Risk Assessment and Risk Management Plan (consultation version) for

**DIR 188**

Limited and controlled release of canola and Indian mustardgenetically modified for altered oil content and herbicide tolerance

Applicant: Nuseed Pty Ltd

**This RARMP is open for consultation until** **26 April 2022**.

Written comments on the risks to human health and safety and the environment posed by this proposed release are invited. You may make your submission

via mail to: The Office of the Gene Technology Regulator

MDP 54, GPO Box 9848, Canberra ACT 2601 or

via email to: [ogtr@health.gov.au](mailto:ogtr@health.gov.au).

Please note that issues regarding food safety and labelling, the use of agricultural chemicals, and marketing and trade implications do **not** fall within the scope of these evaluations as they are the responsibilities of other agencies and authorities.

# Summary of the Risk Assessment and Risk Management Plan

**(consultation version) for**

**Licence Application No. DIR 188**

***Introduction***

The Gene Technology Regulator (the Regulator) has received a licence application for the intentional release of a genetically modified organism (GMO) into the environment. It qualifies as a limited and controlled release application under the *Gene Technology Act 2000* (the Act). The Regulator has prepared a Risk Assessment and Risk Management Plan (RARMP) for this application, which concludes that the proposed field trial poses negligible risks to human health and safety and the environment. Licence conditions have been drafted for the proposed field trial. The Regulator invites submissions on the RARMP, including draft licence conditions, to inform the decision on whether or not to issue a licence.

***The application***

|  |  |
| --- | --- |
| Applicant | Nuseed Pty Ltd |
| Project title | Limited and controlled release of canola and Indian mustard genetically modified for altered oil content and herbicide tolerance |
| Parent organisms | Canola (*Brassica napus* L.)  Indian mustard (*Brassica juncea* (L.) Czern. & Coss.) |
| Introduced genes | Seven genes involved in biosynthesis pathway for long-chain polyunsaturated fatty acids:   * *Lackl-Δ12D* from yeast *Lachancea kluyveri* * *Picpa-* *ω3D* from yeast *Pichia pastoris* * *Micpu-Δ6D* from microalga *Micromonas pusilla* * *Pyrco-Δ6E* from microalga *Pyramimonas cordata* * *Pavsa-Δ5D* from microalga *Pavlova salina* * *Pyrco-Δ5E* from microalga *Pyramimonas cordata* * *Pavsa-Δ4D* from microalga *Pavlova salina*   One selectable marker gene that confers herbicide tolerance:   * *pat* genefrom soil bacterium *Streptomyces viridochromogenes* forglufosinate tolerance |
| Proposed locations | Up to 20 trial sites per year to be selected from 96 possible local government areas in New South Wales, Victoria and Queensland |
| Proposed release size | Up to 150 ha per year |
| Proposed period of release | From November 2022 to December 2027 |
| Principal purpose | To evaluate the altered oil content trait under field conditions |

***Risk assessment***

The risk assessment concludes that risks to the health and safety of people, or the environment, from the proposed release are negligible. No specific risk treatment measures are required to manage these negligible risks.

The risk assessment process considers how the genetic modification and proposed activities conducted with the GMOs might lead to harm to people or the environment. Risks are characterised in relation to both the seriousness and likelihood of harm, taking into account current scientific/technical knowledge, information in the application (including proposed limits and controls) and relevant previous approvals. Both the short- and long-term impacts are considered.

Credible pathways to potential harm that were considered included exposure of people or desirable animals to the GM plant material, potential for persistence or dispersal of the GMOs, and transfer of the introduced genetic material to canola, Indian mustard and related plants outside the field trial. Potential harms associated with these pathways included toxicity or allergenicity to people, toxicity to desirable animals, and environmental harms due to weediness.

The principal reasons for the conclusion of negligible risks are that the proposed limits and controls will effectively minimise exposure to the GMOs, and there is no evidence to suggest the introduced genetic modifications would lead to harm to people or the environment.

***Risk management***

The risk management plan describes measures to protect the health and safety of people and to protect the environment by controlling or mitigating risk. The risk management plan is given effect through licence conditions. Draft licence conditions are detailed in Chapter 4 of the RARMP.

As the level of risk is considered negligible, specific risk treatment is not required. However, since this is a limited and controlled release, the draft licence includes limits on the size, location and duration of the release, as well as controls to prohibit the use of GM plant material in commercial human food or animal feed, to minimise dispersal of the GMOs or GM pollen from the trial sites, to transport the GMOs in accordance with the Regulator’s guidelines, to destroy GMOs at the end of the trial and to conduct post-harvest monitoring at the trial sites to ensure the GMOs are destroyed.

