining two or more small sets into a larger set (e.g., peaches month A + peaches month B or apples month A + peaches month A). If the larger set contains two commodities, then the set shall contain a matrix blank of each commodity and a matrix spike(s) in at least one of the commodities.

5.17.2 Reagent Blank

A reagent blank is intended to demonstrate glassware cleanliness and total system integrity. It shall be prepared by subjecting an amount of distilled water equivalent to that contained in an average sample to the entire analytical process. For consistency in the preparation of the reagent blank, it shall be assumed that an "average" (includes fresh, canned, or frozen) fruit or vegetable sample contains 80% water. If contamination or interferences in the retention time window of the pesticide of interest is present in excess of the calculated LOQ, appropriate action must be taken and documented.

5.17.3 Matrix Blank

A matrix blank is intended to demonstrate the behavior of a substrate within an analytical system. Ideally, a matrix blank should be void of any compounds of interest. A matrix blank may be a previously characterized sample of the same commodity. If a suitable sample is not available, a portion of one of the samples may be randomly selected and used as a matrix blank. If an incurred residue is found in the matrix blank, which has been chosen from the sample set, determine if the same residue is incurred in the actual sample and is not present in other samples in the same set. If this condition cannot be met, appropriate action must be taken, such as reviewing reagent blank information.

5.17.4 Matrix Spike

A matrix spike is intended to reflect the behavior of a chemical in a substrate within an analytical system. The matrix spike indicates the behavior of the chemical for the entire sample set. Analysis of a matrix spike provides valuable information on matrix interference effects as a result of the co-eluted matrix components, affecting the accuracy or detection capability for the analytes of interest.

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5.17.4.1 A second portion of the same material used for the matrix blank shall be used for the matrix spike(s). Laboratories may design their QC spiking schemes to meet their needs. A laboratory may choose to use marker groups as defined in Section 5.7 of this SOP, rotate spike mixtures between analytical sets, or spike all compounds analyzed, as long as each extraction/detection system is adequately represented within each set and the minimum requirement of all compounds reported by the laboratory to be spiked at least quarterly in each commodity, is met.

5.17.4.2 The spike shall be added prior to extraction at approximately 2x LOQ (or less). Additional spikes may be added to satisfy the quarterly spiking of each commodity with all reported compounds, as part of a validation study, or to familiarize a laboratory with pesticides that have not been previously analyzed. More than one matrix spike shall be required if necessary for all spiked compounds to be separated during the chromatographic process. If a laboratory has combined commodities within a set, then the QA/QC Recovery Form shall indicate which commodity was used for the matrix spikes. Results for all spiked compounds shall be reported to USDA/AMS through normal RDE procedures.

5.17.4.3 The matrix spike(s) shall meet the requirements specified in the criteria section below. All reported compounds (markers, required, and any other compound reported by that laboratory) shall be spiked at least quarterly for each commodity. All components of sample sets shall be subject to the same analytical process as detailed in the method SOPs.

5.17.4.4 Recoveries for compounds designated as Marginal Performing Analytes shall be coded with a "P" (Marginal Performing Analyte) in the Exception field of the QA/QC Recovery form.

5.17.4.5 If reported, recoveries for unvalidated compounds shall be coded with a "U" (Unvalidated Residue) in the Exception field of the QA/QC Recovery form.

5.17.4.6 Incurred residue levels may be subtracted from spike recovered prior to calculating the percent recovery if the conditions specified in SOP PDP-DATA are met.

5.17.5 Reporting Fortification Recoveries

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5.17.5.1 "Fresh" spikes are matrix spikes fortified, extracted, and analyzed with that set of analytical samples. Fresh values reported may be the original, re-injected, re-aliquoted, or re-extracted (from homogenate) determination value. The results reported may be the value from primary detection system or the averaged value (e.g., dual column results averaged).

5.17.5.2 "Other" spikes are additional fortifications reported by the laboratory. The laboratory can request that USDA/AMS adds a new spike type code as needed. Examples of "other spike" types are freezer, storage, failed fresh values, or "extra" QA spikes performed by the laboratory.

5.17.6 Quarterly 2xLOQ Spikes

5.17.6.1 All reported compounds (markers, required, and any other compounds reported by the laboratory) shall be spiked at least quarterly at 2x LOQ (or less) for each commodity.

5.17.6.2 The laboratory may choose to rotate spikes on a regular basis as long as the requirements in Subsection 5.17.4.1 are met.

5.17.6.3 The spike results shall be reported to USDA/AMS via RDE (the preferred option) or in Excel spreadsheets. Results shall also be addressed in the semi-annual QA Reports submitted to USADA/AMS.

5.17.7 Process Control Spikes

A process control spike is intended to assure the integrity of a particular sample within an analytical system.

5.17.7.1 Each sample set component, except the reagent and matrix blanks, shall be spiked with a process control at approximately 5x the Limit of Quantitation (LOQ) prior to the extraction step of the analytical procedure. However, if the intent of the process control is to monitor the percent recovery of a clean-up step, or of a derivatization, then the process control shall be added to the extract before the clean-up or derivatization step.

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5.17.7.2 The laboratory shall make an effort to choose a compound that is not expected to be an incurred residue. The value reported as "percent recovery" may be the original, re-injected, re-aliquoted, or re-extracted (from homogenate) determination value [either value from primary detection system or averaged value (e.g., dual column results averaged)].

5.17.8 QA/QA Recovery Form Codes

The following codes shall be entered in the Exception field of the QA/QC Recovery form. See Section 5.17.4 for additional details.

Code	QA Spike Exception
Е	Estimated
Ι	Incurred Residue
Μ	Matrix Interference
Ν	Not Recovered
Р	Marginal Performing Analyte
S	Incurred Residue Subtracted
U	Unvalidated Residue

5.18 Criteria for Method Validation and Continuing QC

5.18.1 Method Validation Criteria

5.18.1.1 PDP criteria for percent recovery for determination of method range and precision and accuracy data collection is 50-150%.

5.18.1.2 Horwitz intralaboratory values are used as a guideline for determining reproducibility acceptability. The laboratory shall indicate any compounds that they feel are not acceptable and/or those that should be classified as Marginal Performing Analytes. These laboratory recommendations are subject to approval by USDA/AMS.

5.18.1.3 Some analytes may not meet method validation criteria for linearity, calibration integrity, ion ratios, recovery (individual or mean), or precision (%CV). Rather than not including them in the laboratory's screening list, USDA/AMS and the

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laboratory may decide that marginal data are preferable to no data. These compounds shall be designated as Marginal Performing Analytes. Details on USDA/AMS review of method validation data can be found in Section 5.15.

5.18.2 Matrix Spike Criteria

5.18.2.1 All spiked compounds shall have recoveries between 50 and 150%, within the statistically calculated range, or within a range agreed upon with USDA/AMS.

5.18.2.2 If a large number of analytes are in the spike, it becomes statistically likely that a few will be outside control limits. This may not indicate that the system is out of control. The laboratory shall have written criteria for when corrective action(s) will be necessary.

5.18.2.2.1 Some analytes may not be optimally recovered during method validation trials. Recoveries may be low and/or erratic and rather than not including them in the laboratory's screening list, the laboratory may consult with USDA/AMS to determine if marginal data may be preferable to no data. If reported by the laboratory, the codes for Marginal Performing Analytes shall be utilized. USDA/AMS will note the use of Marginal Performing Analytes in the Letter of Concurrence and the use of marginal performer codes for particular analyte/commodity pairs. Once a compound is designated as a Marginal Performing Analyte, that designation shall not be changed unless approved by the Technical Director.

5.18.2.2.2 Some analytes that behave acceptably during method validation may behave unacceptably during the analysis of routine batches. This may be due to the fact there is more commodity variability among actual samples than there is in the limited matrix utilized for method validation batches. As above, rather than dropping these analytes from the screening list, the laboratory should consult with USDA/AMS to determine if they should be reclassified as Marginal Performing Analytes. If a compound is reclassified as a Marginal Performing Analyte, an e-mail notification to the Technical Director, with a copy to the USDA/AMS liaison, shall be sent and

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approved/acknowledged by USDA/AMS, and that designation shall not be changed unless an e-mail communication is sent by the Technical Director reversing the previous approval.

5.18.3 Response To Failure To Meet Matrix Spike Criteria Range

If a spike analyte fails, even after re-injection/re-aliquoting, it can and should be reported, because the recovery may reflect normal random variation inherent to pesticide residue analysis. For high recoveries, this practice is defensible. For low recoveries, best professional judgment should be used, although if recovery is 0%, that analyte should be reported as unable to detect in samples.

When a spiked pesticide recovery falls outside the range criteria, any one of the following options, or combination thereof, may be chosen by the TPM or designee. (See Attachment 6 – Matrix Spike and Process Control Recovery Acceptability Flowchart.)

5.18.3.1 The original extract may be re-injected or re-aliquoted. If the spiked pesticide recovery falls within the range criteria, then the results from the re-injected extract shall be reported.

5.18.3.2 The sample set may be re-extracted from the frozen homogenate. If the spiked pesticide recovery falls within the range criteria, the rerun results shall be reported.

5.18.3.3 The original results may be reported with an explanation (e.g., recovery exceed 150% but all samples in the set are non-detects for that analyte; wrong mix spiked; spike spilled but process controls in samples are acceptable; control charts indicate a recurrent analyte/matrix; etc.) The TPM and QAO shall ensure that reported data is not compromised and the explanation shall be conveyed to headquarters (e.g., note in RDE, email message to USDA/AMS liaison chemist and Technical Director).

5.18.3.4 Other options may be acceptable depending on the outcome of investigations and/or consultations with USDA/AMS. An explanation shall be conveyed to headquarters.

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5.18.4 Process Control Criteria

Each laboratory shall decide whether to use the Absolute Range Criteria or the Statistically Calculated Range Criteria. A laboratory may choose different Range Criteria for different test types, but it is intended that a laboratory stay with the chosen criteria unless approved by the laboratory QAO.

5.18.4.1 Absolute Range

Each process control recovery shall fall between 50-150% for all detection systems used to calculate sample data.

5.18.4.2 Statistically Calculated Range

The mean recovery for a sample set's process control shall be calculated. Each process control recovery shall fall within its acceptance recovery range, which is the mean recovery plus and minus three standard deviations.

5.18.5 Response To Failure To Meet Chosen Process Control Criteria Range

If a process control fails, even after re-injection/re-aliquoting/re-extraction, the results may be reported, based on best professional judgment.

When a process control falls outside the chosen range criteria, any one of the following options, or combination thereof, may be chosen by the TPM or designee. (See Attachment 6 - Process Control and Spike Recovery Acceptability Flowchart.)

5.18.5.1 The original extract may be re-injected or re-aliquoted. If the process control recovery falls within the chosen range criteria, then the results from the re-injected or re-aliquoted extract shall be reported.

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5.18.5.2 The sample may be re-extracted from the frozen homogenate. If the process control recovery falls within the chosen range criteria, the re-run results shall be reported.

5.18.5.3 The original results may be reported with an explanation (e.g., pipette error, the PC recovery exceeds 150% but all analytes in the sample are non-detects, etc.). The TPM and QAO shall ensure that reported data is not compromised and the explanation shall be conveyed to headquarters (e.g., note in RDE, email message to USDA/AMS liaison chemist and Technical Director).

5.18.5.4 Other options may be acceptable depending on the outcome of investigations and/or consultations with USDA/AMS. An explanation shall be conveyed to headquarters.

5.18.6 Evaluation of Recoveries

Laboratories shall use control charting or other appropriate statistical tools to evaluate recoveries on a set-to-set basis and monitor trends over time.

5.19 Proficiency Testing

5.19.1 PDP PT Program Overview

5.19.1.1 PDP Fiscal Year (FY) PT program schedules are posted to the PDP Extranet site and are referenced in the applicable PDP Semi-Annual Program Plans.

5.19.1.2 General multi-residue method samples for fruit and vegetables will be supplied by the Food Analysis Performance Assessment Scheme (FAPAS) and the California Department of Food and Agriculture (CDFA).

5.19.1.4 Rounds for commodities other than fruit and vegetables (e.g., meat, milk and dairy products, fish, grains, nuts, etc.) shall be supplied by CDFA. Additionally, applicable FAPAS rounds may be scheduled.

5.19.1.5 PT samples received may be significantly larger than the analytical portion required by the laboratory for analysis. In cases where the PT sample is more than twice the analytical weight needed, the laboratory may subsample duplicate portions for

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extraction and analysis as described below, due to the uncertainty regarding homogeneity of samples. Sample results that meet the QC criteria shall be averaged for reporting.

5.19.1.5.1 Samples shall be mixed in the container they came in, taking care to not spill any sample prior to subsampling.

5.19.1.5.2 Two sub-samples shall be weighed out for extraction and extracted as separate samples.

5.19.1.5.3 Each extract shall be analyzed as an individual sample

5.19.2 Reporting PT Results

5.19.2.1 For FAPAS, lists of potentially spiked pesticides are available from the provider websites. Rounds issued by CDFA are designed to focus only on those compounds validated by the applicable laboratory(ies). For all rounds, participants shall only be evaluated for those residues validated by their laboratory and not declared as Marginal Performing Analytes. The report provided will clearly identify these pesticides. Reporting of the Marginal Performing Analytes shall be optional.

5.19.2.2 For FAPAS, it is recognized that a laboratory may not have validated the commodity scheduled for that specific round. Standards used in routine analyses of assigned commodities should be used. Efforts will be made to provide a matrix blank for each round.

5.19.2.3 Report results according to provider instructions and requirements. Reporting to USDA/AMS via RDE is optional.

5.19.2.4 For FAPAS, LOD/LOQ and recovery values reported may be values obtained from previous routine batches of the laboratory's usual commodity(ies).

5.19.2.5 Reports for each round shall be posted to the PDP Extranet within 10 working days of receipt by USDA/AMS.

5.19.3 Laboratory Response

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5.19.3.1 Upon receipt of PDP PT results, laboratories shall review results and initiate corrective actions when they are considered unacceptable by the PT scheme provider.

5.19.3.2 Where FAPAS is the provider, z-scores whose absolute values are greater than 3 are unsatisfactory.

Note: *FAPAS'* assigned/target value is the consensus value of the submitted results (with appropriate exclusions as noted in the FAPAS reports) and the target standard deviation is determined based on Horwitz.

5.19.3.3 For rounds provided by CDFA, unacceptable results shall be defined as those outside 50-150% recovery, or outside the statistically calculated range defined as $\pm 3xSD$ of the mean of last 20 data points of the laboratory's spike recovery for the compound, or outside a range agreed upon with USDA/AMS. Unvalidated or Marginal Performing Analytes need not meet these criteria, but should be addressed in the PT section of the semi-annual QA report.

5.19.3.4 If any corrective actions are initiated due to the results, USDA/AMS shall be informed within 30 days. Refer to SOP PDP-ADMIN for notification details.

5.20 Measurement Uncertainty

Measurement uncertainty shall be determined on an annual (calendar year) basis by USDA/AMS. USDA/AMS will calculate each year's value using 2x the standard deviation of program recovery data reported with each analytical data set. For example, during calendar year 2003, the mean program matrix spike recovery was 92% and the standard deviation was 26%. Results for 2003 would be expressed as "value \pm 52%." USDA/AMS will be responsible for communicating program measurement uncertainty values to data users.

USDA/AMS does not require individual PDP laboratories to report their measurement uncertainty along with sample results.

