# DATA REQUIREMENTS FOR IMPORT MRLs ON USE SITE CATEGORY

| Data Code | Title  | Data required | Conditions   | Present/<br>Absent |
|-----------|--|---------------|--|--------------------|
| 0         | Index  | R             |  | Hosent             |
| 1         | Label  | R             |  |                    |
| 2         | Chemistry requirements for the registration of a technical grade of active ingredient (TGAI) or an |               |  |                    |
|           | integrated system product.   |               |  |                    |
| 2.1       | Applicant's Name and Office Address  | R             |  |                    |
| 2.2       | Manufacturer's Name and Office Address and   | R             |  |                    |
|           | Manufacturing Plant's Name and Address   |               |  |                    |
| 2.3       | Product Trade Name   | R             |  |                    |
| 2.3.1     | Other Names  | R             |  |                    |
| 2.4       | Common Name  | R             |  |                    |
| 2.5       | Chemical Name  | R             |  |                    |
| 2.6       | Chemical Abstracts Registry Number   | R             |  |                    |
| 2.7       | Structural Formula   | R             |  |                    |
| 2.8       | Molecular Formula  | R             |  |                    |
| 2.9       | Molecular Weight   | R             |  |                    |
| 2.10      | Canadian Patent Status   | R             |  |                    |
| 2.11      | Manufacturing Methods for the TGAI   |               |  |                    |
| 2.11.1    | Manufacturing Summary  | R             |  |                    |
| 2.11.2    | Description of Starting Materials  | R             |  |                    |
| 2.11.3    | Detailed Production Process Description  | R             |  |                    |
| 2.11.4    | Discussion of Formation of Impurities  | R             |  |                    |
| 2.12      | Specifications   |               |  |                    |
| 2.12.1    | Establishing Certified Limits  | CR            | A justification must be provided if standard limits are not met. |                    |
| 2.12.2    | Control Product Specification Form   | R             |  |                    |
| 2.13      | Preliminary Analysis   |               |  |                    |
| 2.13.1    | Methodology/Validation   | R             |  |                    |
| 2.13.2    | Confirmation of Identity   | R             |  |                    |
| 2.13.3    | Batch Data   | R             |  |                    |
| 2.13.4    | Impurities of Toxicological Concern  | CR            | If applicable  |                    |
| 2.14      | Chemical and Physical Properties   |               |  |                    |
| 2.14.1    | Colour   | R             |  |                    |
| 2.14.2    | Physical State   | R             |  |                    |
| 2.14.3    | Odour  | R             |  |                    |
| 2.14.4    | Melting Point / Melting Range  | R             | Solid at room temperature.                                       |                    |
| 2.14.5    | Boiling Point / Boiling Range  | R             | Liquid at room temperature.                                      |                    |
| 2.14.6    | Density or Specific Gravity  | R             |  |                    |
| 2.14.7    | Water Solubility (mg/L)  | R             |  |                    |
| 2.14.8    | Solvent Solubility (mg/L)  | R             |  |                    |
| 2.14.9    | Vapor Pressure   | R             |  |                    |
| 2.14.10   | Dissociation Constant  | R             |  |                    |

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|-----------|--|---------------|--|--------------------|
| 2.14.11   | Octanol/Water Partition Coefficient                                | R             |  |                    |
| 2.14.12   | рН   | R             |  |                    |
| 2.14.13   | UV/Visible Absorption Spectra                                      | R             |  |                    |
| 2.14.14   | Stability (Sunlight, Temperature, Metals)                          | R             |  |                    |
| 2.14.15   | Storage Stability Data   | R             | Ambient 1 year may be submitted                                  |                    |
|           |  |               | later but short term data is                                     |                    |
| 2.15      | Other Studies/Data/Reports/Foreign Reviews                         |               |  |                    |
| 3         | Chemistry Requirements for the Registration of                     |               |  |                    |
|           | Manufacturing Concentrates and End-Use                             |               |  |                    |
|           | Products Formulated from Registered technical                      |               |  |                    |
|           | grade of active ingredients or integrated system                   |               |  |                    |
|           | products.  |               |  |                    |
| 3.1       | Product Identification   |               |  |                    |
| 3.1.1     | Applicant's Name and Office Address                                |               |  |                    |
| 3.1.2     | Formulating Plant's Name and Address                               |               |  |                    |
| 3.1.3     | Trade Name   |               |  |                    |
| 3.1.4     | Other Names  |               |  |                    |
| 3.2       | Formulation Process  |               |  |                    |
| 3.2.1     | Description of Starting Materials                                  | R             |  |                    |
| 3.2.2     | Description of the Formulation Process                             | R             |  |                    |
| 3.2.3     | Discussion of the Formation of Impurities of Toxicological Concern | CR            | If aplicable.  |                    |
| 3.3       | Specifications   |               |  |                    |
| 3.3.1     | Establishing Certified Limits                                      | CR            | A justification must be provided if standard limits are not met. |                    |
| 3.3.2     | Additional Product Specific Requirements                           |               |  |                    |
| 3.3.2.1   | Granules and Baits   | CR            | If applicable.   |                    |
| 3.3.2.2   | Seed Coatings  | CR            | If applicable.   |                    |
| 3.3.3     | Control Product Specification Form                                 | R             |  |                    |
| 3.4       | Product Analysis   |               |  |                    |
| 3.4.1     | Enforcement Analytical Method                                      | R             |  |                    |
| 3.4.2     | Impurities of Toxicological Concern                                | CR            | If applicable.   |                    |
| 3.5       | Chemical and Physical Properties                                   |               |  |                    |
| 3.5.1     | Colour   | R             |  |                    |
| 3.5.2     | Physical State   | R             |  |                    |
| 3.5.3     | Odour  | R             |  |                    |
| 3.5.4     | Formulation Type   | R             |  |                    |
| 3.5.5     | Container Material and Description                                 | R             |  |                    |
| 3.5.6     | Density or Specific Gravity  | R             |  |                    |
| 3.5.7     | рН   | R             |  |                    |
| 3.5.8     | Oxidizing or Reducing Action (Chemical Incompatibility)            | R             |  |                    |