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# Abbreviations

|  |  |
| --- | --- |
| ALA | α-linolenic acid |
| CCI | Confidential Commercial Information |
| DIR | Dealings involving Intentional Release |
| DHA | Docosahexaenoic acid |
| DPA | Docosapentaenoic acid |
| ETA | Eicosatetraenoic acid |
| FSANZ | Food Standards Australia New Zealand |
| GM(O) | Genetically modified (organism) |
| ha | Hectare(s) |
| HGT | Horizontal gene transfer |
| km | Kilometre(s) |
| LC-PUFA | Long chain polyunsaturated fatty acid |
| m | Metre(s) |
| mm | Millimetre(s) |
| NSW | New South Wales |
| OGTR | Office of the Gene Technology Regulator |
| PAT | Phosphinothricin N-acetyltransferase |
| PUFA | Polyunsaturated fatty acid |
| RARMP | Risk Assessment and Risk Management Plan |
| Regulations | Gene Technology Regulations 2001 |
| Regulator | Gene Technology Regulator |
| the Act | The *Gene Technology Act 2000* |
| ω | Omega |

1. Risk assessment context
   1. Background
2. An application has been made under the *Gene Technology Act 2000* (the Act) for Dealings involving the Intentional Release (DIR) of genetically modified organisms (GMOs) into the Australian environment.
3. The Act and the Gene Technology Regulations 2001 (the Regulations), together with corresponding State and Territory legislation, comprise Australia’s national regulatory system for gene technology. Its objective is to protect the health and safety of people, and to protect the environment, by identifying risks posed by or as a result of gene technology, and by managing those risks through regulating certain dealings with GMOs.
4. Section 50 of the Act requires that the Gene Technology Regulator (the Regulator) must prepare a Risk Assessment and Risk Management Plan (RARMP) in response to an application for release of GMOs into the Australian environment. Sections 50, 50A and 51 of the Act and sections 9 and 10 of the Regulations outline the matters which the Regulator must take into account and who must be consulted when preparing the RARMP.
5. The *Risk Analysis Framework* (OGTR, 2013a) explains the Regulator’s approach to the preparation of RARMPs in accordance with the Act and the Regulations. The Regulator has also developed operational policies and guidelines that are relevant to DIR licences. These documents are available from the Office of the Gene Technology Regulator (OGTR) [website](http://www.ogtr.gov.au/).
6. Figure 1 shows the information that is considered, within the regulatory framework, in establishing the risk assessment context. This information is specific for each application. Risks to the health and safety of people or the environment posed by the proposed release are assessed within this context. Chapter 1 provides the specific information for establishing the risk assessment context for this application.



**Figure 1.** Summary of parameters used to establish the risk assessment context

1. In accordance with section 50A of the Act, this application is considered to be a limited and controlled release application, as the Regulator was satisfied that it meets the criteria prescribed by the Act. Therefore, the Regulator was not required to consult with prescribed experts, agencies and authorities before preparation of the RARMP.
2. The GMOs and any proposed dealings may also be subject to regulation by other Australian government agencies that regulate GMOs or GM products, including Food Standards Australia New Zealand (FSANZ), the Australian Pesticides and Veterinary Medicines Authority, the Therapeutic Goods Administration and the Department of Agriculture and Water Resources. Proposed dealings may also be subject to the operation of State legislation declaring areas to be GM, GM free, or both, for marketing purposes.
   1. The proposed dealings
3. Nuseed Pty Ltd (Nuseed) proposes to release up to 80 GM canola and Indian mustard lines into the environment under limited and controlled conditions. The GM plants have been genetically modified for altered seed oil content and herbicide tolerance.
4. The purpose of the release is to evaluate the altered oil content trait under field conditions. The field trial will gather research and regulatory data about agronomic performance, oil content and profile, nutritional assessment, compositional analysis, molecular analysis and genetic stability.
5. The dealings involved in the proposed intentional release are:

* conducting experiments with the GMOs
* breeding the GMOs
* propagating the GMOs
* using the GMOs in the course of manufacture of a thing that is not a GMO
* growing the GMOs
* importing the GMOs
* transporting the GMOs
* disposing of the GMOs

and the possession, supply or use of the GMOs in the course of any of these dealings.

1. GM plant material would not be used in commercial human food or animal feed.
2. GM plant material may be exported and used in a human nutritional study outside Australia. Proposed dealings with GM plant material in countries other than Australia do not fall within the jurisdiction of Australia’s Gene Technology Regulator and will not be considered in this RARMP.
3. GM plant material may be used in human sensory trials to test the feel, smell, taste and appearance of the seed oil or food products containing the oil. Taste testing trials would not result in consumption of the oil. These trials would only occur if Nuseed has the appropriate approvals for each trial in accordance with the National Statement on Ethical Conduct in Human Research.
4. GM plant material may be used in animal feeding studies. These could include toxicology trials with rodents, bioavailability trials with rodents, broiler chicken feeding trials and aquaculture feeding trials. These trials would only occur if Nuseed has the appropriate approvals for each trial in accordance with the Australian Code of Practice for the Care and Use of Animals for Scientific Purposes.