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ORIGINAL SIGNATURE PAGE MAINTAINED BY USDA, AMS, SCIENCE & TECHNOLOGY, MONITORING PROGRAMS DIVISION ELECTRONICALLY REPRODUCED SIGNATURE

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Revision 8February 2018Monitoring Programs Division

- Updated the heading for section 5.11 throughout the document
- Updated guidance to section 5.11 for LOQ verification
- In section 5.13.2 replaced "single PDP analyte" with "compound being validated"
- Updated Attachment 2 by adding the following compounds: aspon, aclonifen, anilofos, atraton, beflubutamid, bioallethrin, bromophos ethyl, chlordimeform, chlorthiophos, desmetryn, dichlofenthion, dimepiperate, dipropetryn, ditalimfos, diphacinone, dithiopyr, dioxacarb, etrimfos, famphur, fluensulfone, flazasulfuron, flucythrinate, fluorodifen, iprobenfos, isoprothiolane, mephosfolan, methacrifos, mefenacet, simetryn, prodiamine, pretilachlor, pyraclofos, pyridaphenthion, terbutryn, trichloronate
- Added commodity codes for canned tomatoes, canned garbanzo beans, canned peaches, dried plum/prunes and honey to section 5.9
- Updated attachment 3 with new commodities: canned peaches and dried plum/prunes

Rev	ision 7	February 2017	Monitoring Programs Division
• (Jpdated Attachment 2 by a	dding the following compounds: 1,3	3 dichloropropene, 2,6 DIPN, 5-(4-
С	hlorophenyl) oxazole-2-pr	opionic acid (CPOPA), acetamide, a	allidochlor, amicarbazone,
a	minopyralid, asulam, barb	an, chlorpyrifos methyl O-analog, cl	loquintocet methyl, cloquintocet
n	nexyl, cumyluron. cyclanil	ide, dinocap, ethephon, flumetralin,	flupyradifurone, glufosinate,
h	hexaflumuron, isofetamid, kasugamycin, melamine, merphos, metrafenone, niclosamide,		
C	oxythioquinox, perthane, pi	peralin, prohexadione calcium, prop	poxycarbazone, proquinazid,
p	prothioconazole, quinchlora	c, thidiazuron, thiencarbazone meth	nyl
• F	Removed oxytetracycline fi	rom Attachment 2	
• /	Added commodity codes fo	r canned olives kale canned ninear	onle to section 5.9

- Added commodity codes for canned olives, kale, canned pineapple to section 5.9
- Updated section 4: link to Codex Alimentarius Commission, Pesticide Residues in Food and Feed
- Updated attachment 3 with new commodities: canned pineapples and canned olives
- Updated attachment 4 with new commodity: canned olives

R	evision 6	February 2016	Monitoring Programs Division
•	Updated Attachment 2 by adding t	he following compound	s: acequinocyl, ametoctradin, benalaxyl,
	benazolin, BHC-delta, BHC-epsile	on, bifenox, carbophenot	hion methyl, chlorobenzilate, cloransulam
	methyl, cyantraniliprole, cyflufenamid, cyflumetofen, cyprosulfamide, dichlormid, diclosulam,		
	diethofencarb, diniconazole, EPN, ethiofencarb sulfone, ethiofencarb sulfoxide, ethiprole, ethylan,		
	etofenprox, fenoxycarb, fenpropid	in, fenpyrazamine, fenth	ion sulfone, fenthion sulfoxide, fluazifop,

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flufenpyr ethyl, flumiclorac pentyl, fluopyram, fluthiacet methyl, fomesafen, furalaxyl, heptenophos, imazosulfuron, ipconazole, isocarbophos, isofenphos methyl, isoprocarb, isoproturon, isoxadifen ethyl, mecarbam, mefenpyr diethyl, mesotrione, metolcarb, monolinuron, nitrofen, penflufen, phorate OA sulfone, phorate OA sulfoxide, picoxystrobin, profluralin, profoxydim, pyraflufen, pyroxasulfone, quizalofop, rotenone, sedaxane, sulfallate, sulfoxaflor, terbufos OA sulfone, terbufos sulfoxide, thionazin, tolfenpyrad, topramezone, and tricyclazole

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- Included additional matrix spike language to sections 5.17.4 and 5.18.2
- Removed references to AOAC PTs in section 5.19

Revision 5August 2014Monitoring Programs Division• Updated references to USDA/AMS throughout document

- Updated prioritization protocols in sections 5.1.2.1, 5.1.2.2.2, 5.1.2.2.3, and 5.1.2.2.4
- Changed standard checking requirement to 15% RPD in sections 5.2.6, 5.2.6.2, 5.2.7.2.1, and 5.2.7.2.2
- Combined laboratory Letter of Intent requirements into section 5.15.3
- Updated USDA/AMS address in section 5.15.4
- Updated USDA/AMS Letter of Intent procedures in section 5.16.1

Revision 4 July 2013 M	Ionitoring Programs Division
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- Updated references to USDA/AMS throughout document
- Added commodity codes for avocado, catfish, dairy-based infant formula, raspberries, salmon, and soy-based infant formula to section 5.9
- Added procedures for subsampling PT samples to section 5.19.1.5
- Updated Attachment 2 by adding the following compounds: acrinathrin, AMPA, aviglycine HCl, bromopropylate, bupirimate, butocarboxim, butocarboxim sulfone, butocarboxim sulfoxide, chlorsulfuron, chlozolinate, clethodim 5 hydroxy sulfone, clethodim sulfone, clethodim sulfoxide, clofencet, crotoxyphos, crufomate, demeton-S, demeton-S sulfone, dichlofluanid, DMST,fenbutatin oxide, fenchlorphos, fenpropimorph, fensulfothion, fenthion o-analog, fipronil sulfone, fluquinconazole, flusilazole, flutriafol, fluxapyroxad, glyphosate, haloxyfop, iodosulfuron methyl, lenacil, lufenuron, mesosulfuron methyl, metaflumizole, methiocarb sulfone, methiocarb sulfoxide, oxytetracyline, paclobutrazol, penconazole, pencyuron, penthiopyrad, phoxim, pirimicarb desmethyl, primisulfuron, propaquizafop, prosulfuron, tebufenpyrad, teflubenzuron, terbuthylazine, thifensulfuron methyl, thymol, toxaphene, and tribenuron methyl
- Updated Attachments 3 and 4 for the following commodities: avocado, catfish, infant formula (dairybased and soy based), raspberries, and salmon

Revision 3	March 2012	Monitoring Programs Division
		0 0

- Updated prioritization rationale in section 5.1.2
- Added exemption for CoA and current expiration date for revoked compounds to section 5.2.1.2

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- Defined extraction/detection system in section 5.4.3
- Clarified process control compound requirements in section 5.8
- Combined PDP Commodity Groupings in section 5.9
- Added guidance for LOD establishment for zero noise instruments in section 5.10.2.2
- Changed multi-peak compound LOD requirements in section 5.10.2.5
- Changed LOD verification requirements in section 5.11.4
- Specified PT samples larger than routine analytical samples may be run in duplicate in section 5.19.1.5
- Clarified what constitutes unacceptable PT scores in section 5.19.3.2
- Updated Attachment 2 by adding the following compounds: 2,4-DMPF, DEET, Dialofos, Dioxathion, Endothall, Indaziflam, Leptophos o-analog, Metconazole, Quinalphos, Saflufencil, Triazophos

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Revision 2
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July 2011

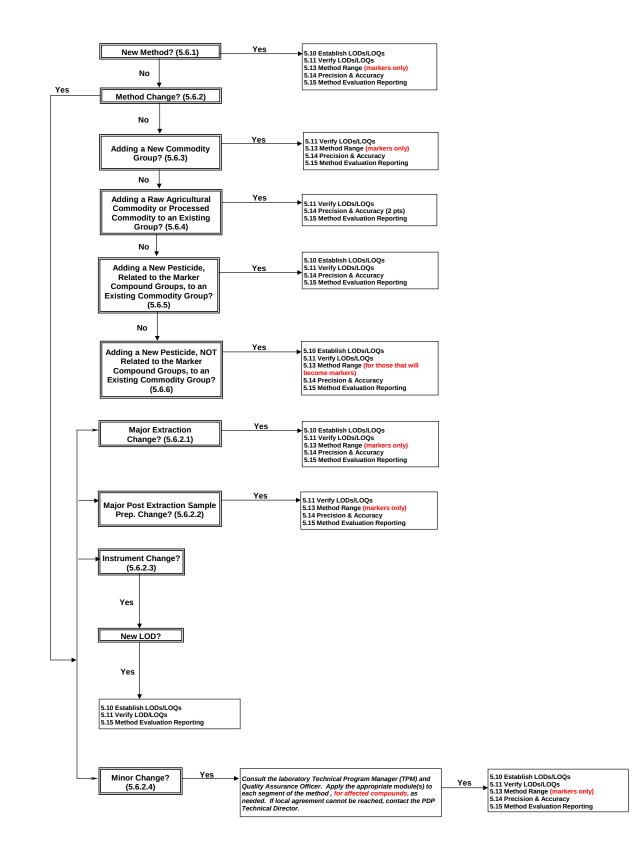
Monitoring Programs Division

- Updated the entire SOP with the new program's name Monitoring Programs Division or MP instead of MPO
- Updated the Reference list.
- Removed the word "Note" in sections 5.2.1.2; 5.6.2.3; 5.6.4; 5.10.2; 5.20 leaving the paragraphs as instructions
- Updated section 5.2.7 by renumbering the subsections
- Added new commodities (Baby Foods, Papaya, Tangerines, Cherry Tomatoes, Snap Peas, Canned Beets) to 5.9
- Updated sections 5.15.5.3, 5.16.2.5 about approving MPAs
- Added the E code in section 5.17.8
- Updated requirements for section 5.18.2 by eliminating subsection 5.18.2.1
- Updated sections 5.18.2.2.1, 5.18.2.2.2 about MPAs (re)designation, replacing the letter of deviation with e-mail communication
- Updated section 5.18.2.2.2 by replacing the letter of deviation requirement with an e-mail communication
- Updated sections 5.19.1 and 5.19.2, by replacing "Ultra/GLEC" with "Ultra"
- Updated section 5.19.3.1 with new FAPAS requirements regarding z-scores
- In section 5.19.1.3 replaced "collected by GLEC" with "provided by MP"
- Updated Attachment 2 by adding a Group 4, Benzothiazoles/triazolones, to 'PDP Compound Groups for Fruit and Vegetables' list
- Updated Attachment 2 by adding the following compounds: Fosthiazate, Iprovalicarb, Rimsulfuron, Trifloxysulfuron, Uniconazole to PDP Pesticides Codes list
- Updated Attachment 3, 4 with new commodities: Baby Foods, Papaya, Tangerines, Cherry Tomatoes, Snap Peas, Canned Beets

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Revision 1	July 2010	Monitoring Programs Office

- Renumbered the entire SOP replacing the sections' letters with numbers.
- Redefined Priority 1-4 compounds.
- Updared section 5.2.8 for Working Dilutions/Mixed Standards.
- Changed and updated section 5.5 Method Validation Evaluation Guidelines.
- Changed and updated section 5.6 Method Validation Scenarios.
- Updated section 5.7 Marker Pesticides to remove mandatory markers.
- Updated section 5.9 PDP Commodity Groupings.
- Removed section 5.12 LOD Check.
- Added section 5.13.6 Method Range Extension.
- Updated section 5.15.3 as part of Method Evaluation Reporting.
- Removed section 5.16.e.4
- Removed section 5.18.d.1.b
- Updated section 5.19.c (now 5.18.2) Matrix Spike Criteria.
- Updated section 5.19.e (now 5.18.3) Response to Failure To Meet Matrix Spike Criteria Range.
- Updated section 5.19.d (now 5.18.5) Response to Failure To Meet Chosen Process Control Criteria Range.
- Updated Attachment 1.
- Updated Attachment 2, by adding the following new compounds: Avermectin B₁, Bensulide oxygen analog, Cyhalofop butyl, Dimethipin, Disulfoton oxygen analog, Disulfoton oxygen analog sulfone, Disulfoton oxygen analog sulfoxide, Eprinomectin, Fenobucarb (BPMC), Flubendiamine, Flufenoxuron, Fluopicolide, Imidacloprid urea, Mandipropamid, Metaldehyde, Milbemectin, Pinoxaden, Promecarb, Prothioconazole, Pyrasulfotole, Pyridalyl, Tepraloxydim.
- Updated Attachments 3, 4 and 5.



PDP Compound	Groups for Fruit	and Vegetables
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Group	Description
1	Phthalimides, conazoles and metabolites, carbamaldehydes, phenyl pyrroles, methoxy-acetamides, and neonicotinyls
2	Cyano/nitrile group(s) attached to double bond
3	Halogenated aromatics and chlorinated cyclics/cyclodienes
4	Benzothiazoles/triazolones
7	Dinitroanilines
8	Pyrethroids and metabolites and synergists
9	Triazines
11	Organophosphates and metabolites
14	Carbamates, thiocarbamates and metabolites
16	Uracils/ureas, imidazolinones, diacylhydrazines, and sulfonyl ureas
17	Nitrogenous heterocyclics
20	Phenoxy acids, ethanesulfonic acids (ESA), and oxanilic acids(OA)
21	Oxyhydrocarbons
22	Strobilurins
27	Tetronic acids
28	Cyclohexenone oxime
29	Macrocyclic lactones
99	Single

Note: Missing group numbers are attributed to the consolidation of groups. For example, Group 15, Thiocarbamates, was consolidated into Group 14, Carbamates.

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code
1-naphthol	90-15-3	C ₁₀ H ₈ O	carbamate metabolite	14	382
1,2,4-triazole	288-88-0	$C_2H_3N_3$	triazole metabolite	1	A68
1,3-dichloropropene	542-75-6	$C_3H_4Cl_2$	fumigant nematicide		251
2,4-DB	94-82-6	$C_{10}H_{10}CI_2O_3$	phenoxy acid	20	317
2,4-D	94-75-7	$C_8H_6Cl_2O_3$	phenoxy acid	20	026
2,4-dimethylphenyl formamide (DMPF)	60397-77-5	$C_9H_{11}NO$	amidine	2	AGR
2,4,5-T	93-76-5	$C_8H_5CI_3O_3$	phenoxy acid	20	312
2,6-DIPN	24157-81-1	$C_{16}H_{20}$	substituted naphthalene	99	AFZ
3-hydroxycarbofuran	16655-82-6	$C_{12}H_{15}NO_4$	carbamate metabolite	14	512
4-dimethylaminosulphotoluidide (DMST); tolylfluanid metabolite	66840-71-9	$C_9H_{14}N_2O_2S$	phenylsulfamidemetabolite	1	AJU
5-hydroxythiabendazole	948-71-0	$C_{10}H_8N_3OS$	carbamate	1	B28
Abamectin	71751-41-2	$C_{48}H_{72}O_{14} + C_{47}H_{70}O_{14}$	avermectin (macrocyclic lactone)	29	948
Acephate	30560-19-1	$C_4H_{10}NO_3PS$	phosphoramidothioic acid	11	204
Acequinocyl	57960-19-7	$C_{24}H_{32}O_4$	unclassified acaricides	99	AKS
Acetamide	60-35-5	C ₂ H ₅ NO	amide	1	AAT
Acetamiprid	160430-64-8	$C_{10}H_{11}CIN_4$	neonicotinyls	1	B80
Acetochlor	34256-82-1	$C_{14}H_{20}CINO_2$	chloroacetanilide	1	807
Acetochlor ethanesulfonic acid	187022-11-3	$C_8H_{21}NO_5S$	chloroacetanilide metabolite	20	ABN
Acetochlor oxanilic acid	194992-44-4	$C_{14}H_{19}NO_4$	chloroacetanilide metabolite	20	ABO
Acibenzolar-S-methyl	135158-54-2	$C_8H_6N_2OS_2$	thiadiazole	1	B51
Acifluorfen	50594-66-6	$C_{14}H_{21}NO_5S$	diphenyl ether	3	727
Aclonifen	74070-46-5	$C_{12}H_9CIN_2O_3$	nitrophenyl ether	3	D58
Acrinathrin	103833-18-7	$C_{26}H_{21}F_6NO_5$	pyrethroid	8	A03
Alachlor	15972-60-80	$C_{14}H_{20}CINO_2$	acetamide	1	227
Alachlor ethanesulfonic acid	142363-53-9	$C_8H_{21}NO_5S$	chloroacetanilide metabolite	20	ABP
Alachlor oxanilic acid	171262-17-2	$C_{14}H_{19}NO_4$	chloroacetanilide metabolite	20	ABQ
Aldicarb	116-06-3	$C_7H_{14}N_2O_2S$	carbamate	14	167