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| Data Code | Title  | Data     | Conditions                         | Present/ |
|-----------|--|----------|------------------------------------|----------|
|           |  | required |                                    | Absent   |
| 3.5.9     | Viscosity                                    | R        |                                    |          |
| 3.5.10    | Storage Stability Data                       | R        | Ambient 1 year may be submitted    |          |
|           |  |          | later but short term data is       |          |
| 3.5.11    | Flammability                                 | R        |                                    |          |
| 3.5.12    | Explodability                                | R        |                                    |          |
| 3.5.13    | Miscibility                                  | R        |                                    |          |
| 3.5.14    | Corrosion Characteristics                    | R        |                                    |          |
| 3.5.15    | Dielectric Breakdown Voltage                 | R        |                                    |          |
| 3.6       | Other Studies/Data/Reports/Foreign Reviews   |          |                                    |          |
| 4         | Toxicology                                   |          |                                    |          |
| 4.1       | Summaries                                    | R        |                                    |          |
| 4.2       | Acute Studies — TGAI                         |          |                                    |          |
| 4.2.1     | Acute Oral                                   | R        |                                    |          |
| 4.2.2     | Acute Dermal                                 | NR       |                                    |          |
| 4.2.3     | Acute Inhalation                             | NR       |                                    |          |
| 4.2.4     | Primary Eye Irritation                       | NR       |                                    |          |
| 4.2.5     | Primary Dermal Irritation                    | NR       |                                    |          |
| 4.2.6     | Dermal Sensitization                         | NR       |                                    |          |
| 4.2.7     | Potentiation/Interaction                     | CR       | If available                       |          |
| 4.2.8     | Antidote                                     | CR       | If available                       |          |
| 4.2.9     | Other Acute Studies                          | CR       | If available                       |          |
| 4.3       | Short-term Studies — TGAI                    |          |                                    |          |
| 4.3.1     | Short-term Oral (90-day rodent)              | R        |                                    |          |
| 4.3.2     | Short-term Oral (90-day and/or 12-month dog) | R        |                                    |          |
| 4.3.3     | Short-term Oral (28-day)                     | CR       | If available                       |          |
| 4.3.4     | Short-term Dermal (90-day)                   | NR       |                                    |          |
| 4.3.5     | Short-term Dermal (21/28-day)                | NR       |                                    |          |
| 4.3.6     | Short-term Inhalation (90-day)               | NR       |                                    |          |
| 4.3.7     | Short-term Inhalation (21/28-day)            | NR       |                                    |          |
| 4.3.8     | Other Short-term Studies                     | CR       | If available                       |          |
| 4.4       | Long-term Studies — TGAI                     |          |                                    |          |
| 4.4.1     | Chronic (rodent)                             | R        | 4.4.1 and 4.4.2 could be submitted |          |
|           |  |          | as a combined study under 4.4.4    |          |
| 4.4.2     | Oncogenicity (rodent species 1)              | R        | 4.4.1 and 4.4.2 could be submitted |          |
|           |  |          | as a combined study under 4.4.4    |          |
| 4.4.3     | Oncogenicity (rodent species 2)              | R        | ·                                  |          |
| 4.4.4     | Combined Chronic/Oncogenicity (rodent)       | CR       | 4.4.1 and 4.4.2 could be submitted |          |
|           |  |          | as a combined study under 4.4.4    |          |
| 4.4.5     | Other Long-term Studies                      | CR       | If available                       |          |
| 4.5       | Special Studies — TGAI                       |          |                                    |          |
| 4.5.1     | Multigeneration Reproduction (rodent)        | R        |                                    |          |
| 4.5.2     | Prenatal Developmental Toxicity (rodent)     | R        |                                    |          |