2.1 The proposed limits of the dealings (duration, size, location and people)

1. The field trial is proposed to take place over five years, between November 2022 and December 2027. In each year there would be up to ten trial sites of 10 ha and ten trial sites of 5 ha, with a total trial area of up to 150 ha per year.
2. The trial sites would be selected from 96 local government areas in New South Wales (NSW), Victoria and Queensland (Table 1). The trial sites would be located on private land in rural areas. Details of site locations would be provided to the Regulator prior to each planting season.

**Table 1.** Local government areas where proposed trial sites may be located

|  |  |  |
| --- | --- | --- |
| New South Wales | Victoria | Queensland |
| Albury City Council | Ararat Rural City Council | Goondiwindi Regional Council |
| Balranald Shire Council | Ballarat City Council | Lockyer Valley Regional Council |
| Berrigan Shire Council | Benalla Rural City Council | Somerset Regional Council |
| Bland Shire Council | Buloke Shire Council | Southern Downs Regional Council |
| Blayney Shire Council | Campaspe Shire Council | Toowoomba Regional Council |
| Cabonne Shire Council | Central Goldfields Shire Council | Western Downs Regional Council |
| Carrathool Shire Council | Colac-Otway Shire Council |  |
| Coolamon Shire Council | Corangamite Shire Council |  |
| Coonamble Shire Council | Gannawarra Shire Council |  |
| Cootamundra- Gundagai Regional Council | Glenelg Shire Council |  |
| Cowra Shire Council | Golden Plains Shire Council |  |
| Dubbo Regional Council | Greater Bendigo City Council |  |
| Edward River Council | Greater Geelong City Council |  |
| Federation Council | Greater Shepparton City Council |  |
| Forbes Shire Council | Hepburn Shire Council |  |
| Gilgandra Shire Council | Hindmarsh Shire Council |  |
| Greater Hume Shire Council | Horsham Rural City Council |  |
| Griffith City Council | Indigo Shire Council |  |
| Gunnedah Shire Council | Latrobe City Council |  |
| Gwydir Shire Council | Loddon Shire Council |  |
| Hay Shire Council | Macedon Ranges Shire Council |  |
| Hilltops Council | Melton Shire Council |  |
| Junee Shire Council | Mildura Rural City Council |  |
| Lachlan Shire Council | Mitchell Shire Council |  |
| Leeton Shire Council | Moira Shire Council |  |
| Liverpool Plains Shire Council | Moorabool Shire Council |  |
| Lockhart Shire Council | Mount Alexander Shire Council |  |
| Mid-Western Regional Council | Moyne Shire Council |  |
| Moree Plains Shire Council | Murrindindi Shire Council |  |
| Murray River Council | Northern Grampians Shire Council |  |
| Murrumbidgee Council | Pyrenees Shire Council |  |
| Muswellbrook Shire Council | South Gippsland Shire Council |  |
| Narrabri Shire Council | Southern Grampians Shire Council |  |
| Narrandera Shire Council | Strathbogie Shire Council |  |
| Narromine Shire Council | Surf Coast Shire Council |  |
| Orange City Council | Swan Hill Rural City Council |  |
| Parkes Shire Council | Towong Shire Council |  |
| Snowy Valleys Council | Wangaratta Rural City Council |  |
| Tamworth Regional Council | Warrnambool City Council |  |
| Temora Shire Council | Wellington Shire Council |  |
| Upper Hunter Shire Council | West Wimmera Shire Council |  |
| Wagga Wagga City Council | Wodonga City Council |  |
| Walgett Shire Council | Wyndham City Council |  |
| Warren Shire Council | Yarriambiack Shire Council |  |
| Warrumbungle Shire Council |  |  |
| Weddin Shire Council |  |  |

1. Only trained and authorised staff would be permitted to deal with the GM plants.

2.2 The proposed controls to restrict the spread and persistence of the GMOs in the environment

1. The applicant has proposed a number of controls to restrict the spread and persistence of the GM canola and Indian mustard and the introduced genetic material in the environment. These include:

* locating each proposed trial site at least 50 m away from the nearest natural waterway
* restricting gene flow from the GMOs using one of the combinations of controls shown in Figure 2
* treating any non-GM canola or Indian mustard plants grown in planting areas or pollen traps like the GMOs
* harvesting the GMOs separately from other crops
* after harvest, destroying GMOs not required for further evaluation or future trials
* cleaning equipment used in connection with the GMOs as soon as practicable and before use for any other purpose
* transporting and storing GMOs in accordance with the current Regulator’s [Guidelines for the Transport, Storage and Disposal of GMOs](https://www.ogtr.gov.au/resources/publications/guidelines-transport-storage-and-disposal-gmos)
* post-harvest tilling of planting areas, pollen traps and other areas where GMOs were dispersed to encourage seed germination
* post-harvest monitoring of each trial site monthly for at least 2 years and until the site is free of volunteer canola or Indian mustard plants for at least 12 months, with any volunteer plants destroyed prior to flowering
* during the post-harvest monitoring period, planting only crops permitted on GM brassica trial sites by the Regulator’s [Policy on Post-Harvest Crops](https://www.ogtr.gov.au/resources/publications/policy-post-harvest-crops).