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code
Aldicarb sulfone	1646-88-4	$C_7H_{14}N_2O_4S$	carbamate	14	168
Aldicarb sulfoxide	1646-87-3	$C_7H_{14}N_2O_3S$	carbamate	14	169
Aldrin	309-00-2	$C_{12}H_8CI_6$	cyclodiene	3	001
Allethrin	584-79-2	$\mathrm{C_{19}H_{26}O_{3}}$	pyrethroid	8	002
Allidochlor	93-71-0	$C_8H_{12}CINO$	amide	1	768
Ametoctradin	865318-97-4	$\mathrm{C_{15}H_{25}N_5}$	triazolopyrimidine fungicides	1	АКС
Ametryn	834-12-8	$C_9H_{17}N_5S$	triazine	9	156
Amicarbazone	129909-90-6	$C_{10}H_{19}N_5O_2$	triazolone	4	AGK
Aminomethylphosphonic acid (AMPA)	1066-51-9	CH ₆ NO ₃ P	organophosphate metabolite	99	957
Aminopyralid	150114-71-9	$C_6H_4Cl_2N_2O_2$	pyradine	20	AGO
Amitraz	33089-61-1	$\mathrm{C_{19}H_{23}N_{3}}$	amidine	2	233
Anilazine	101-05-3	$C_9H_5CI_3N_4$	triazine	9	033
Anilofos	64249-01-0	$C_{13}H_{19}CINO_3PS_2$	organophosphate	11	D62
Aspon	3244-90-4	$C_{12}H_{28}O_5P_2S_2$	organophosphate	11	816
Asulam	3337-71-1	$C_8H_{10}N_2O_4S$	sulfonamide	14	ANG
Atraton	1610-17-9	$C_9H_{17}N_5O$	methoxytriazine	9	D64
Atrazine	1912-24-9	$C_8H_{14}CIN_5$	triazine	9	305
Avermectin B ₁	71751-41-2	$C_{48}H_{72}O_{14}$ (avermectin B_{1a}) + $C_{47}H_{70}O_{14}$ (avermectin B_{1b})	macrocyclic lactone	29	AHQ
Aviglycine HCl	55720-26-8	$C_6H_{12}N_2O_3$	ethylene inhibitors	99	AKT
Azinphos ethyl	2642-71-9	$\mathrm{C_{12}H_{16}N_{3}O_{3}PS_{2}}$	organophosphate	11	547
Azinphos methyl	86-50-0	$C_{10}H_{12}N_{3}O_{3}PS_{2}$	benzotriazine	11	042
Azinphos methyl oxygen analog	7643-80-3	$C_{10}H_{12}N_3O_4PS$	oxon	11	769
Azoxystrobin	997888-88-8	$C_{22}H_{17}N_{3}O_{5}$	strobilurin	22	B48
Barban	101-27-9	$C_{11}H_9CI_2NO_2$	carbanilate	14	716

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code
Benalaxyl	71626-11-4	C ₂₀ H ₂₃ NO ₃	anilide fungicides	1	B45
Benazolin	3813-05-6	C ₉ H ₆ CINO ₃ S	benzothiazole herbicides	4	832
Bendiocarb	22781-23-3	$C_{11}H_{13}NO_4$	carbamate	14	658
Beflubutamid	113614-08-7	$\mathrm{C_{18}H_{17}F_4NO_2}$	amide	1	D69
Benfluralin	1861-40-1	$\mathrm{C}_{13}H_{16}F_3N_3O_4$	dinitroaniline	7	191
Benomyl	17804-35-2	$\mathrm{C}_{14}H_{18}N_4O_3$	benzimidazole	14	192
Benoxacor	98730-04-2	$C_{11}H_{11}CI_2NO_2$	benzoxazine	1	A05
Bensulfuron methyl	83055-99-6	$\mathrm{C_{16}H_{18}N_4O_7S}$	sulfonyl urea	16	ABR
Bensulide	741-58-2	$C_{14}H_{24}NO_4PS_3$	organophosphate	11	239
Bensulide oxygen analog	20243-81-6	$C_{14}H_{24}NO_4PS_3$	organophosphate	11	740
Bentazon	25057-89-0	$C_{10}H_{12}N_2O_3S$	thiadiazinone dioxide	17	758
Benthiavalicarb-isopropyl	177406-68-7	$C_{15}H_{18}FN_3O_3S$	benzothiazole	4	AGP
BHC alpha	319-84-6	$C_6H_6CI_6$	hexane ring	3	903
BHC beta	319-85-7	$C_6H_6Cl_6$	hexane ring	3	904
BHC, delta	319-86-8	$C_6H_6Cl_6$	hexane ring	3	905
BHC, epsilon	6108-10-7	C ₆ H ₆ Cl ₆	hexane ring	3	ALH
Bifenazate	149877-41-8	$C_{17}H_{20}N_2O_3$	hydrazine carboxylate	14	B82
Bifenox	42576-02-3	$C_{14}H_9Cl_2NO_5$	nirophenyl ether herbicides	3	728
Bifenthrin	82657-04-3	$C_{23}H_{22}CIF_3O_2$	pyrethroid	8	930
Bioallethrin	260359-57-7	$C_{19}H_{26}O_3$	pyrethroid	8	ANP
Bitertanol	55179-31-2	$C_{20}H_{23}N_{3}O_{2}$	triazole	1	850
Boscalid	188425-85-6	$C_{18}H_{12}CI_2N_2O$	anilide/pyridine	1	B75
Bromacil	314-40-9	$C_9H_{13}BrN_2O_2$	uracil	16	153
Bromophos ethyl	4824-78-6	$C_8H_8BrCl_2O_3PS$	organophosphate	11	602
Bromopropylate	18181-80-1	$\mathrm{C_{17}H_{16}Br_2O_3}$	bridged diphenyl	3	523
Bromoxynil	1689-84-5	$C_7H_3Br_2NO$	phenol	20	729

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code
Bromuconazole-46	NA ^[2]	$C_{13}H_{12}BrCl_2N_3O$	conazole	1	ADU
Bromuconazole-47	NA ^[2]	$C_{13}H_{12}BrCl_2N_3O$	conazole	1	ADV
Bupirimate	41483-43-6	$\mathrm{C_{13}H_{24}N_4O_3S}$	pyrimidine	17 or 3	872
Buprofezin	69327-76-0	$\mathrm{C_{16}H_{23}N_{3}OS}$	thiadiazinone	17	B52
Butachlor	23184-66-9	$C_{17}H_{26}CINO_2$	chloroacetanilide	1	806
Butocarboxim	34681-10-2	$C_7H_{14}N_2O_2S$	oxime carbamate	14	857
Butocarboxim sulfone	NA ^[2]	$C_7H_{14}N_2O_4S$	oxime carbamate metabolite	14	AKN
Butocarboxim sulfoxide	34681-24-8	$C_7H_{14}N_2O_3S$	oxime carbamate metabolite	14	АКО
Butylate	2008-41-5	$C_{11}H_{23}NOS$	thiocarbamate	14	783
Cadusafos	95465-99-9	$C_{10}HOPS_2$	phosphorodithionate	11	953
Captafol	2939-80-2	$\mathrm{C_{10}H_9Cl_4NO_2S}$	phthalimide	1	174
Captan	133-06-2	$C_9H_8CI_3NO_2S$	phthalimide	1	011
Carbaryl	63-25-2	$C_{12}H_{11}NO_2$	carbamate	14	102
Carbendazim	10605-21-7	$C_9H_9N_3O_2$	benzimidazole	14	666
Carbofuran	1563-66-2	$C_{12}H_{15}NO_3$	carbamate	14	180
Carbophenothion	786-19-6	$\mathrm{C}_{11}H_{16}CIO_{2}PS_{3}$	organophosphate	11	202
Carbophenothion methyl	953-17-3	$C_9H_{12}CIO_2PS_3$	organophosphate	11	AGZ
Carboxin	5234-68-4	$C_{12}H_{13}NO_2S$	carboxamide	1	210
Carfentrazone ethyl	128639-02-1	$\mathrm{C_{15}H_{14}Cl_2F_3N_3O_3}$	fluorophenyl triazole	4	B21
Chloramben	133-90-4	$C_7H_5Cl_2NO_2$	benzoic acid	20	952
Chlorantraniliprole	500008-45-7	$\mathrm{C_{18}H_{14}BrCl_2N_5O_2}$	diamide; pyrazole	1	AGW
Chlordane cis	5103-71-9	$C_{10}H_6Cl_8$	cyclodiene	3	173
Chlordane trans	5103-74-2	$C_{10}H_6CI_8$	cyclodiene	3	172
Chlordimeform	6164-98-3	$C_{10}H_{13}CIN_2$	formamidine	3	278
Chlorethoxyfos	54593-83-8	$C_6H_{11}CI_4O_3PS$	phosphorothioate	11	A15
Chlorfenapyr	122453-73-0	$\mathrm{C_{15}H_{11}BrClF_{3}N_{2}O}$	pyrrole	1	B13
Chlorfenvinphos total	470-90-6	$C_{12}H_{14}CI_3O_4P$	organophosphate	11	AAK

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code
Chlorimuron ethyl	90982-32-4	$C_{15}H_{15}CIN_4O_6S$	sulfonyl urea	16	717
Chlorobenzilate	510-15-6	$\mathrm{C_{16}H_{14}Cl_{2}O_{3}}$	bridged dipehnyl	3	015
Chloroneb	2675-77-6	$C_8H_8Cl_2O_2$	chlorobenzene	3	196
Chlorothalonil	1897-45-6	$C_8Cl_4N_2$	phthalimide	2	164
Chlorpropham	101-21-3	$C_{10}H_{12}CINO_2$	carbamate	14	114
Chlorpyrifos	2921-88-2	$C_9H_{11}CI_3NO_3PS$	phosphorothionic acid	11	160
Chlorpyrifos methyl	5598-13-0	$C_7H_7CI_3NO_3PS$	phosphorothionic	11	235
Chlorpyrifos methyl O-analog	5598-52-7	$C_7H_7CI_3NO_4P$	oxon	11	AAZ
Chlorpyrifos oxygen analog	5598-15-2	$C_9H_{11}CI_3NO_4P$	oxon	11	772
Chlorsulfuron	64902-72-3	$C_{12}H_{12}CIN_5O_4S$	triazinylsulfonyl urea	16	718
Chlorthiophos	60238-56-4	$\mathrm{C_{11}H_{15}Cl_{2}O_{3}PS_{2}}$	organophosphate	3	545
Chlozolinate	84332-86-5	$\mathrm{C_{13}H_{11}Cl_2NO_5}$	dichlorophenyl dicarboxamide; oxazole	1	AJS
Clethodim	99129-21-2	$C_{17}H_{26}CINO_3S$	cyclohexene oxime	28	AER
Clethodim 5-hydroxy sulfone	111031-11-9	$C_{17}H_{26}CINO_6S$	cyclohexene oxime metabolite	28	AJM
Clethodim sulfone	111031-17-5	$C_{17}H_{26}CINO_5S$	cyclohexene oxime metabolite	28	AJN
Clethodim sulfoxide	111031-14-2	$C_{17}H_{26}CINO_4S$	cyclohexene oxime metabolite	28	AJO
Clodinafop propargyl	105512-06-9	$C_{17}H_{13}CIFNO_4$	aryloxyphenoxypropionic acid	20	B38
Clofencet	129025-54-3	$C_{13}H_{11}CIN_2O_3$	unclassified	99	AET
Clofentezine	74115-24-5	$C_{14}H_8CI_2N_4$	tetrazine	99	699
Clomazone	81777-89-1	$C_{12}H_{14}CINO_2$	pyridazone	17	719
Clopyralid	1702-17-6	$C_6H_3Cl_2NO_2$	pyridinecarboxylic acid	20	B46
Cloquintocet methyl	99607-70-2	$C_{18}H_{22}CINO_3$	herbicide safener	1	AKE
Cloquintocet mexyl	99607-70-2	$C_{18}H_{22}CINO_3$	herbicide safener	1	B39
Cloransulam methyl	147150-35-4	$C_{15}H_{13}CIFN_5O_5S$	triazolopyrimidine herbicides	1	ALP
Clothianidin	210880-92-5	$C_6H_8CIN_5O_2S$	neonicotinyl	1	AEP
Coumaphos	56-72-4	$C_{14}H_{16}CIO_5PS$	phosphorothioate	11	124

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code
Coumaphos oxygen analog	321-54-0	$C_{14}H_{16}CIO_6P$	oxon	11	614
CPOPA (5-(4-Chlorophenyl)oxazole-2- propionic Acid)	23464-95-1	$C_{12}H_{10}CINO_3$	propionic acid	20	ALQ
Crotoxyphos	7700-17-6	$C_{14}H_{19}O_6P$	organophosphate	11	267
Crufomate	299-86-5	$C_{12}H_{19}CINO_3P$	phosphoramidate	11	667
Cumyluron	99485-76-4	$C_{17}H_{19}CIN_2O$	Urea	16	ANJ
Cyanazine	21725-46-2	$C_9H_{13}CIN_6$	triazine	9	228
Cyantraniliprole	736994-63-1	$C_{19}H_{14}BrCIN_6O_2$	pyrazole	1	AMB
Cyazofamid	120116-88-3	$C_{13}H_{13}CIN_4O_2S$	imidazole	1	AGA
Cyclanilide	113136-77-9	$C_{11}H_9Cl_2NO_3$	unclassified plant growth regulator	1	A81
Cycloate	1134-23-2	$C_{11}H_{21}NOS$	thiocarbamate	14	232
Cyflufenamid	180409-60-3	$C_{20}H_{17}F_5N_2O_2$	amide	1	AKU
Cyflumetofen	400882-07-7	$C_{24}H_{24}F_3NO_4$	bridged dipehnyl	3	AMC
Cyfluthrin	68359-37-5	$C_{22}H_{18}CI_2FNO_3$	pyrethroid	8	781
Cyhalofop butyl	122008-85-9	$C_{20}H_{20}FNO_4$	aryloxyphenoxypropionic herbicide	17	B59
Cyhalothrin (lambda)	91465-08-6	$\mathrm{C_{23}H_{19}CIF_{3}NO_{3}}$	pyrethroid	8	AEM
Cyhalothrin (lambda epimer R157836)	68085-85-8	$\mathrm{C_{23}H_{19}CIF_{3}NO_{3}}$	pyrethroid	8	AEN
Cyhalothrin total (L-cyhalothrin + R157836 epin	68085-85-8	$C_{23}H_{19}CIF_3NO_3$	pyrethroid	8	AEL
Cymoxanil	57966-95-7	$C_7H_{10}N_4O_3$	cyanoacetamide	2	877
Cypermethrin	52315-07-8	$C_{22}H_{19}CI_2NO_3$	pyrethroid	8	597
Cyphenothrin	39515-40-7	$C_{24}H_{25}NO_3$	pyrethroid	8	ADH
Cyproconazole	94361-06-5	$C_{15}H_{18}CIN_3O$	conazole	1	A22
Cyprodinil	121552-61-2	$C_{14}H_{15}N_3$	anilinopyrimidine	17	B22
Cyprosulfamide	221667-31-8	$C_{18}H_{18}N_2O_5S$	herbicide safeners	1	AMD
Cyromazine	66215-27-8	$C_6H_{10}N_6$	triazine	9	255
DCPA	1861-32-1	$C_{10}H_6Cl_4O_4$	phthalic acid	3	134
DCPA mono acid	887-54-7	$C_9H_4Cl_4O_4$	dicarboxylic acid	20	ABV