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| Data Code | Title   | Data     | Conditions   | Present/ |
|-----------|---|----------|--|----------|
| 1.7.0     |   | required |  | Absent   |
| 4.5.3     | Prenatal Developmental Toxicity (non-rodent)              | R        |  |          |
| 4.5.4     | Genotoxicity: Bacterial Reverse Mutation Assay            | R        |  |          |
| 4.5.5     | Genotoxicity: In vitro Mammalian Cell Assay               | R        |  |          |
| 4.5.6     | Genotoxicity: In vitro Mammalian Clastogenicity           | CR       | Required if not addressed in study submitted for 4.5.5   |          |
| 4.5.7     | Genotoxicity: In vivo Cytogenetics                        | R        |  |          |
| 4.5.8     | Other Genotoxicity Studies                                | CR       | If available   |          |
| 4.5.9     | Metabolism/Toxicokinetics in Mammals (laboratory animals) | R        |  |          |
| 4.5.10    | Acute Delayed Neurotoxicity (hen)                         | CR       | Required if the test substance is an organophosphorus substance or is structurally related to other substances that may cause delayed neurotoxicity                  |          |
| 4.5.11    | 28-day Delayed Neurotoxicity (hen)                        | CR       | Required if results of acute delayed neurotoxicity study indicates effects, or if other available data indicate the potential for this type of delayed neurotoxicity |          |
| 4.5.12    | Acute Neurotoxicity (rat)                                 | CR       | Required if there is neurotoxic potential  |          |
| 4.5.13    | 90-day Neurotoxicity (rat)                                | CR       | Required if there is neurotoxic potential  |          |
| 4.5.14    | Developmental Neurotoxicity                               | CR       | Required if neurological effects are<br>observed in other studies<br>Should be considered if test  |          |
|           |   |          | substance: i) causes neuropathology or neurotoxicity in adults;  |          |
|           |   |          | ii) is hormonally active <i>in vivo</i> ; or iii) causes other types of nervous system involvement at a developmental stage  |          |
| 4.6       | Acute Studies — EP  | <u> </u> | 1  |          |
| 4.6.1     | Acute Oral  | CR       | If available   |          |
| 4.6.2     | Acute Dermal  | NR       |  |          |
| 4.6.3     | Acute Inhalation  | NR       |  |          |
| 4.6.4     | Primary Eye Irritation                                    | NR       |  |          |
| 4.6.5     | Primary Dermal Irritation                                 | NR       |  |          |
| 4.6.6     | Dermal Sensitization                                      | NR       |  |          |
| 4.6.7     | Potentiation/Interaction                                  | CR       | If available   |          |
| 4.6.8     | Other Acute Studies                                       | CR       | If available   |          |

# DATA REQUIREMENTS FOR IMPORT MRLs ON USE SITE CATEGORY

(USC # 8): Livestock for Food

| Data Code | Title   | Data     | Conditions                                  | Present/ |
|-----------|---|----------|---|----------|
|           |   | required |   | Absent   |
| 4.7       | Short-term Studies — EP                         | CR       | Depending on use pattern, required          |          |
|           |   |          | if any component of the EP may              |          |
|           |   |          | increase absorption of the active           |          |
|           |   |          | ingredient(s) or increase toxic or          |          |
| 4.7.1     | Shout town Oral (00 day as dant)                | CR       | pharmacologic effects<br>See 4.7            |          |
|           | Short-term Oral (90-day rodent)                 |          |   |          |
| 4.7.2     | Short-term Oral (90-day and/or 12-month dog)    | CR       | See 4.7                                     |          |
| 4.7.3     | Short-term Dermal (90-day)                      | NR       |   |          |
| 4.7.4     | Short-term Dermal (21/28-day)                   | NR       |   |          |
| 4.7.5     | Short-term Inhalation (21/28-day)               | NR       |   |          |
| 4.7.6     | Short-term Inhalation (90-day)                  | NR       |   |          |
| 4.7.7     | Other Special Studies                           | CR       | See 4.7                                     |          |
| 4.8       | Other Studies/Data/Reports                      | CR       | If available                                |          |
| 6         | Metabolism/Toxicokinetics Studies (TGAI or EP)  |          |   |          |
| 6.1       | Summaries                                       | R        | Input to assessment of toxicity to wildlife |          |
| 6.2       | Livestock                                       | R        |   |          |
| 6.4       | Other Studies/Data/Reports/Foreign Reviews      | CR       |   |          |
| 7         | Food, Feed and Tobacco Residue Studies EP       |          |   |          |
| 7.1       | Summaries                                       | R        | Input to assessment of wildlife             |          |
|           |   |          | exposure                                    |          |
| 7.2       | Analytical Methodology (Food Crops & Tobacco)   |          |   |          |
| 7.2.1     | Supervised Residue Trial Analytical Methodology | R        |   |          |
| 7.3       | Freezer Storage Stability Tests                 | CR       | Depends on length of storage                |          |
| 7.4       | Crop Residue Data                               |          |   |          |
| 7.4.5     | Processed Food/Feed                             | CR       |   |          |
| 7.6       | Livestock, Poultry, Egg and Milk Residue Data   | R        | Some use patterns are not directly          |          |
|           | (external application)                          |          | dermal                                      |          |
| 7.8       | Other Studies/Data/Reports/Foreign Reviews      | CR       |   |          |

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