Figure showing three options for trial layout, including some of the controls to restrict gene flow.


**Figure 2.** Options for restricting gene flow from the GM canola and Indian mustard (not to scale).Site layout (a) with Insect-proof tent covering the GMOs during flowering, (b) without Insect‑proof tent and with Pollen Trap surrounding the Planting Area, and (c) without Insect-proof tent or Pollen Trap. Monitoring and Isolation Zones must be kept free of related plants.

1. The proposed limits and controls are taken into account in the risk assessment (Chapter 2) and their suitability for containing the release will be evaluated in the risk management plan (Chapter 3).
   1. The parent organism
2. The parent organisms are *Brassica napus* L., which is commonly known as canola, rapeseed or oilseed rape, and *Brassica juncea* (L.) Czern. & Coss., which is commonly known as Indian mustard or juncea canola. *B. napus* and *B. juncea* are both exotic to Australia.
3. Canola is the third-most widely grown crop in Australia. It is grown mainly in Western Australia, NSW, Victoria and South Australia (ABARES, 2021). Canola oil is used as food and the canola meal remaining after oil extraction is used as animal feed. Almost all commercial canola grown in Australia is *B. napus*, but a small amount is canola-quality *B. juncea*, which is adapted to low-rainfall areas. Other varieties of *B. juncea* are grown in Australia to produce condiment mustard (GRDC, 2017). The GM Indian mustard proposed for release is derived from canola-quality *B. juncea* varieties.
4. Both *B. napus* and *B. juncea* are naturalised in Australia. In areas where they are grown, they can be agricultural weeds in subsequent crops. There are isolated reports of *B. napus* as an environmental weed in Western Australia and *B. napus* and *B. juncea* as environmental weeds in Victoria (Randall, 2017). However, the most recent Western Australian state government environmental weed risk assessment gives *B. napus* a weed risk rating of negligible ([Environmental weed risk assessments](https://www.agric.wa.gov.au/rangelands/environmental-weed-risk-assessments), accessed 10 Jan 2022), and the most recent Victorian state government environmental weed list gives both *B. napus* and *B. juncea* risk ranking scores of zero (White et al., 2018).
5. Detailed information about the parent organisms is contained in the document *The Biology of Brassica napus L. (canola) and Brassica juncea (L.) Czern. & Coss. (Indian mustard)* (OGTR, 2017), which was produced to inform the risk analysis process and is available from the [Resources page](https://www.ogtr.gov.au/resources) on the OGTR website. Baseline information from this document will be used and referred to throughout the RARMP.
6. Some information about the specific parent organisms used for this application has been declared Confidential Commercial Information (CCI). Under section 185 of the Act, the confidential information is made available to the prescribed experts and agencies that are consulted on the RARMP for this application.
   1. The GMOs, nature and effect of the genetic modification
7. The applicant proposes to grow up to 80 lines of GM canola and Indian mustard with altered oil content and herbicide tolerance. Some information about the categories of GMOs proposed for release has been declared Confidential Commercial Information (CCI). Under section 185 of the Act, the confidential information is made available to the prescribed experts and agencies that are consulted on the RARMP for this application.

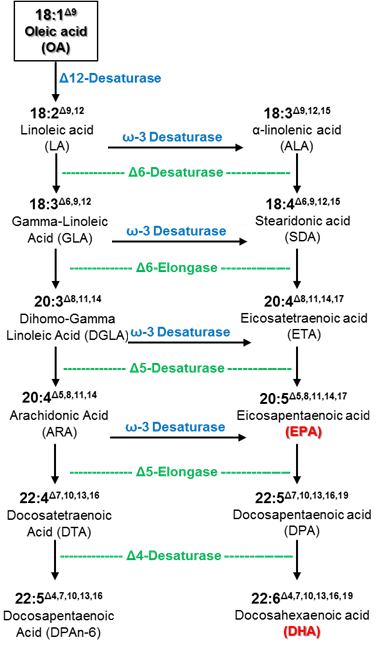
4.1 The genetic modifications in the GMOs proposed for release

1. The GMOs contain up to seven introduced genes involved in fatty acid biosynthesis and one introduced selectable marker gene that confers herbicide tolerance.
2. The seven introduced fatty acid biosynthesis genes (Table 2) were sourced from yeast and marine microalgae, and codon optimised for expression in higher plants.