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code
DDD o,p'	53-19-0	$C_{14}H_{10}CI_4$	bridged biphenyl	3	909
DDD p,p'	72-54-8	$C_{14}H_{10}CI_4$	bridged biphenyl	3	908
DDE o,p'	3424-82-6	$C_{14}H_8CI_4$	bridged biphenyl	3	911
DDE p,p'	72-55-9	$C_{14}H_8Cl_4$	bridged biphenyl	3	910
DDT o,p'	789-02-6	$C_{14}H_9Cl_5$	bridged biphenyl	3	907
DDT p,p'	50-29-3	$C_{14}H_9Cl_5$	bridged biphenyl	3	906
DEET (N,N-diethyl-m-toluamide)	134-62-3	$C_{12}H_{17}NO$	amide	2	PBS
DEF (Tribufos)	78-48-8	$C_{12}H_{27}OPS_3$	organophosphate	11	217
Deltamethrin (includes parent Tralomethrin)	52918-63-5	$C_{22}H_{19}Br_2NO_3$	pyrethroid	8	612
Demeton	8065-48-3	$C_8H_{19}O_3PS_2$	phosphorothioate	11	023
Demeton-S	126-75-0	$C_8H_{19}O_3PS_2$	organothiophosphate	11	558
Demeton-S sulfone	2496-91-5	$C_8H_{19}O_5PS_2$	organothiophosphate metabolite	11	226
Desethyl atrazine	6190-65-4	$C_6H_{10}CIN_5$	triazine metabolite	9	964
Desethyl-desisopropyl atrazine	3397-62-4	$C_3H_4CIN_5$	triazine metabolite	9	784
Desisopropyl atrazine	1007-28-9	$C_5H_8CIN_5$	triazine metabolite	9	785
Desmedipham	13684-56-5	$\mathrm{C_{16}H_{16}N_2O_4}$	carbamate	14	786
Desmetryn	1014-69-3	$C_8H_{15}N_5S$	methylthiotriazine	9	A88
Dialifos	10311-84-9	$C_{14}H_{17}CINO_4PS_2$	organothiophosphate	11	244
Diazinon	333-41-5	$C_{12}H_{21}N_2O_3PS$	phosphorothioate	11	024
Diazinon oxygen analog	962-58-3	$C_{12}H_{21}N_2O_4P$	oxon	11	395
Dicamba	1918-00-9	$C_8H_6Cl_2O_3$	benzoic acid	20	155
Dichlobenil	1194-65-6	$C_7H_3Cl_2N$	nitrile	2	324
Dichlofenthion	97-17-6	$C_{10}H_{13}CI_2O_3PS$	phenylorganothiophosphate	11	664
Dichlofluanid	1085-98-9	$C_9H_{11}CI_2FN_2O_2S_2$	phenylsulfamide	1	588
Dichlormid	37764-25-3	$C_8H_{11}Cl_2NO$	herbicide safeners	1	A43
Dichlorprop	120-36-5	$C_9H_8Cl_2O_3$	phenoxy acid	20	A25
Dichlorvos (DDVP)	62-73-7	$C_4H_7CI_2O_4P$	phosphoric acid	11	338

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code
Diclofop methyl	51338-27-3	$\mathrm{C_{16}H_{14}Cl_{2}O_{4}}$	aryloxyphenoxypropionic acid	20	299
Dicloran	99-30-9	$C_6H_4Cl_2N_2O_2$	nitroaniline	7	144
Diclosulam	145701-21-9	$\mathrm{C_{13}H_{10}Cl_2FN_5O_3S}$	triazolopyrimidine herbicides	1	ALU
Dicofol o,p'	10606-46-9	$C_{14}H_9CI_5O$	bridged biphenyl	3	253
Dicofol p,p'	115-32-2	$C_{14}H_9CI_5O$	bridged biphenyl	3	254
Dicrotophos	141-66-2	$C_8H_{16}NO_5P$	organophosphate	11	209
Dieldrin	60-57-1	$C_{12}H_8Cl_6O$	cyclodiene	3	028
Diethofencarb	87130-20-9	$C_{14}H_{21}NO_4$	carbanilite fungicides	22	B62
Difenoconazole	119446-68-3	$\mathrm{C_{19}H_{17}Cl_2N_3O_3}$	triazole	1	B58
Diflubenzuron	35367-38-5	$C_{14}H_9CIF_2N_2O_2$	urea	16	651
Diflufenzopyr	109293-97-2	$C_{15}H_{12}F_2N_4O_3$	urea	16	AFY
Dimepiperate	61432-55-1	$C_{15}H_{21}NOS$	unclassified	99	E13
Dimethenamid	87674-68-8	$C_{12}H_{18}CINO_2S$	acetamide	1	ADD
Dimethenamid ethanesulfonic acid	205939-58-8	$\mathrm{C_{12}H_{19}NO_5S_2}$	acetamide metabolite	20	AEX
Dimethenamid oxanilic acid	NA ^[2]	$C_{12}H_{17}NO_4S$	acetamide metabolite	20	AEY
Dimethenamid P	87674-68-8	$\mathrm{C_{12}H_{18}CINO_{2}S}$	amide	1	AEB
Dimethipin	55290-64-7	$C_6H_{10}O_4S_2$	urea	16	787
Dimethoate	60-51-5	$C_5H_{12}NO_3PS_2$	phosphorodithionic acid	11	171
Dimethomorph	110488-70-5	$C_{21}H_{22}CINO_4$	chlorophenyl morpholine	3	B77
Diniconazole	83657-24-3	$\mathrm{C_{15}H_{17}Cl_2N_3O}$	conazole	1	AFN
Dinocap	131-72-6	$C_{18}H_{24}N_2O_6$	dinitrophenol	20	315
Dinoseb	88-85-7	$C_{10}H_{12}N_2O_5$	phenol	20	031
Dinotefuran	165252-70-0	$C_7H_{14}N_4O_3$	neonicotinyl	1	AFO
Dioxacarb	6988-21-2	$C_{11}H_{13}NO_4$	phenylmethyl carbamate	14	656
Dioxathion	78-34-2	$C_{12}H_{26}O_6P_2S_4$	organothiophosphate	11	103
Diphacinone	82-66-6	$C_{23}H_{16}O_{3}$	indandione	99	B99
Diphenamid	957-51-7	$C_{16}H_{17}NO$	acetamide	1	330

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code
Diphenylamine (DPA)	122-39-4	$C_{12}H_{11}N$	amine	3	125
Dipropetryn	4147-51-7	$C_{11}H_{21}N_5S$	triazine	9	735
Disulfoton	298-04-4	$C_8H_{19}O_2PS_3$	phosphorodithioate	11	117
Disulfoton oxygen analog	NA ^[2]	$C_8H_{19}O_3PS_2$	organophosphate	11	AHN
Disulfoton oxygen analog sulfone	NA ^[2]	$C_6H_{15}O_5PS_2$	organophosphate	11	AHV
Disulfoton oxygen analog sulfoxide	NA ^[2]	$C_6H_{15}O_4PS_2$	organophosphate	11	AHW
Disulfoton sulfone	2497-06-5	$C_8H_{19}O_4PS$	sulfone	11	216
Disulfoton sulfoxide	2497-07-6	$C_8H_{19}O_3PS_3$	sulfoxide	11	706
Ditalimfos	5131-24-8	$C_{12}H_{14}NO_4PS$	organophosphorus	11	E22
Dithianon	3347-22-6	$C_{14}H_4N_2O_2S_2$	quinine	17	АНО
Dithiopyr	97886-45-8	$\mathrm{C_{15}H_{16}F_5NO_2S_2}$	pyridine	20	E24
Diuron	330-54-1	$C_9H_{10}Cl_2N_2O$	urea	16	032
Dodine	2439-10-3	$C_{15}H_{33}N_{3}O_{2}$	aliphatic nitrogenous fungicide	2	104
Emamectin benzoate	155569-91-8	$C_{49}H_{75}NO_{13} + C_{48}H_{73}NO_{13}$	avermectin (macrocyclic lactone)	29	AGH
Endosulfan I	959-98-8	$C_9H_6CI_6O_3S$	cyclodiene	3	900
Endosulfan II	33213-65-9	$C_9H_6CI_6O_3S$	cyclodiene	3	901
Endosulfan sulfate	1031-07-8	$C_9H_6CI_6O_4S$	cyclodiene	3	902
Endrin	72-20-8	$C_{12}H_8Cl_6O$	cyclodiene	3	034
Endothall	145-73-3	$C_8H_{10}O_5$	dicarboxylic acid	21	AKV
Epoxiconazole	135319-73-2	$C_{17}H_{13}CIFN_3O$	conazole	1	B53
Eprinomectin	123997-26-2	$C_{50}H_{75}NO_{14}$ (eprinomectin B_{1a}) + $C_{49}H_{73}NO_{14}$ (eprinomectin B_{1b})	macrocyclic lactone	29	AHR
EPN	2104-64-5	$C_{14}H_{14}NO_4PS$	phenyl phenylphosphonothioate insecticides	11	035
EPTC	759-94-4	$C_9H_{19}NOS$	thiocarbamate	14	200
Esfenvalerate	66230-04-4	$C_{25}H_{22}CINO_3$	pyrethroid	8	714

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Ethalfluralin	55283-68-6	$\mathrm{C}_{13}H_{14}F_3N_3O_4$	dinitroaniline	7	721
Ethephon	16672-87-0	$C_2H_6CIO_3P$	plant growth regulator	11	730
Ethiofencarb	29973-13-5	$C_{11}H_{15}NO_2S$	carbamate	14	858
Ethiofencarb sulfone	53380-23-7	$C_{11}H_{15}NO_4S$	carbamate	14	AMX
Ethiofencarb sulfoxide	53380-22-6	$C_{11}H_{15}NO_3S$	carbamate	14	AMY
Ethion	563-12-2	$C_9H_{22}O_4P_2S_4$	phosphorodithioic acid	11	107
Ethion di oxon	22756-17-8	$\mathrm{C_9H_{22}O_6P_2S_2}$	oxon	11	538
Ethion mono oxon	17356-42-2	$C_9H_{22}O_5P_2S_3$	oxon	11	AAX
Ethiprole	181587-01-9	$\mathrm{C_{13}H_9Cl_2F_3N_4OS}$	phenylpyrazole	1	AME
Ethofumesate	26225-79-6	$C_{13}H_{18}O_5S$	benzofuranyl alkylsulfonate	11	945
Ethoprop	13194-48-4	$C_8H_{19}O_2PS_2$	dipropyl phosphorodithioate	11	175
Ethoxyquin	91-53-2	$C_{14}H_{19}NO$	quinoline	99	111
Ethylan	72-56-0	$\mathrm{C_{18}H_{20}Cl_2}$	organochlorine	3	066
Etofenprox	80844-07-1	$C_{25}H_{28}O_3$	pyrethroid ether	8	ADI
Etoxazole	1532333-91-1	$\mathrm{C}_{21}H_{23}F_2NO_2$	oxazole	1	B84
Etridiazole	2593-15-9	$C_5H_5CI_3N_2OS$	thiadiazole	1	722
Etrimfos	38260-54-7	$\mathrm{C_{10}H_{17}N_2O_4PS}$	pyrimidine organothiophosphate	11	293
Famoxadone	131807-57-3	$\mathrm{C}_{22}H_{18}N_{2}O_{4}$	dicarboximide/oxazole	1	AEW
Famphur	52-85-7	$\mathrm{C_{10}H_{16}NO_5PS_2}$	phenylorganothiophosphate	11	603
Fenamidone	161326-34-7	$C_{17}H_{17}N_3OS$	imidazole	1	B64
Fenamiphos	22224-92-6	$C_{13}H_{22}NO_3PS$	phosphoramidate	11	236
Fenamiphos sulfone	31972-44-8	$C_{13}H_{22}NO_5PS$	sulfone	11	745
Fenamiphos sulfoxide	31972-43-7	$C_{13}H_{22}NO_4PS$	sulfoxide	11	746
Fenarimol	60168-88-9	$\mathrm{C_{17}H_{12}Cl_2N_2O}$	pyrimidine	3	271
Fenazaquin	120928-09-8	$C_{20}H_{22}N_2O$	unclassified acaricide	27	B73
Fenbuconazole	114369-43-6	$C_{19}H_{17}CIN_4$	conazole	1	A30
Fenbutatin oxide	13356-08-6	$C_{60}H_{28}OSn_2$	organotin acaride	99	639

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code
Fenchlorphos (ronnel)	299-84-3	$C_8H_8CI_3O_3PS$	phenyl organothiophosphate	11	105
Fenhexamid	126833-17-8	$C_{14}H_{17}CI_2NO_2$	chlorocarboximide	1	B41
Fenitrothion	122-14-5	$C_9H_{12}NO_5PS$	phosphorothioate	11	391
Fenitrothion oxygen analog	2255-17-6	$C_9H_{12}NO_6P$	oxon	11	648
Fenobucarb (BPMC)	3766-81-2	$C_{12}H_{17}NO_2$	phenyl methylcarbamate	14	856
Fenoxaprop ethyl	66441-23-4	$C_{18}H_{16}CINO_5$	aryloxyphenoxypropionic acid	20	777
Fenoxycarb	72490-01-8	$\mathrm{C_{17}H_{19}NO_4}$	carbamate	14	811
Fenpropathrin	39515-41-8	$C_{22}H_{23}NO_3$	pyrethroid	8	808
Fenpropidin	67306-00-7	$C_{19}H_{31}N$	nitrogenous hetercyclic	17	AMF
Fenpropimorph	67564-91-4	$C_{20}H_{33}NO$	morpholine	3	886
Fenpyrazamine	473798-59-3	$C_{17}H_{21}N_{3}O_{2}S$	pyrazole	1	AMG
Fenpyroximate	111812-58-9	$\mathrm{C}_{24}\mathrm{H}_{27}\mathrm{N}_{3}\mathrm{O}_{4}$	phenoxypyrazol	1	AFS
Fensulfothion	115-90-2	$C_{11}H_{17}O_4PS_2$	phenyl organothiophosphate	11	243
Fenthion	55-38-9	$\mathrm{C_{10}H_{15}O_{3}PS_{2}}$	phosphorothioate	11	177
Fenthion oxygen analog	6552-12-1	$C_{10}H_{15}O_4PS$	oxon	11	691
Fenthion sulfone	3761-42-0	$\mathrm{C_{10}H_{15}O_5PS_2}$	organophosphate	11	660
Fenthion sulfoxide	3761-41-9	$\mathrm{C_{10}H_{15}O_4PS_2}$	organophosphate	11	AKP
Fenuron	101-42-8	$C_9H_{12}N_2O$	urea	16	840
Fenvalerate	51630-58-1	$C_{25}H_{22}CINO_3$	pyrethroid	8	546
Fipronil	120068-37-3	$\mathrm{C_{12}H_4C_{l2}F_6N_4OS}$	phenyl pyrazole	1	A82
Fipronil sulfone	120068-36-2	$\mathrm{C}_{12}H_4CI_2F_6N_4O_2S$	phenylpyrazole	1	A84
Flazasulfuron	104040-78-0	$\mathrm{C}_{13}H_{12}F_{3}N_{5}O_{5}S$	pyridiminyl sulfonylurea	16	AMH
Flonicamid	158062-67-0	$C_9H_6F_3N_3O$	nicotinoid	1	AGG
Fluazifop	69335-91-7	$C_{15}H_{12}F_3NO_4$	pyridine	17	ALW
Fluazifop butyl	69806-50-4	$\mathrm{C_{15}H_{12}F_{3}NO_{4}}$	pyridine	17	292
Fluazinam	79622-59-6	$\mathrm{C_{13}H_4C_{l2}F_6N_4O_4}$	pyridine	17	B54
Flubendiamide	272451-65-7	$C_{23}H_{22}F_7IN_2O_4S$	diamide	17 or 1	AHS

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Flucythrinate	70124-77-5	$\mathrm{C_{26}H_{23}F_2NO_4}$	pyrethroid	8	229
Fludioxonil	131341-86-1	$\mathrm{C}_{12}H_{6}F_{2}N_{2}O_{2}$	phenyl pyrrole	1	B23
Fluensulfone	318290-98-1	$C_7H_5ClF_3NO_2S_2$	unclassified nematicide	99	ANK
Flufenacet	142459-58-3	$\mathrm{C}_{14}H_{13}F_4N_3O_2S$	anilide	1	B30
Flufenacet ethanesulfonic acid	NA ^[2]	$C_{11}H_{14}FNO_4S$	anilide metabolite	20	AFH
Flufenacet oxanilic acid	201668-31-7	$C_{11}H_{12}FNO_3$	anilide metabolite	20	AEZ
Flufenoxuron	101463-69-8	$\mathrm{C}_{21}H_{11}CIF_6N_2O_3$	urea	16	AHG
Flufenpyr ethyl	188489-07-8	$\mathrm{C_{16}H_{13}CIF_4N_2O_4}$	pyridazinone	17	ALR
Flumetralin	62924-70-3	$\mathrm{C_{16}H_{12}CIF_4N_3O_4}$	growth inhibitor	7	834
Flumetsulam	98967-40-9	$C_{12}H_9F_2N_5O_2S$	pyrimidine	1	AAU
Flumiclorac pentyl	87546-18-7	$C_{21}H_{23}CIFNO_5$	dicarboximide	1	AAV
Flumioxazin	103361-09-7	$\mathrm{C_{19}H_{15}FN_2O_2}$	N-phenylphthalimide	1	AFF
Fluometuron	2164-17-2	$C_{10}H_{11}F_{3}N_{2}O$	urea	16	701
Fluopicolide	239110-15-7	$\mathrm{C_{14}H_8Cl_3F_3N_2O}$	pyridine	17 or 1	AHT
Fluopyram	658066-35-4	$\mathrm{C_{16}H_{11}CIF_6N_2O}$	benzamide	1	AKG
Fluorodifen	15457-05-3	$\mathrm{C}_{13}H_{7}F_{3}N_{2}O_{5}$	nitrophenyl ether	3	836
Fluoxastrobin	361377-29-9	$C_{21}H_{16}CIFN_4O_5$	strobilurin	22	AGJ
Flupyradifurone	951659-40-8	$\mathrm{C}_{12}H_{11}CIF_2N_2O_2$	unclassified	1	ANE
Fluquinconazole	136426-54-5	$C_{16}H_8Cl_2FN_5O$	aryloxyphenoxypropionic acid	1	B78
Fluridone	59756-60-4	$C_{19}H_{14}F_3NO$	pyridine	17	736
Fluroxapyr-1-methylheptyl ester	81406-37-3	$\mathrm{C_{15}H_{22}Cl_2FN_2O_3}$	pyridine	17	ADJ
Flusilazole	85509-19-9	$\mathrm{C_{16}H_{15}F_2N_3Si}$	conazole	1	950
Fluthiacet methyl	117337-19-6	$\mathrm{C_{15}H_{15}CIFN_{3}O_{3}S_{2}}$	phenylpyrazole	1	AGM
Flutolanil	66332-96-5	$\mathrm{C}_{17}H_{16}F_3NO_2$	caboxamide	1	B63
Flutriafol	76674-21-0	$C_{16}H_{13}F_2N_3O$	conazole	1	AFM
Fluvalinate	69409-94-5	$\mathrm{C_{26}H_{22}CIF_3N_2O_3}$	pyrethroid	8	297
Fluxapyroxad	907204-31-3	$\mathrm{C}_{18}H_{12}F_{5}N_{3}O$	anilide; pyrazole	22	AKW

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Folpet	133-07-3	$C_9H_4CI_3NO_2S$	phthalimide	1	126
Fomesafen	72178-02-0	$\mathrm{C_{15}H_{10}CIF_{3}N_{2}O_{6}S}$	amide or nitrophenyl ether herbicide	1 or 3	ALX
Fonofos	944-22-9	$C_{10}H_{15}OPS_2$	phosphorodithioic acid	11	163
Fonofos oxygen analog	944-21-8	$\mathrm{C_{10}H_{15}O_{2}PS}$	oxon	11	692
Forchlorfenuron	68157-60-8	$C_{12}H_{10}CIN_3O$	phenyl urea	16	B32
Formetanate hydrochloride	23422-53-9	$\mathrm{C_{11}H_{15}N_3O_2}$	formamidine	1	723
Fosthiazate	98886-44-3	$C_9H_{18}NO_3PS_2$	organothiophosphate	11	B09
Furalaxyl	57646-30-7	$C_{17}H_{19}NO_4$	furanilide	1	AMZ
Furathiocarb	65907-30-4	$C_{18}H_{26}N_2O_5S$	benzofurayl methyl carbamate	14	AMR
Glufosinate	77182-82-2	$C_5H_{12}NO_4P$	quaternary ammonium	99	AJL
Glyphosate	1071-83-6	$C_3H_8NO_5P$	organophosphate	99	653
Halosulfuron	135397-30-7	$C_{12}H_{13}CIN_6O_7S$	sulfonyl urea	16	AFK
Halosulfuron methyl	100784-20-1	$C_{12}H_{13}CIN_6O_7S$	sulfonyl urea	16	AEH
Haloxyfop	69806-34-4	$C_{15}H_{11}CIF_3NO_4$	aryloxyphenoxypropionic acid	20	798
Heptachlor	76-44-8	$C_{10}H_5CI_7$	cyclodiene	3	044
Heptachlor epoxide	1024-57-3	$C_{10}H_5CI_7O$	cyclodiene	3	143
Heptenophos	23560-59-0	$C_9H_{12}CIO_4P$	organophosphate	11	841
Hexachlorobenzene (HCB)	118-74-1	C_6Cl_6	benzene ring	3	321
Hexaconazole	79983-71-4	$\mathrm{C_{14}H_{17}Cl_2N_3O}$	conazole	1	954
Hexaflumuron	86479-06-3	$\mathrm{C_{16}H_8Cl_2F_6N_2O_3}$	benzoylphenylurea chitin synthesis inhib	16	AMA
Hexazinone	51235-04-2	$C_{12}H_{20}N_4O_2$	triazine	9	633
Hexythiazox	78587-05-0	$C_{17}H_{21}CIN_2O_2S$	thiazolidine carboxamide	1	B10
Hydroprene	41096-46-2	$C_{17}H_{30}O_2$	oxyhydrocarbon	21	AEC
Hydroxy atrazine	2163-68-0	$C_8H_{15}N_5O$	triazine metabolite	9	AED
Imazalil	35554-44-0	$C_{14}H_{14}CI_2N_2O$	conazole	1	604
Imazamethabenz acid	NA ^[2]	$C_{15}H_{18}N_2O_3$	imidazolinone	16	AEE
Imazamethabenz methyl	81405-85-8	$C_{16}H_{20}N_2O_3$	imidazolinone	16	753

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Imazamox	114311-32-9	$\mathrm{C_{15}H_{19}N_{3}O_{4}}$	imidazolinone	16	ACA
Imazapic	104098-48-8	$\mathrm{C}_{14}H_{17}N_3O_3$	imidazolinone	16	ACZ
Imazapyr	81334-34-1	$\mathrm{C_{13}H_{15}N_{3}O_{3}}$	imidazolinone	16	ACB
Imazaquin	81335-37-7	$\mathrm{C_{17}H_{17}N_{3}O_{3}}$	imidazolinone	16	ACC
Imazethapyr	81335-77-5	$\mathrm{C_{15}H_{19}N_{3}O_{3}}$	imidazolinone	16	ACD
Imazosulfuron	122548-33-8	$\mathrm{C_{14}H_{13}CIN_6O_5S}$	sulfonyl urea	16	AMK
Imidacloprid	138261-41-3	$C_9H_{10}CIN_5O_2$	neonicotinyl	1	967
Imidacloprid urea	120868-66-8	$C_9H_{10}CIN_3O$	neonicotinyl metabolite	1	AHF
Imiprothrin	72963-72-5	$\mathrm{C_{17}H_{22}N_2O_4}$	pyrethroid	8	ADK
Indaziflam	950782-86-2	$\mathrm{C_{16}H_{20}FN_5}$	triazine	9	AJP
Indoxacarb	173584-44-6	$\mathrm{C}_{22}H_{17}CIF_3N_3O_7$	carbamate	14	ADG
Iodosulfuron methyl	144550-36-7	$\mathrm{C_{14}H_{14}IN_5O_6S}$	triazinylsulfonyl urea	16	AKB
Ipconazole	125225-28-7	$C_{18}H_{24}CIN_3O$	conazole	1	AHY
Iprobenfos	26087-47-8	$C_{13}H_{21}O_3PS$	organophosphate	11	867
Iprodione	36734-19-7	$\mathrm{C_{13}H_{13}Cl_2N_3O_3}$	dicarboximide	1	626
Iprodione metabolite isomer	63637-89-8	$\mathrm{C_{13}H_{13}Cl_2N_3O_3}$	dicarboximide	1	231
Iprovalicarb	140923-17-7	$C_{18}H_{28}N_2O_3$	carbamates	14	AGE
Isocarbophos	24353-61-5	$\mathrm{C_{11}H_{16}NO_4PS}$	phosphoramidothioate	11	ALE
Isofenphos	25311-71-1	$C_{15}H_{24}NO_4PS$	organophosphate	11	258
Isofenphos methyl	99675-03-3	$C_{14}H_{22}NO_4PS$	phosphoramidothioate	11	ANA
Isofenphos oxygen analog	31120-85-1	$C_{15}H_{24}NO_5P$	oxon	11	655
Isofetamid	875915-78-9	$C_{20}H_{25}NO_3S$	amide/thiophene	1	ANH
Isoprocarb	2631-40-5	$C_{11}H_{15}NO_2$	carbamate	14	637
Isoprothiolane	50512-35-1	$C_{12}H_{18}O_4S_2$	unclassified	99	855
Isoproturon	34123-59-6	$C_{12}H_{18}N_2O$	phenyl urea	16	843
Isoxadifen ethyl	163520-33-0	$C_{18}H_{17}NO_3$	herbicide safeners	1	AGL
Isoxaflutole	141112-29-0	$\mathrm{C_{15}H_{12}F_{3}NO_{4}S}$	cyclopropylisoxazole	17	B15

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Kasugamycin	6980-18-3	$C_{14}H_{25}N_3O_9$	bactericide	99	AKY
Kresoxim methyl	143390-89-0	$C_{18}H_{19}NO_4$	strobilurin	22	B42
Lactofen	77501-63-4	$\mathrm{C_{19}H_{15}CIF_{3}NO_{7}}$	flurodiphenyl ether	3	A38
Leptophos oxygen analog	25006-32-0	$\mathrm{C_{13}H_{10}BrCl_2O_3P}$	oxon	11	A40
Lenacil	2164-08-1	$C_{13}H_{18}N_2O_2$	uracil	16	859
Lindane (BHC gamma)	58-89-9	$C_6H_6CI_6$	hexane ring	3	050
Linuron	330-55-2	$C_9H_{10}CI_2N_2O_2$	urea	16	129
Lufenuron	103055-07-8	$\mathrm{C_{17}H_8Cl_2F_8N_2O_3}$	benzoylphenylurea	16	AJV
Malathion	121-75-5	$C_{10}H_{19}O_6PS_2$	phosphorodithioate	11	052
Malathion oxygen analog	1634-78-2	$C_{10}H_{19}O_7PS$	oxon	11	208
Mandipropamid	374726-62-2	$C_{23}H_{22}CINO_4$	amide	1	AGX
MCPA	94-74-6	$C_9H_9CIO_3$	phenoxy	20	318
МСРВ	94-81-5	$C_{11}H_{13}CIO_3$	phenoxy acid	20	620
Mecarbam	2595-54-2	$\mathrm{C_{10}H_{20}NO_5PS_2}$	organophosphate	11	662
Mecoprop (MCPP)	7085-19-0	$C_{10}H_{11}CIO_3$	phenoxy acid	20	A42
Mefenacet	73250-68-7	$C_{16}H_{14}N_2O_2S$	anilide	1	D21
Mefenpyr diethyl	135590-91-9	$C_{16}H_{18}CI_2N_2O_4$	herbicide safeners	1	AKH
Melamine	108-78-1	$C_3H_6N_6$	trimer of cyanamide	9	260
Mepanipyrim	1102335-47-7	$\mathrm{C}_{14}H_{13}N_{3}$	pyrimidine	17	AGF
Mephosfolan	950-10-7	$C_8H_{16}NO_3PS_2$	phosphoramidate	11	242
Merphos	150-50-5	$C_{12}H_{27}PS_3$	plant growth regulator	11	121
Mesosulfuron methyl	208465-21-8	$C_{17}H_{21}N_5O_9S_2$	pyrimidinylsulfonyl urea	16	AKJ
Mesotrione	104206-82-8	$C_{14}H_{13}NO_7S$	benzoylcyclohexanedione	1	AJA
Metaflumizone	139968-49-3	$\mathrm{C}_{24}H_{16}F_{6}N_{4}O_{2}$	unclassified	1	AJW
Metalaxyl	57837-19-1	$C_{15}H_{21}NO_4$	acylalanine	1	607
Metaldehyde	108-62-3	$C_8H_{16}O_4$	polyaldehyde	99	B07
Metconazole	125116-23-6	$C_{17}H_{22}CIN_3O$	conazole	1	AHX