**Table 2.** Introduced genes involved in fatty acid biosynthesis

| **Gene** | **Source organism** | **Encoded protein** | **Reference** |
| --- | --- | --- | --- |
| *Lackl-*Δ*12D* | *Lachancea kluyveri* yeast | Δ12-desaturase | (Petrie et al., 2012) |
| *Picpa-ω3D* | *Pichia pastoris* yeast | ω-3 desaturase | (Zhang et al., 2008) |
| *Micpu-*Δ*6D* | *Micromonas pusilla* microalgae | Δ6-desaturase | (Petrie et al., 2010b) |
| *Pyrco-*Δ*6E* | *Pyramimonas cordata* microalgae | Δ6-elongase | (Petrie et al., 2010a) |
| *Pavsa-*Δ*5D* | *Pavlova salina* microalgae | Δ5-desaturase | (Zhou et al., 2007) |
| *Pyrco-*Δ*5E* | *Pyramimonas cordata* microalgae | Δ5-elongase | (Petrie et al., 2010a) |
| *Pavsa-*Δ*4D* | *Pavlova salina* microalgae | Δ4-desaturase | (Zhou et al., 2007) |

1. The purpose of the introduced fatty acid biosynthesis genes is to convert oleic acid, which is an abundant fatty acid in canola and Indian mustard seed oil, into ω-3 long-chain polyunsaturated fatty acids (LC-PUFAs), which are not naturally present in plant seed oil (Ruiz-Lopez et al., 2015; Saini et al., 2021). ω-3 LC-PUFAs are fatty acids of 20 or more carbons in length with multiple cis double bonds in their backbone, with the first double bond on the third carbon from the methyl end. The fatty acid biosynthesis pathways are shown in Figure 3.



**Figure 3.** Outline of pathways for biosynthesis of ω-3 LC-PUFAs in GM plants, adapted from Ruiz-Lopez et al. (2013). The main ω‑3 LC-PUFAs with importance for human health are highlighted in red. The *Picpa-ω3D* ω-3 desaturase introduced into the GMOs has similar conversion rates for all 18-carbon and 20‑carbon substrates (Zhang et al., 2008). However, the right-most biosynthesis pathway in the diagram is likely to be preferred in the GMOs, as Δ6-desaturation is reported to be the rate-limiting step (Petrie et al., 2020) and the *Micpu-Δ6D* Δ6-desaturase has 3.5‑fold greater conversion efficiency with an α-linolenic acid (ALA) substrate than with a linoleic acid (LA) substrate (Petrie et al., 2010b).

1. The GMOs may also contain the *pat* selectable marker gene, which was used during initial development of the GM plants in the laboratory to select plant cells containing the introduced genes. The *pat* gene is sourced from the soil bacterium *Streptomyces viridochromogenes.* It encodes the phosphinothricin N‑acetyltransferase (PAT) enzyme, which confers tolerance to glufosinate (phosphinothricin) herbicide.
2. Short regulatory sequences that control gene expression have also been introduced into the GMOs (Table 3). The expression of the introduced fatty acid biosynthesis genes is targeted to the seed with seed-specific promoters, while the expression of the selectable marker gene *pat* is driven by a constitutive promoter, which is active in all plant tissues. Other short regulatory elements used include enhancers of gene expression and terminators.

**Table 3.** Introduced regulatory sequences

| **Sequence** | **Source** | **Intended function** |
| --- | --- | --- |
| PRO\_Arath-FAE1 | Promoter of *Arabidopsis thaliana* fatty acid elongase 1 | Seed specific promoter |
| PRO\_Brana-FP1 | Promoter of *Brassica napus* napin | Seed specific promoter |
| PRO\_Linus-Cnl1 | Promoter of *Linum usitatissimum* conlinin1 | Seed specific promoter |
| PRO\_Linus-Cnl2 | Promoter of *Linum usitatissimum* conlinin2 | Seed specific promoter |
| PRO\_35S×2 | Promoter of Cauliflower mosaic virus 35S RNA | Constitutive promoter |
| Tobacco mosaic virus 5' UTR leader | Enhancer from Tobacco mosaic virus 59 | Increase gene expression |
| MAR\_Nicta- RB7 | Rb7 matrix attachment region from *Nicotiana tabacum* | Increase gene expression |
| TER\_Agrtu-NOS | Terminator of *Agrobacterium tumefaciens* nopaline synthase | Terminator |
| TER\_Glyma-Lectin | Terminator of *Glycine max* lectin *Le1* | Terminator |
| TER\_Linus-Cnl1 | Terminator of *Linum usitatissimum* conlinin1 | Terminator |
| TER\_Linus-Cnl2 | Terminator of *Linum usitatissimum* conlinin2 | Terminator |

1. Gene constructs were introduced into the GMOs using *Agrobacterium*–mediated transformation. This method has been widely used in Australia and overseas for introducing genetic modifications into plants. More information can be found in the document *Methods of Plant Genetic Modification* which is available from the [Resources page](https://www.ogtr.gov.au/resources) on the OGTR website.