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Metolcarb	1129-41-5	$C_9H_{11}NO_2$	carbamate	14	860
Metrafenone	220899-03-6	$C_{19}H_{21}BrO_5$	arylphenylketone	3	ANF
Methamidophos	10265-92-6	$C_2H_8NO_2PS$	phosphoramidothioic acid	11	170
Methacrifos	62610-77-9	$C_7H_{13}O_5PS$	organothiophosphate	11	E74
Methidathion	950-37-8	$C_6H_{11}N_2O_4PS_3$	phosphorodithioate	11	197
Methidathion oxygen analog	39856-16-1	$C_6H_{11}N_2O_5PS_2$	oxon	11	ACE
Methiocarb	2032-65-7	$C_{11}H_{15}NO_2S$	carbamate	14	195
Methiocarb sulfone	2179-25-1	$C_{11}H_{15}NO_4S$	carbamate metabolilte	14	634
Methiocarb sulfoxide	2635-10-1	$C_{11}H_{15}NO_3S$	carbamate metabolilte	14	256
Methomyl	16752-77-5	$C_5H_{10}N_2O_2S$	carbamate	14	159
Methoprene	40596-69-8	$C_{19}H_{34}O_3$	oxyhydrocarbon	21	ACV
Methoxychlor olefin	2132-70-9	$C_{16}H_{14}CI_2O_2$	bridged biphenyl	3	276
Methoxychlor p,p'	72-43-5	$C_{16}H_{15}CI_3O_2$	bridged biphenyl	3	275
Methoxychlor Total	72-43-5	$C_{16}H_{15}CI_3O_2$	bridged biphenyl	3	055
Methoxyfenozide	161050-58-4	$C_{22}H_{28}N_2O_3$	diacylhydrazine	16	AES
Methyl pentachlorophenyl sulfide (MPCPS)	1825-19-0	C ₇ H ₃ Cl ₅ S	benzene ring	3	388
Metolachlor	51218-45-2	$C_{15}H_{22}CINO_2$	acetamide	1	283
Metolachlor ethanesulfonic acid	NA ^[2]	$C_{15}H_{23}NO_5S$	chloroacetanilide metabolite	20	ACG
Metolachlor oxanilic acid	152019-73-3	$C_{15}H_{21}NO_4$	chloroacetanilide metabolite	20	ACH
Metribuzin	21087-64-9	$C_8H_{14}N_4OS$	triazines	9	181
Metsulfuron methyl	74223-64-6	$\mathrm{C}_{14}\mathrm{H}_{15}\mathrm{N}_{5}\mathrm{O}_{6}\mathrm{S}$	sulfonyl urea	16	ACI
Mevinphos E/Z	298-01-1	$C_7H_{13}O_6P$	butenoic acid	11	579
MGK-264	113-48-4	$C_{17}H_{25}NO_2$	synergist	8	058
MGK-326 (Dipropyl isocinchomeronate)	136-45-8	$C_{13}H_{17}NO_4$	synergist	8	ADL
Milbemectin	51596-10-2	$C_{31}H_{44}O_7$ (milbemycin A ₃) + $C_{32}H_{46}O_7$ (milbemycin A ₄)	macrocyclic lactone	29	AHP

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Mirex	2385-85-5	C ₁₀ Cl ₁₂	cyclodiene	3	352
Molinate	2212-67-1	$C_9H_{17}NO_5$	thiocarbamate	14	778
Monocrotophos	6923-22-4	$C_7H_{14}NO_5P$	phosphoric acid	11	343
Monolinuron	1746-81-2	$\mathrm{C_9H_{11}CIN_2O_2}$	phenyl urea	16	682
Monuron	150-68-5	$C_9H_{11}CIN_2O$	urea	16	046
Myclobutanil	88671-89-0	$C_{15}H_{17}CIN_4$	triazole	1	679
Naled	300-76-5	$C_4H_7Br_2Cl_2O_4P$	organophosphate	11	303
Napropamide	15299-99-7	$C_{17}H_{21}NO_2$	amide	1	594
Naptalam (Alanap)	132-66-1	$C_{18}H_{13}NO_3$	amide	1	B18
Neburon	555-37-3	$C_{12}H_{16}C_{12}N_2O$	urea	16	061
Niclosamide	50-65-7	$C_{13}H_8Cl_2N_2O_4$	molluscicide	3	ACL
Nicosulfuron	111991-09-4	$\mathrm{C_{15}H_{18}N_6O_6S}$	sulfonyl urea	16	ACM
Nitrapyrin	1929-82-4	$C_6H_3Cl_4N$	pyridine	17	725
Nitrofen	1836-75-5	$C_{12}H_7Cl_2NO_3$	nitrophenyl ether herbicides	3	158
Norflurazon	27314-13-2	$C_{12}H_9CIF_3N_3O$	pyridazinone	17	596
Norflurazon desmethyl	23576-24-1	$C_{11}H_7CIF_3N_3O$	pyridazinone	17	720
Novaluron	116714-46-6	$\mathrm{C_{17}H_9ClF_8N_2O_4}$	benzoyl urea	16	AFX
Omethoate	1113-02-6	$C_5H_{12}NO_4PS$	phosphorothioate	11	178
o-Phenylphenol	90-43-7	$C_{12}H_{10}O$	biphenyl	3	083
Oryzalin	19044-88-3	$\mathrm{C_{12}H_{18}N_4O_6S}$	dinitroaniline	7	737
Oxadiazon	19666-30-9	$\mathrm{C_{15}H_{18}Cl_2N_2O_3}$	oxadiazon	1	625
Oxadixyl	77732-09-3	$C_{14}H_{18}N_2O_4$	oxazolidine	1	A46
Oxamyl	23135-22-0	$C_7H_{13}N_3O_3S$	carbamate	14	537
Oxamyl oxime	30558-43-1	$C_5H_{10}N_2O_2S$	carbamate	14	A47
Oxychlordane	27304-13-8	$C_{10}H_4CI_8O$	cyclodiene	3	349
Oxydemeton methyl	301-12-2	$\mathrm{C_6H_{15}O_4PS_2}$	organophosphate	11	219
Oxydemeton methyl sulfone	17040-19-6	$C_6H_{15}O_5PS_2$	phosphorothioate	11	245

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code
Oxyfluorfen	42874-03-3	$\mathrm{C_{15}H_{11}CIF_{3}NO_{4}}$	diphenyl ether	3	713
Oxythioquinox	2439-01-2	$C_{10}H_6N_2OS_2$	quionoxaline	17	246
Paclobutrazol	76738-62-0	$C_{15}H_{20}CIN_3O$	growth inhibitor	1	A48
Parathion ethyl	56-38-2	$C_{10}H_{14}NO_5PS$	phosphorothionic acid	11	065
Parathion ethyl oxygen analog	NA ^[2]	$C_{10}H_{14}NO_6P$	oxon	11	370
Parathion methyl	298-00-0	$C_8H_{10}NO_5PS$	phosphorothionic acid	11	057
Parathion methyl oxygen analog	950-35-6	$C_8H_{10}NO_6P$	oxon	11	779
Pebulate	1114-71-2	$C_{10}H_{21}NOS$	thiocarbamate	14	161
Penconazole	66246-88-6	$\mathrm{C_{13}H_{15}Cl_2N_3}$	conazole	1	956
Pencycuron	66063-05-6	$C_{19}H_{21}CIN_2O$	urea	16	AJX
Pendimethalin	40487-42-1	$\mathrm{C_{13}H_{19}N_{3}O_{4}}$	dinitroaniline	7	230
Penflufen	494793-67-8	$\mathrm{C_{18}H_{24}FN_{3}O}$	pyrazole	1	AKZ
Penoxsulam	219714-96-2	$\mathrm{C_{16}H_{14}F_5N_5O_5S}$	triazolpyrimidine	1	AMS
Pentachloroaniline (PCA)	527-20-8	$C_6H_2CI_5N$	aniline	3	351
Pentachlorobenzene (PCB)	608-93-5	C_6HCl_5	benzene ring	3	387
Penthiopyrad	183675-82-3	$\mathrm{C_{16}H_{20}F_{3}N_{3}OS}$	pyridazinone	22	AKD
Permethrin cis	61949-76-6	$C_{21}H_{20}CI_2O_3$	pyrethroid	8	222
Permethrin total	52645-53-1	$C_{21}H_{20}CI_2O_3$	pyrethroid	8	539
Permethrin trans	61949-77-7	$C_{21}H_{20}CI_2O_3$	pyrethroid	8	223
Perthane	72-56-0	$C_{18}H_{20}CI_2$	organochlorine	3	AGN
Phenmedipham	13684-63-4	$\mathrm{C_{16}H_{16}N_2O_4}$	carbamate	14	791
Phenothrin	26002-80-2	$C_{23}H_{26}O_{3}$	pyrethroid	8	848
Phenthoate	2597-03-7	$\mathrm{C_{12}H_{17}O_4PS_2}$	organophosphate	11	377
Phorate	298-02-2	$C_7H_{17}O_2PS_3$	phosphorodithionic acid	11	148
Phorate oxygen analog	2600-69-3	$C_7H_{17}O_3PS_2$	oxon	11	928
Phorate oxygen analog sulfone	2588-06-9	$C_7H_{17}O_5PS_2$	organophosphate	11	966
Phorate oxygen analog sulfoxide	2588-05-8	$C_7H_{17}O_4PS_2$	organophosphate	11	951

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code
Phorate sulfone	2588-04-7	$C_7H_{17}O_4PS_3$	sulfone	11	189
Phorate sulfoxide	2588-03-6	$C_7H_{17}O_3PS_2$	sulfoxide	11	190
Phosalone	2310-17-0	$C_{12}H_{15}CINO_4PS_2$	phosphorodithionic acid	11	166
Phosalone oxygen analog	2275-06-1	$C_{12}H_{15}CINO_5PS$	oxon	11	929
Phosmet	732-11-6	$\mathrm{C_{11}H_{12}NO_4PS_2}$	phosphorodithionic acid	11	165
Phosmet oxygen analog	3735-33-9	$C_{11}H_{12}NO_5PS$	oxon	11	237
Phosphamidon	13171-21-6	$C_{10}H_{19}CINO_5P$	dimethyl phosphate	11	203
Phoxim	14816-18-3	$\mathrm{C}_{12}H_{15}N_2O_3PS$	organothiophosphate	11	247
Picloram	1918-02-1	$\mathrm{C_6H_3Cl_3N_2O_2}$	carboxylic acid	20	329
Picoxystrobin	117428-22-5	$\mathrm{C}_{18}H_{16}F_3NO_4$	strobilurin	22	ALA
Pinoxaden	243973-20-8	$\mathrm{C}_{23}H_{32}N_2O_4$	phenylpyrazole	22	AHH
Piperalin	3478-94-2	$\mathrm{C_{16}H_{21}Cl_2NO_2}$	unclassified	99	AGV
Piperonyl butoxide	51-03-6	$C_{19}H_{30}O_5$	benzodioxole	8	070
Pirimicarb	23103-98-2	$\mathrm{C}_{11}\mathrm{H}_{18}\mathrm{N}_4\mathrm{O}_2$	carbamate	14	580
Pirimicarb desmethyl	30614-22-3	$C_{10}H_{16}N_4O_2$	dimethylcarbamate metabolite	14	873
Pirimiphos methyl	29232-93-7	$\mathrm{C}_{11}\mathrm{H}_{20}\mathrm{N}_{3}\mathrm{O}_{3}\mathrm{PS}$	phosphorothioate	11	562
Prallethrin	23031-36-9	$C_{19}H_{24}O_3$	pyrethroid	8	ADC
Pretilachlor	51218-49-6	$C_{17}H_{26}CINO_2$	chloroacetanilide	1	892
Primisulfuron	113036-87-6	$\mathrm{C}_{14}H_{10}F_4N_4O_7S$	pyrimidinylsulfonyl urea	16	AHA
Prochloraz	67747-09-5	$\mathrm{C_{15}H_{16}Cl_{3}N_{3}O_{2}}$	imidazole	9	833
Procymidone	32809-16-8	$C_{13}H_{11}CI_2NO_2$	dicarboximide	1	593
Prodiamine	29091-21-2	$\mathrm{C}_{13}H_{17}F_3N_4O_4$	dinitroaniline	7	814
Profenofos	41198-08-7	$C_{11}H_{15}BrClO_3PS$	phosphorothioate	11	224
Profluralin	26399-36-0	$\mathrm{C}_{14}H_{16}F_3N_3O_4$	dinitroaniline	7	A53
Profoxydim	139001-49-3	$C_{24}H_{32}CINO_4S$	cyclohexene oxime	28	ANB
Prohexadione calcium	127277-53-6	$\mathrm{C_{20}H_{22}CaO_{10}}$	unclassified plant growth regulator	99	ALO
Promecarb	2631-37-0	$C_{12}H_{17}NO_2$	phenyl methylcarbamate	14	385

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Prothioconazole	178928-70-6	$\mathrm{C_{14}H_{15}Cl_2N_3OS}$	conazole	1	AHJ
Prometon	1610-18-0	$C_{10}H_{19}N_5O$	triazine	9	942
Prometryn	7287-19-6	$C_{10}H_{19}N_5S$	triazine	9	249
Pronamide (propyzamide)	23950-58-5	$C_{12}H_{11}CI_2NO$	amide	1	540
Propachlor	1918-16-7	$C_{11}H_{14}CINO$	chloroacetanilide	1	675
Propachlor oxanilic acid	70628-36-3	$C_{11}H_{13}NO_3$	chloroacetanilide metabolite	20	AFA
Propamocarb hydrochloride	25606-41-1	$C_9H_{20}N_2O_2$	carbamate	14	AFU
Propanil	709-98-8	$C_9H_9Cl_2NO$	anilide	1	341
Propaquizafop	111479-05-1	$C_{22}H_{22}CIN_3O_5$	aryloxyphenoxypropionic acid	17	ALK
Propargite	2312-35-8	$C_{19}H_{26}O_4S$	sulfite	1	623
Propazine	139-40-2	$C_9H_{16}CIN_5$	triazine	9	333
Propetamphos	31218-83-4	$C_{10}H_{20}NO_4PS$	phosphorothioate	11	636
Propham	122-42-9	$C_{10}H_{13}NO_2$	carbamate	14	310
Propiconazole	60207-90-1	$C_{15}H_{17}CI_2N_3O_2$	conazole	1	264
Propoxur	114-26-1	$C_{11}H_{15}NO_3$	carbamate	14	162
Propoxycarbazone	145026-81-9	$C_{15}H_{18}N_4O_7S$	triazolone	1	AKK
Proquinazid	189278-12-4	$C_{14}H_{17}IN_2O_2$	unclassified	17	AMM
Prosulfuron	94125-34-5	$\mathrm{C_{15}H_{16}F_{3}N_{5}O_{4}S}$	triazinylsulfonyl urea	16	AEG
Prothioconazole	178928-70-6	C ₁₄ H ₁₅ Cl ₂ N ₃ OS	conazole	1	AHJ
Prothiofos	34643-46-4	$\mathrm{C_{11}H_{15}Cl_{2}O_{2}PS_{2}}$	phenylorganothiophosphate	11	613
Pymetrozine	123312-89-0	$C_{10}H_{11}N_5O$	azomethine	9	ABF
Pyraclofos	77458-01-6	$C_{14}H_{18}CIN_2O_3PS$	heterocyclic organothiophosphate	11	F01
Pyraclostrobin	175013-18-0	$C_{19}H_{18}CIN_3O_4$	strobilurin	22	B61
Pyraflufen	129630-17-7	$\mathrm{C_{13}H_9Cl_2F_3N_2O_4}$	phenylpyrazole	1	ALY
Pyraflufen ethyl	129630-19-9	$\mathrm{C_{15}H_{13}Cl_2F_3N_2O_4}$	phenoxypyrazole	1	AGB
Pyrasulfotole	365400-11-9	$\mathrm{C}_{14}H_{13}F_3N_2O_4S$	benzoylpyrazole	1	AHK
Pyrazon (Chloridazon)	1698-60-8	$C_{10}H_8CIN_3O$	pyridazinone	17	595

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code
Pyrazophos	13457-18-6	$\mathrm{C_{14}H_{20}N_{3}O_{5}PS}$	organophosphate	11	553
Pyrethrins	8003-34-7	$C_{21}H_{27}O_4$	pyrethrum, botanical	8	075
Pyridaben	96489-71-3	$C_{19}H_{25}CIN_2OS$	pyridazinone	17	B56
Pyridalyl	179101-81-6	$\mathrm{C_{18}H_{14}Cl_{4}F_{3}NO_{3}}$	pyridine	17	AHU
Pyridaphenthion	119-12-0	$\mathrm{C_{14}H_{17}N_2O_4PS}$	organophosphate	11	961
Pyrimethanil	53112-28-0	$\mathrm{C_{12}H_{13}N_{3}}$	pyrimidine	17	B16
Pyriproxyfen	95737-68-1	$\mathrm{C_{20}H_{19}NO_{3}}$	pyridine	17	B24
Pyroxasulfone	447399-55-5	$\mathrm{C}_{12}H_{14}F_{5}N_{3}O_{4}S$	pyrazole	1	AMO
Quinalphos	13593-03-8	$\mathrm{C_{12}H_{15}N_2O_3PS}$	organothiophosphate	11	661
Quinchlorac	84087-01-4	$C_{10}H_5CI_2NO_2$	quinolinecarboxylic acid	20	B29
Quinoxyfen	124495-18-7	$C_{15}H_8Cl_2FNO$	pyridine	17	B57
Quintozene (PCNB)	82-68-8	$C_6 Cl_5 NO_2$	benzene ring	3	304
Quizalofop	76578-12-6	$\mathrm{C_{17}H_{13}CIN_2O_4}$	aryloxyphenoxypropionic hebicide	20	ALZ
Quizalofop ethyl	76578-14-8	$C_{19}H_{17}CIN_2O_4$	aryloxyphenoxypropionic acid	20	750
Resmethrin	10453-86-8	$C_{22}H_{26}O_3$	pyrethroid	8	556
RH 9129	146887-38-9	$\mathrm{C_{19}H_{16}N_{3}ClO_{2}}$	fenbuconazole metabolite	1	A54
RH 9130	146887-37-8	$\mathrm{C_{19}H_{16}N_3ClO_2}$	fenbuconazole metabolite	1	A55
Rimsulfuron	122931-48-0	$C_{14}H_{17}N_5O_7S_2$	sulfonyl urea	16	AJF
Rotenone	83-79-4	$C_{23}H_{22}O_{6}$	botanical insecticide	8	020
S-(2-hydroxy)propyl EPTC	759-94-4	$C_9H_{19}NOS$	thiocarbamate	14	ACO
Saflufenacil	372137-35-4	$\mathrm{C_{17}H_{17}CIF_4N_4O_5S}$	urea	16	AHZ
Sedaxane	874967-67-6	$\mathrm{C_{18}H_{19}F_2N_3O}$	pyrazole	1	ALB
Sethoxydim	74051-80-2	$C_{17}H_{29}NO_3S$	cyclohexene oxime	28	AEV
Sethoxydim sulfoxide	114480-24-9	$C_{17}H_{29}NO_4S$	cyclohexene oxime	28	AJR
Siduron	1982-49-6	$C_{14}H_{20}N_2O$	urea	16	ACT
Simazine	122-34-9	$C_7H_{12}CIN_5$	triazine	9	149
Simetryn	1014-70-6	$C_8H_{15}N_5S$	triazine	9	837

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code
Spinetoram	187166-40-1	$C_{42}H_{69}NO_{10} + C_{43}H_{69}NO_{10}$	spinosyn (macrocyclic lactone)	29	AGY
Spinosad	168316-95-8	$C_{41}H_{65}NO_{10} + C_{42}H_{67}NO_{10}$	spinosyn (macrocyclic lactone)	29	ABB
Spirodiclofen	148477-71-8	$C_{21}H_{24}CI_2O_4$	tetronic acid	27	B85
Spiromesifen	283594-90-1	$C_{23}H_{30}O_4$	tetronic acid	27	AGT
Spiromesifen enol metabolite	148476-30-6	$C_{17}H_{20}O_3$	tetronic acid metabolite	27	AGU
Spiromesifen, total (including enol metabolite)	283594-90-1	$C_{23}H_{30}O_4 + C_{17}H_{20}O_3$	tetronic acid	27	AFW
Spirotetramat	203313-25-1	$C_{21}H_{27}NO_5$	tetramic acid insecticide	27	AHM
Spiroxamine	118134-30-8	$C_{18}H_{35}NO_2$	unclassified	99	AJY
Sulfallate	95-06-7	$C_8H_{14}CINS_2$	thiocarbamate	14	323
Sulfentrazone	122836-35-5	$\mathrm{C_{11}H_{10}Cl_2F_2N_4O_3S}$	triazole sulfonamide	1	AAY
Sulfometuron methyl	74222-97-2	$\mathrm{C_{15}H_{16}N_4O_5S}$	sulfonyl urea	16	ACP
Sulfosulfuron	141776-32-1	$\mathrm{C_{16}H_{18}N_6O_7S_2}$	pyrimidinylsulfonyl urea	16	ADS
Sulfotep	3689-24-5	$C_8H_{20}O_5P_2S_2$	organophosphate	11	311
Sulfoxaflor	946578-00-3	$\mathrm{C_{10}H_{10}F_{3}N_{3}OS}$	sulfoximine	99	ALS
Sulprofos	35400-43-2	$\mathrm{C_{12}H_{19}O_{2}PS_{3}}$	organophosphate	11	609
Sulprofos oxygen analog	38527-90-1	$\mathrm{C_{12}H_{19}O_{3}PS_{2}}$	oxon	11	ACQ
ТСМТВ	21564-17-0	$C_9H_6N_2S_3$	benzothiazole	17	793
Tebuconazole	107534-96-3	$C_{16}H_{23}CIN_3O$	conazole	1	A58
Tebufenozide	112410-23-8	$C_{22}H_{28}N_2O_2$	diacylhydrazine	16	ABG
Tebufenpyrad	119168-77-3	$C_{18}H_{24}CIN_3O$	pyrazole	1	AJZ
Tebupirimfos	96182-53-5	$C_{13}H_{23}N_2O_3PS$	organophosphate	11	A59
Tebupirimfos oxygen analog	NA ^[2]	$\mathrm{C}_{13}H_{23}N_{2}O_{4}P$	oxon	11	ACR
Tebuthiuron	34014-18-1	$C_9H_{16}N_4OS$	urea	16	780
Tecnazene	117-18-0	$C_6HCl_4NO_2$	nitrobenzene	3	147
Teflubenzuron	83121-18-0	$\mathrm{C}_{14}H_{6}CI_{2}F_{4}N_{2}O_{2}$	benzoylphenylurea	16	AKA
Tefluthrin	79538-32-2	$C_{17}H_{14}CIF_7O_2$	pyrethroid	8	B26
TEPP	107-49-3	$C_8H_{20}O_7P_2$	organophosphate	11	088

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code	
Tepraloxydim	149979-41-9	$C_{17}H_{24}CINO_4$	cyclohexene oxime	28	AHL	
Terbacil	5902-51-2	$C_9H_{13}CIN_2O_2$	uracil	16	152	
Terbufos	13071-79-9	$C_9H_{21}O_2PS_3$	phosphorothioate	11	205	
Terbufos oxygen analog	56070-14-5	$C_9H_{21}O_3PS_2$	oxon	11	A60	
Terbufos oxygen analog sulfone	56070-15-6	$C_9H_{21}O_5PS_2$	organophosphate	11	752	
Terbufos sulfone	56070-16-7	$C_9H_{21}O_4PS_3$	sulfone	11	963	
Terbufos Sulfoxide	10548-10-4	$C_9H_{21}O_4PS_2$	organophosphate	11	AMP	
Terbutryn	886-50-0	$C_{10}H_{19}N_5S$	methylthiotriazine	9	738	
Terbuthylazine	5915-41-3	$C_9H_{16}CIN_5$	chlorotriazine	9	678	
Tetrachlorvinphos	22248-79-9	$C_{10}H_9CI_4O_4P$	chlorethylene phosphate	11	176	
Tetraconazole	112281-77-3	$\mathrm{C_{13}H_{11}Cl_{2}F_{4}N_{3}O}$	conazole	1	B72	
Tetradifon	116-29-0	$C_{12}H_6CI_4O_2S$	bridged biphenyl	3	108	
Tetrahydrophthalimide (THPI) ^[1]	1469-48-3	$C_8H_9NO_2$	phthalimide	1	624	
Tetramethrin	7696-12-0	$\mathrm{C_{19}H_{25}NO_4}$	pyrethroid	8	947	
Thiabendazole	148-79-8	$C_{10}H_7N_3S$	benzimidazole	1	157	
Thiacloprid	111988-49-9	$C_{10}H_9CIN_4S$	neonicotinyl	1	B68	
Thiamethoxam	153719-23-4	$C_8H_{10}CIN_5O_3S$	neonicotinyl	1	B43	
Thiazopyr	117718-60-2	$C_{16}H_{17}F_5N_2O_2S$	pyridine	17	B12	
Thidiazuron	51707-55-2	C ₉ H ₈ N ₄ OS	plant growth regulator	16	794	
Thiencarbazone methyl	317815-83-1	$C_{12}H_{14}N_4O_7S_2$	triazolone	4	AKL	
Thifensulfuron	79277-67-1	$C_{11}H_{11}N_5O_6S_2$	sulfonyl urea	16	AEF	
Thifensulfuron methyl	79227-27-3	$C_{12}H_{13}N_5O_6S_2$	triazinylsulfonyl urea	16	AEQ	
Thiobencarb	28249-77-6	C ₁₂ H ₁₆ CINOS	thiocarbamate	14	726	
Thiodicarb	59669-26-0	$\mathrm{C_{10}H_{18}N_4O_4S_3}$	carbamate	14	943	
Thionazin	297-97-2	$C_8H_{13}N_2O_3PS$	organophosphate	11	250	
Thiophanate methyl	23564-05-8	$\mathrm{C}_{12}H_{14}N_4O_4S_2$	carbamate	14	611	
Thymol	89-83-8	C ₁₀ H ₁₄ O	phenol	3	ALG	

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Tolclofos methyl	57018-04-9	$C_9H_{11}CI_2O_3PS$	organophosphate	11	B70
Tolfenpyrad	129558-76-5	$\mathrm{C}_{21}\mathrm{H}_{22}\mathrm{CIN_3O_2}$	pyrazole	1	ANC
Tolyfluanid	731-27-1	$\mathrm{C_{10}H_{13}Cl_2FN_2O_2S_2}$	phenylsulfamide	1	649
Topramezone	210631-68-8	$\mathrm{C_{16}H_{17}N_{3}O_{5}S}$	benzoylpyrazole	1	AMO
Toxaphene	8001-35-2	$C_{10}H_{10}CI_8$	organochlorine	3	090
Tralomethrin	66841-25-6	$C_{22}H_{19}Br_4NO_3$	pyrethroid	8	755
Triadimefon	43121-43-3	$\mathrm{C}_{14}H_{16}CIN_3O_2$	conazole	1	608
Triadimenol	55219-65-3	$\mathrm{C}_{14}H_{18}CIN_3O_2$	conazole	1	638
Triallate	2303-17-5	$C_{10}H_{16}CI_3NOS$	thiocarbamate	14	621
Triasulfuron	82097-50-5	$\mathrm{C_{14}H_{16}CIN_5O_5S}$	sulfonyl urea	16	ADP
Triazole acetic acid	28711-29-7	$C_4H_6N_3O_2$	triazole metabolite	1	ADX
Triazole alanine	86362-20-1	$C_5H_8N_4O_2$	triazole metabolite	1	ADW
Triazophos	24017-47-8	$\mathrm{C_{12}H_{16}N_3O_3PS}$	organothiophosphate	11	536
Tribenuron methyl	101200-48-0	$\mathrm{C_{15}H_{17}N_5O_6S}$	triazinylsulfonyl urea	16	ACS
Trichlorfon (as dichlorvos)	52-68-6	$C_4H_8CI_3O_4P$	phosphate	11	130
Trichloronate	327-98-0	$\mathrm{C_{10}H_{12}Cl_{3}O_{2}PS}$	organothiophosphate	11	569
Triclopyr	55335-06-3	$C_7H_4CI_3NO_3$	acetic acid	20	731
Tricyclazole	41814-78-2	$C_9H_7N_3S$	benzothiazole	4	804
Trifloxystrobin	221007-60-9	$C_{20}H_{19}F_3N_2O_4$	strobilurin	22	B79
Trifloxysulfuron	145099-21-4	$\mathrm{C}_{14}H_{14}F_3N_5O_6S$	sulfonyl urea	16	AJG
Triflumizole	68694-11-1	$\mathrm{C_{15}H_{15}CIF_{3}N_{3}O}$	conazole	1	A61
Trifluralin	1582-09-8	$\mathrm{C_{13}H_{16}F_3N_3O_4}$	dinitroaniline	7	151
Triforine	26644-46-2	$\mathrm{C_{10}H_{14}Cl_6N_4O_2}$	formamide	1	915
Triticonazole	131983-72-7	$C_{17}H_{20}CIN_3O$	conazole	1	ADR
Uniconazole	83657-22-1	$C_{15}H_{18}CIN_3O$	conazole	1	AJJ
Vernolate	1929-77-7	$C_{10}H_{21}NOS$	thiocarbamate	14	201
Vinclozolin	50471-44-8	$C_{12}H_9CI_2NO_3$	dichloroanilide	1	529

Compound Name	CAS#	Molecular Formula	Chemical Family	Group	Pesticide Code
Zoxamide	156052-68-5	$\mathrm{C_{14}H_{16}Cl_{3}NO_{2}}$	benzamide	1	B44

[1] Metabolite of captan and captafol.

[2] Not available.

EPA, Codex, and Food and Drug Administration (FDA) Pesticide Analytical Manual (PAM) Commodity Groupings

(Commodity	PDP Code	PDP Commodity Group	EPA	Codex		PAM	
Almonds		AL	Cereal Grains (High Oil)	Tree nuts	Tree nuts		N/A	
	Fruit	AP			Pome fruits		Med. Sugar	
	Juice	AJ			Fruit juice	-	Med. Sugar	
Apples	Sauce	AC	Fruits and Vegetables	Pome fruits	Manufactured food single ingredient	Non-fatty	High Sugar	
	Baby Food	IA			Manufactured food single ingredient			
	Single serving	AX	•		Pome fruits	1	Med. Sugar	
Asparagus	olligio solving	AS	Fruits and Vegetables	Miscellaneous	Stalk & stem vegs.	Non-fatty	Low Sugar	
Avocado		AV	Fruits and Vegetables	Miscellaneous		Fatty	Low Sugar	
Bananas		BN	Fruits and Vegetables	Miscellaneous	Assorted tropical & sub-tropical fruits - inedible peel		High Sugar	
Barley		BY	Cereal Grains (Low Oil)	Cereal grains	Cereal grains	Non-fatty	Low Water	
Dariey	Black	BC		Cereal grains		Non-fatty	Low Water	
	Garbanzo (Chick	ZB	•			-	Low Water	
Beans,		BC	Fruits and Vegetables	Legume vegs.	Legume vegs.	Fatty Non-fatty	Low Sugar	
canned	Kidney	BC	Fruits and vegetables	Leguine vegs.	Leguine vegs.			
	Pinto	BC				Non-fatty	Low Water	
						Non-fatty	Low Sugar	
Beets		BT	Fruits and Vegetables	Root & Tuber vegs.	Root & tuber vegs.	Non-fatty	N/A	
	Adipose	BA					-	
Beef	Liver	BL	Animal Tissue/High Protein	Meat	Meat		N/A	
	Muscle	BM	, j					
Blueberry		BB	Fruits and Vegetables	Berry & Small Fruit	Berries & other small fruits	Non-fatty	Med. Sugar	
Broccoli		BR	Fruits and Vegetables	Brassica leafy vegs.	Brassica leafy vegs.	Non-fatty	Low Sugar	
Butter		BU	Dairy Products	Dairy	Derived milk products	Fatty	Low Water	
Cabbage		CG	Fruits and Vegetables	Brassica leafy vegs.	Brassica leafy vegs.	Non-fatty	Low Sugar	
Cantaloupe		CN	Fruits and Vegetables	Cucurbits	Cucurbits	Non-fatty	Med. Sugar	
	Fresh	CR		Root & tuber	Root & tuber vegs.	· · · ·		
Carrots	Baby Food	IC	Fruits and Vegetables	vegs.	Manufactured food single ingredient	Non-fatty	Med. Sugar	
Cauliflower	Babyrood	CF	Fruits and Vegetables	Brassica leafy vegs.	Brassica leafy vegs.	Non-fatty	Low Sugar	
Celery		CE	Fruits and Vegetables	Leafy vegs.	Stalk & stem vegs.	Non-fatty	Low Sugar	
Cherries, Sw	(oot	CH	Fruits and Vegetables	Stone fruits	Stark & Sterri Vegs.	Non-fatty	Med. Sugar	
Cilantro	/eel	CL	Fruits and Vegetables	Herbs & Spices	Herbs & Spices	Non-fally	N/A	
Cilantio	0			Herbs & Spices		F - 44 -		
Carra	Grain	CO	Cereal Grains (Low Oil)		Cereal grains	Fatty	Low Water	
Corn	Sweet	CS	Fruits and Vegetables	Cereal grains	Cereal grains	Non-fatty	Med. Sugar	
	Syrup	CY	Single Commodities		Derived edible plant products		N/A	
Cranberry		CA	Fruits and Vegetables	Berry & Small Fruit	Berries & other small fruits	Non-fatty	N/A	
Cream, heav	/y	CM	Dairy Products	Dairy	Derived milk products	Fatty	Low Sugar	
Cucumbers		CU	Fruits and Vegetables	Cucurbits	Cucurbits	Non-fatty	Low Sugar	
Egg		EG	Animal Tissue/High Protein	Miscellaneous	Poultry products	Non-fatty	N/A	
Eggplant		EP	Fruits and Vegetables	Fruiting vegs.	Fruiting vegs.	Non-fatty	Low Sugar	
Fish, Catfish		FC	Single Commodities or	Miscellaneous	Aquatic animal products	Fatty	No sugar	
Fish, Salmor		FS	Animal Tissue/High Protein	Miscellaneous	Aquatic animal products	Fatty	No sugar	
Grapefruit		GF	Fruits and Vegetables	Citrus fruits	Citrus fruits		Med. Sugar	