4.2 Toxicity/allergenicity of the proteins associated with the introduced genes

1. None of the source organisms of the introduced genes are known to be toxic, allergenic or pathogenic.
2. Bioinformatic analysis may assist in the assessment process by predicting, on a theoretical basis, the toxic or allergenic potential of a protein. The sequences of the eight introduced proteins were compared to all proteins in the NCBI Entrez Protein database as well as the AllergenOnline.org database (version 18). The searches did not find any biologically relevant similarity between the introduced proteins and any known toxin or allergen (MacIntosh et al., 2021).
3. All introduced proteins were readily digested by pepsin in a standard assay of the digestibility of proteins in simulated gastric fluid (MacIntosh et al., 2021).
4. FSANZ has assessed the safety of a GM canola line containing all of the introduced genes included in this application. FSANZ concluded that food derived from the GM canola line is considered to be as safe for human consumption as food derived from conventional (non-GM) canola cultivars (FSANZ, 2017).

4.3 Toxicity due to the altered oil content trait

1. The GMOs are intended to produce ω-3 LC-PUFAs in seed oil. The applicant states that the GMOs will contain some or all of the fatty acid biosynthesis genes listed in Table 2. Depending on which introduced genes are present in each GM transformant, the main ω-3 LC-PUFA produced could be eicosatetraenoic acid (ETA), eicosapentaenoic acid (EPA), docosapentaenoic acid (DPA) or docosahexaenoic acid (DHA), as shown in Figure 3.
2. As enzymatic conversion efficiency is not 100%, the seed oil of the GMOs is expected to contain some fatty acid intermediates as well as the target ω-3 LC-PUFAs. For example, in a GM canola line containing all of the fatty acid biosynthesis genes listed in Table 2 and designed to produce a high level of DHA (Petrie et al., 2020), the fatty acid profile included 9.7% DHA, 1.0% DPA, 1.3% ETA and 2.2% stearidonic acid (SDA), while none of these fatty acids were present in the non-GM parent canola. The GM canola line was also enriched in ALA, with 20% ALA in the GM line compared to 9.5% ALA in the non-GM parent.
3. ω-3 LC-PUFAs are naturally present in seafood and are particularly abundant in oily fish. Many plant food products are rich in ALA, and human biosynthetic pathways can convert ALA into ω-3 LC‑PUFAs, although at low conversion efficiencies. As ω-3 LC-PUFAs are considered to be beneficial to human health, fish oil is commonly consumed as a dietary supplement (Zarate et al., 2017; Shahidi and Ambigaipalan, 2018; Saini et al., 2021). Therefore, the target ω-3 LC-PUFAs and fatty acid intermediates produced by the GMOs are normally present in the human diet and/or synthesized in humans, and are not expected to be toxic.
4. ω-3 LC-PUFAs are highly susceptible to oxidation, with their oxidative stability inversely related to the number of carbon double bonds. Their primary oxidation products are lipid peroxides and lipid free radicals, which further decompose into a mix of secondary oxidation products including aldehydes and ketones. The rate of oxidation depends on temperature, exposure to light and exposure to oxygen (Arab-Tehrany et al., 2012; Albert et al., 2013; Miyashita, 2019). Even very low levels of oxidation produce volatile secondary oxidation products with offensive (rancid) odours and tastes. Flavour deterioration due to oxidation is a common problem in commercial fish oil products, despite use of antioxidant additives (Arab-Tehrany et al., 2012; Miyashita, 2019). In animal studies, highly oxidised PUFAs are reported to have toxic effects (Albert et al., 2013; Albert et al., 2016). For instance, pregnant rats fed fish oil where approximately 10% of the ω-3 LC‑PUFAs were oxidized had much greater newborn mortality than control rats fed unoxidized fish oil or water (Albert et al., 2016).
5. A study of a GM canola line containing all of the fatty acid biosynthesis genes listed in Table 2 (Petrie et al., 2020) tested the stability of the seed oil profile when the seeds were stored at 24°C or 32°C for six months after harvest. There was no measured difference between the DHA levels of freshly harvested seeds and these stored seeds, although it is noted that the error bars of the measurements were up to ±4%, so it is possible that a small proportion of the DHA was oxidized during storage.

4.4 Characterisation of the GMOs

1. The introduced genes are not known to confer any phenotypic changes other than altered seed oil profile and herbicide tolerance. The applicant states that no unexpected phenotype has been observed while growing the GMOs in glasshouses.
2. A study of a GM canola line containing all of the introduced genes included in the current application (Petrie et al., 2020) evaluated agronomic parameters in the field. The GM canola line had a small reduction in total seed oil content compared to the non-GM parent canola cultivar, but there were no significant changes to yield, crop emergence, time to flowering and maturity, pod shattering, disease incidence or pest predation.
3. Gene constructs were introduced into the GMOs using *Agrobacterium*–mediated transformation. The GM lines will be propagated by seed to at least the third generation from the transformation within glasshouses before the field trial. *Agrobacterium* is not normally transmitted from one generation to the next via seed, so is not expected to be present in the GMOs proposed for release.
   1. The receiving environment
4. The receiving environment forms part of the context in which the risks associated with dealings involving the GMOs are assessed. Relevant information about the receiving environment includes abiotic and biotic interactions of the crop with the environment where the release would occur; agronomic practices for the crop; presence of plants that are sexually compatible with the GMOs; and background presence of the gene(s) used in the genetic modification (OGTR, 2013a).
5. Detailed information about the commercial cultivation and distribution of canola and Indian mustard in Australia is presented in the document *The Biology of Brassica napus L. (canola) and Brassica juncea (L.) Czern. & Coss. (Indian mustard)* (OGTR, 2017).

5.1 Relevant abiotic factors

1. The proposed release would occur in a range of geographic and climatic regions in NSW, Victoria and Queensland. The most important abiotic factor limiting the geographical distribution of commercial canola and Indian mustard cultivation in Australia is water availability. Typically, canola can be grown in areas with annual rainfall over 325 mm and Indian mustard needs over 300 mm, although water requirements increase in hotter climates. Other abiotic stresses that can reduce canola or Indian mustard yield include soil acidity, waterlogging, frost and heat stress (GRDC, 2015, 2017).

5.2 Relevant biotic factors

1. The most important disease affecting canola and Indian mustard in Australia is blackleg, caused by the fungal pathogen *Leptosphaeria maculans.* Blackleg is most problematic in higher rainfall regions where a lot of canola is grown. Canola and Indian mustard can also be seriously damaged by stem rot caused by the fungus *Sclerotinia sclerotiorum* in wet springs. A range of other fungal or viral diseases sometimes reduce crop yield (McCaffery et al., 2009; GRDC, 2015, 2017).
2. Pests of canola and Indian mustard in eastern Australia include earth mites, aphids, moths and Rutherglen bugs (McCaffery et al., 2009; GRDC, 2015, 2017).
3. Canola is highly susceptible to weed competition during the early stages of growth (GRDC 2015). Indian mustard and hybrid canola have greater seedling vigour than open-pollinated canola and so are more competitive with weeds (McCaffery et al., 2009; GRDC, 2015, 2017). Common weeds of Australian canola crops include grassy weeds, volunteer cereals, and weeds from the *Brassicaceae* family including wild radish (*Raphanus raphanistrum*), Indian hedge mustard (*Sisymbrium orientale*), shepherds purse (*Capsella bursa‑pastoris*), wild turnip (*Brassica tournefortii*), charlock (*Sinapis arvensis*), turnip weed (*Rapistrum rugosum*) and Buchan weed (*Hirschfeldia incana*) (GRDC, 2015, 2017). To facilitate weed management, most canola varieties available in NSW and Victoria have a herbicide tolerance trait: imidizolinone tolerance, triazine tolerance, glyphosate tolerance (a GM trait) or dual-herbicide tolerance (Brown, 2021; Matthews et al., 2021).

5.3 Relevant agricultural practices

1. The applicant proposes that crop management practices for the GMOs would be the same as for commercial canola and Indian mustard crops, except for the proposed controls to restrict spread and persistence of the GMOs (see Section 2.2). Standard cultivation practices for canola and Indian mustard in eastern Australia are discussed elsewhere (GRDC, 2015, 2017).
2. The applicant specifies that small areas/rows would be hand-planted or planted with a small plot seeder, while larger areas would be planted with commercial equipment. Harvesting may occur by hand or with commercial equipment. Herbicides, pesticides and drip/pipe irrigation may be used as necessary to manage the health of the GM crop.

5.4 Presence of related plants in the receiving environment

1. Canola and Indian mustard are primarily self-pollinating, but approximately 30% of seeds are produced by cross‑pollination. Cross-pollination can be mediated by insects, wind or physical contact (OGTR, 2017).
2. Canola or Indian mustard have been reported to outcross in the field with the following species: *Brassica carinata*, *B. napus*, *B. juncea*, *B. oleracea,* *B. rapa*, *Hirschfeldia incana*, *Raphanus raphanistrum* and *Sinapis arvensis* (Ford et al., 2006; Warwick et al., 2009; Warwick and Martin, 2013). *B. carinata* is not known to be present in Australia ([Atlas of Living Australia](https://www.ala.org.au/), accessed 27 Jan 2022).
3. Canola (*B. napus*) is widely grown as an oilseed crop in NSW and Victoria, but rarely grown in Queensland (ABARES, 2021). The proposed trial sites in NSW and Victoria, but not Queensland, are likely to be located in commercial canola growing regions. Indian mustard (*B. juncea*) is a minor oilseed crop grown in similar areas to canola. Cabbage (*B. oleracea*) and turnip (*B. rapa*) are cultivated as horticultural crops. These four species are also naturalised in parts of NSW, Victoria and Queensland ([VICFLORA](https://vicflora.rbg.vic.gov.au/), accessed 27 Jan 2022).
4. Buchan weed (*H. incana*), wild radish (*R. raphanistrum*) and charlock (*S. arvensis*) are widespread weeds in NSW, Victoria and south-east Queensland ([New South Wales Flora Online](https://plantnet.rbgsyd.nsw.gov.au/floraonline.htm), accessed 27 Jan 2022; [Weeds Australia](https://weeds.org.au/), accessed 27 Jan 2022). As discussed in section 5.2, these species are common weeds in canola crops.