EPA, Codex, and Food and Drug Administration (FDA) Pesticide Analytical Manual (PAM) Commodity Groupings

Co	ommodity	PDP Code	PDP Commodity Group	EPA	Codex		PAM
Grapes	Fruit	GR	Fruits and Vegetables	Berry & Small	Berries & other small fruits	Non-fatty	High Sugar
Juice		GJ		Fruit	Fruit juice	Non-fatty	Med. Sugar
Green Beans	Raw, fresh	GB	Fruits and Vegetables	Legume vegs.	Legume vegs.	Non-fatty	Low Sugar
Oreen Deans	Baby Food	IG	Traits and vegetables	Leguine vegs.	Manufactured food single ingredient	Non-latty	LOW Ougar
Greens	Collard	GS	Fruits and Vegetables	Brassica leafy	Leafy vegs. (including Brassica leafy vegs.)	Non-fatty	Low Sugar
Greens	Kale	GK	Truits and vegetables	vegs.	Leary vegs. (including brassica leary vegs.)	Non-latty	LOW Sugar
Honey		HY	Single Commodities	Miscellaneous		Non-fatty	High Sugar
Honey Dew Me	elon	HD	Fruits and Vegetables	Cucurbits	Cucurbits	Non-fatty	N/A
Infant formula,	dairy-based	DF	Single Commodities		Manufactured food multiple ingredient	N/A	N/A
Infant formula,	soy-based	YF	Single Commodities		Manufactured food multiple ingredient	N/A	N/A
Kiwi Fruit, Fres	sh	KW	Fruits and Vegetables	Berry & Small Fruit	Assorted tropical & sub-tropical fruits - inedible peel	Non-fatty	Med. Sugar
Lemons		LM	Fruits and Vegetables	Citrus fruits	Citrus fruits	Non-fatty	Low Sugar
	Bunch	LT		Leafy vegs.	Leafy vegs.		
Lettuce	Bagged	LB	Fruits and Vegetables			Non-fatty	Low Sugar
Mangoes		MA	Fruits and Vegetables	Miscellaneous	Assorted tropical & sub-tropical fruits - inedible peel	Non-fattv	Med. Sugar
Milk, whole		MK	Dairy Products	Dairy	Milks	Fatty	Low Sugar
Mushrooms		MU	Fruits and Vegetables	Edible fungi	Fruiting vegs.	Non-fatty	Low Sugar
Nectarines	-	NE	Fruits and Vegetables	Stone fruits	Stone fruits	Non-fatty	Med. Sugar
Oats			Cereal Grains (Low Oil)	Cereal grains	Cereal grains	Fatty	Low Water
0	Bulb	ON		Bulb vegs.	Dulh years	Non-fatty	
Onions	Green	GO	Fruits and Vegetables		Bulb vegs.		Low Sugar
0	Fruit	OG	Fruits and Vegetables	Citrus fruits	Citrus fruits	Non foth	
Orange	Juice	OJ			Fruit juice	Non-fatty	Med. Sugar
Olives	Fruit	OL	Fruits and Vegetables	Tropical & sub- tropical fruits -	Tropical & sub-tropical fruits - edible peel	N/A	N/A
	Canned	CC			Stone fruits		
B	Fruit	PC					
Peaches	Single serving	CX	Fruits and Vegetables	Stone fruits		Non-fatty	Med. Sugar
	Baby Food	IH			Manufactured food single ingredient		
Papaya	• •	YA	Fruits and Vegetables	Miscellaneous		Non-fatty	Med. Sugar
Peanut Butter		PB	Cereal Grains (High Oil)	Miscellaneous	Manufactured food single ingredient	Fatty	Med. Sugar
	Fruit	PE			Pome fruits	Non-fatty	Med. Sugar
	Juice	PJ			Derived edible plant products	Í	N/A
Pears	Canned	CP	Fruits and Vegetables	Pome fruits			
	Single serving	PX	-		Pome fruits	Non-fatty	Med. Sugar
	Baby Food	IP			Manufactured food single ingredient	1	j j
D	Vegetable	PS		1	Legume vegs.	Nam fatt	
Peas	Baby Food	IE	Fruits and Vegetables	Legume vegs.	Manufactured food single ingredient	Non-fatty	Low Sugar
D	Bell	PP		E		Nam Car	1
Peppers	Hot	HP	Fruits and Vegetables	Fruiting vegs.	Fruiting vegs.	Non-fatty	Low Sugar
D: .	Fruit	PN		Tropical & sub-			
Pineapples	Canned	NC	Fruits and Vegetables	tropical fruits -	Assorted tropical & sub-tropical truits - inedible peel		Med. Sugar

EPA, Codex, and Food and Drug Administration (FDA) Pesticide Analytical Manual (PAM) Commodity Groupings

C	ommodity	PDP Code	PDP Commodity Group	EPA	Codex	[PAM	
Plums	Dried	PD	Fruits and Vegetables	Stone fruits	Stone fruits	Non-fatty	Med. Sugar	
lullis	Fruit	PU				Non latty	Nicu. Ougai	
Pork	Adipose	KA	Animal Tissue/High Protein	Meat	Meat		N/A	
-	Muscle	KM	-				-	
Potatoes	- I	PO	Fruits and Vegetables	Root & tuber	Root & tuber vegs.	Non-fatty	Low Sugar	
	Adipose	PA						
	Breast	PR						
Poultry	Liver	PL	Animal Tissue/High Protein	Meat	Poultry meat		N/A	
	Muscle	PM						
	Thigh	PT						
Raspberries	Fresh	RS	Fruits and Vegetables	Berry & Small	Berries & other small fruits	Non-fatty	N/A	
	Frozen	RZ		Fruit				
Raisins		RA	Single Commodities	Berry & Small Fruit	Dried fruits	Non-fatty	High sugar	
Rice		RI	Cereal Grains (Low Oil)	Cereal grains	Cereal grains	Non-fatty	Low Sugar	
Snap Peas		SN	Fruits and Vegetables	Legume vegs.	Legume vegs.	Non-fatty	Low Sugar	
Soybeans, Gr	ain	SY	Cereal Grains (High Oil)	Legume vegs.	Legume vegs.	Fatty	Med Sugar	
	Leafy	SP	Fruits and Vegetables	Leafy vegs.	Leafy vegs.	Non-fatty	Low Sugar	
	Canned	SC	Truits and vegetables	Leary vegs.	Manufactured food single ingredient			
	Summer	SS	Fruits and Vegetables	Cucurbits				
Squash	Winter	WS			Cucurbits	Non-fatty	Low Sugar	
	Winter, frozen	WZ						
Strawberries	Fresh	ST	Fruits and Vegetables	Berry & Small	Berries & other small fruits	Non-fatty	Med. Sugar	
	Frozen	SZ		Fruit		ŗ		
Sweet	Raw, fresh	SW	Fruits and Vegetables	Root & tuber	Root & tuber vegs.		Med. Sugar	
Potatoes	Baby Food	IS			Manufactured food single ingredient	Non-fatty	Med. Sugar	
Tangerines	-	TA	Fruits and Vegetables	Citrus fruits	Citrus fruits			
	Cherry/Grape	СТ						
Tomatoes	Fresh	TO	Fruits and Vegetables	Fruiting vegs.	Fruiting vegs.	Non-fatty	Low Sugar	
- on allo o o	Canned	TC						
	Paste	TP	Single Commodities		Manufactured food single ingredient		N/A	
	Bottled	WB						
Water	Drinking	WR	Water	Miscellaneous	N/A	N/A	N/A	
	Ground	WG					,	
Untreated		WU						
Watermelon	-	WM	Fruits and Vegetables	Cucurbits	Cucurbits	Non-fatty	Med. Sugar	
Wheat	Grain	WH	Cereal Grains (Low Oil)	Cereal grains	Cereal grains		N/A	
	Flour	WF			Cereal grains, milling fraction			

Food and Drug Administration Information

C	Commodity		% Water ¹	% Sugar ¹	рН ²	
Almonds						
	Fruit	0.36	83.93	11.5	3.30 -4.00	
Apples	Juice	0.11	87.93	10.9	3.35-4.00	
	Sauce	0.18	79.58	16.5	3.10-3.60	
Asparagus		0.22	92.25	2.1	6.00-6.70	
Avocado		8.87-17.33	72.56-79.73	0.9	6.27-6.58	
Bananas		0.48	74.26	18.4	4.50-5.20	
Barley		1.16	10.09		5.19-5.32	
	Black	1.42	11.02		5.78-6.02	
	Garbanzo (Chick pea)	6.04	11.53	3.8	6.48-6.80	
Beans	Kidney	1.06	11.75	0.0	5.40-6.00	
Deans	Pinto	1.13	10.95		3.40-0.00	
		0.1				
Beef	Baby Food	0.1	92.5			
		0.00	00.45		5 00 0 00	
Beets		0.06	92.15	7.0	5.30-6.60	
Blueberry		0.38	84.61	7.3	3.12-3.33	
Broccoli		0.35	90.69	1.6	6.30-6.52	
Butter		81.11	17.94			
Cabbage		0.18	92.52	2.7	5.20-6.80	
Cantaloupe		0.28	89.78	8.1	6.13-6.58	
Carrots		0.19	87.79	6.6	5.88-6.40	
Cauliflower		0.18	92.26	2.2	5.60	
Celery		0.14	94.64	1	5.70-6.00	
Cherries, sweet		0.96	80.76	14.6	4.01-4.54	
Cilantro						
	Grain	2.08	10			
Corn	Sweet	1.18	75.96	5.4	5.90-7.30	
	Syrup					
Cranberry		0.2	86.54			
Cream, heavy		37	57.71	2.8	6.50-6.68	
Cucumbers		0.13	96.05	2.3	5.12-5.78	
Eggplant		0.1	91.93	3.4	5.50-6.50	
Fish, catfish		4.26	76.39	0	0.00 0.00	
Fish, salmon		3.4-10.44	68.5-76.35	0		
Grapefruit		0.1	90.89	6.2	3.00-3.75	
Giapeiruit	En .: it					
Grapes	Fruit	0.35	81.3	16.4	2.90-3.82	
Croop Beans	Juice	0.08	84.12	14.2	E 00	
Green Beans	Callard	0.12	90.27	2.6	5.60	
Greens	Collard	0.22	90.55		0.00.0.00	
	Kale	0.7	84.46	2.2	6.36-6.80	
Honey		0	17.2	81.9	3.70-4.20	
Honey Dew Melon		0.1	89.66		6.00 - 6.67	
Infant formula, dai						
Infant formula, soy	/-based	0.19				
	Lettuce		95.89	1.8	5.80-6.15	
Mangoes		0.27	81.71	14.8	3.40 - 4.80	
Milk, whole		3.66	87.69	4.9	6.40-6.80	
Mushrooms		0.42	91.81	1.8	6.00-6.70	
Nectarines		0.46	86.28	8.5	3.92-4.18	
Oats		6.9	8.22	5.9		
Olives						
	Bulb	0.16	89.68	4.1	5.30-5.85	
Onions	Green	0.10	89.83	3.2	6.20	
<u> </u>	Fruit	0.19	86.75	8.9	3.60-4.34	
Oranges	Juice	0.12	88.3	10.2	3.30-4.19	
	Juice	0.2	00.3	10.2	3.30-4.19	

Food and Drug Administration Information

	Commodity	% Fat ¹	% Water ¹	% Sugar ¹	pH ²
Papaya		0.14	88.83		
Peaches		0.09	87.66	8.7	3.30-4.05
Peanut Butter		49.98	1.42	7.8	6.28
Pears	Fruit	0.4	83.81	10.5	3.50-4.60
	Juice				
Peas		0.4	78.86	4.5	5.70-6.70
Peppers	Bell	0.19	92.19	2.5	5.20-5.93
reppers	Hot	0.2	87.74		4.65 - 5.45
Pineapples		0.43	86.5	11.9	3.20-4.00
Plums		0.62	85.2	7.5	2.80-4.30
Pork					
Potatoes	Potatoes		78.96	1.0	5.40-5.90
Poultry					
Raisins		0.46	15.42	61.7	3.80-4.10
Raspberries		0.55	86.57		3.18-3.95
Rice		0.58	12.89	0.5	6.06.70
Soybeans		19.94	8.54	6.6	
Spinach		0.35	91.58	0.4	5.50-6.80
Squash	Summer	0.21	93.68	2.2	5.79-6.10
Squash	Winter	0.23	88.72	2.2	5.18-6.49
Strawberries		0.37	91.57	5.7	3.00-3.90
Sweet Potatoes	i	0.3	72.84	5.0	5.30-5.60
Tangerines		0.19	87.6		
Tomatoes	Fresh	0.33	93.76	3.0	4.30-4.90
TUTTALUES	Paste				3.50-4.70
Watermelon		0.43	91.51	9	5.18-5.60
Wheat	Grain				
VITEAL	Flour				

1 = Pesticide Analytical Manual (PAM) data

2 = Center for Food Safety and Applied Nutrition data

Data not avalilable Fatty (>2% fat) Non-fatty (<2% fat)

Low H₂O (<75%)

Low sugar (<5%)

Med sugar (5-15%) High sugar (>15%)

USDA, AMS Pesticide Data Program Verification of Limits of Detection (LODs)

Comm	nodity: _
Date:	
Lab:	

Note: During method validation, two spikes are required; if this form is used to record annual spike verification, only one spike is required.

Pesticide/Compound	Amt Spk	LOD Spike Recovered (yes/no or +/-)			
	Units =	Spike 1	Spike 2		