5.5 Presence of similar genes and their products in the environment

1. Five of the introduced genes involved in fatty acid biosynthesis are sourced from microalgae that are present in the marine environment (*Pavlova salina*, *Micromonas pusilla* and *Pyramimonas cordata)*. People may naturally encounter the genes and encoded proteins through contact with sea water or seafood. In addition, humans and other mammals have similar, endogenous genes encoding enzymes responsible for converting dietary ALA into ω‑3 LC PUFAs (Zarate et al., 2017; Shahidi and Ambigaipalan, 2018; Saini et al., 2021).
2. Two of the introduced genes involved in fatty acid biosynthesis are sourced from the yeasts *Pichia pastoris* and *Lachancea kluyveri*. These yeasts were isolated from trees and soil in the Northern Hemisphere. Both of these yeasts are present in the New Zealand environment (Zhang et al., 2010), likely due to import of host tree species from Europe, and are also expected to be present in Australia.
3. The *pat* gene was obtained from the common soil bacterium *Streptomyces viridochromogenes.* The *pat* gene or the similar *bar* gene from *S. hygroscopicus* are also present in many types of GM canola or cotton authorised for commercial release in Australia (licences DIR 021/2002, DIR 062/2005, DIR 091, DIR 108, DIR 138, DIR 143, DIR 145, DIR 155, DIR 173, DIR 175 and DIR 178).
   1. Relevant Australian and international approvals

6.1 Australian approvals

Approvals by the Regulator

1. GM canola lines containing all introduced genes proposed for release under the current application were previously approved for field trials under licences DIR 123 and DIR 163 and for commercial release under licence DIR 155. The GM canola line approved for commercial release under licence DIR 155 is known as DHA canola (NS-B50027-4).
2. GM Indian mustard lines containing all introduced genes proposed for release under the current application were previously approved for field trials under licence DIR 149.
3. There are no reported adverse effects from these previous releases.

Approvals by other government agencies

1. FSANZ is responsible for human food safety assessment and food labelling, including GM food. FSANZ has assessed the safety of food derived from DHA canola and approved its food products for commercial sale (FSANZ, 2017).

6.2 International approvals

1. DHA canola was deregulated for commercial cultivation in the United States in 2018. DHA canola was approved for food, feed and commercial cultivation in Canada in 2020.
2. Risk assessment
   1. Introduction
3. The risk assessment identifies and characterises risks to the health and safety of people or to the environment from dealings with GMOs, posed by or as the result of gene technology (Figure 4). Risks are identified within the established risk context (Chapter 1), taking into account current scientific and technical knowledge. A consideration of uncertainty, in particular knowledge gaps, occurs throughout the risk assessment process.



**Figure 4.** The risk assessment process

1. The Regulator uses a number of techniques to identify risks, including checklists, brainstorming, reported international experience and consultation (OGTR, 2013a). A weed risk assessment approach is used to identify traits that may contribute to risks from GM plants, as this approach addresses the full range of potential adverse outcomes associated with plants. In particular, novel traits that may increase the potential of the GMO to spread and persist in the environment or increase the level of potential harm compared with the parental plant(s) are used to postulate risk scenarios (Keese et al., 2014). Risk scenarios examined in RARMPs prepared for licence applications for the same or similar GMOs are also considered.
2. Risk identification first considers a wide range of circumstances in which the GMO, or the introduced genetic material, could come into contact with people or the environment. This leads to postulating plausible causal pathways that may give rise to harm for people or the environment from dealings with a GMO. These are risk scenarios. These risk scenarios are screened to identify those that are considered to have a reasonable chance of causing harm in the short or long term. Pathways that do not lead to harm, or those that could not plausibly occur, do not advance in the risk assessment process (Figure 4), i.e., the risk is considered to be no greater than negligible.
3. Risks identified as being potentially greater than negligible are characterised in terms of the potential seriousness of harm (Consequence assessment) and the likelihood of harm (Likelihood assessment). Risk evaluation then combines the Consequence and Likelihood assessments to estimate the level of risk and determine whether risk treatment measures are required. The potential for interactions between risks is also considered.
   1. Risk identification
4. Postulated risk scenarios have three components (Figure 5):
5. the source of potential harm (risk source)
6. a plausible causal linkage to potential harm (causal pathway)
7. potential harm to people or the environment.

**source of**

**potential harm**

(a novel GM trait)

**plausible causal linkage**

**potential harm to**

**an object of value**

(people/environment)

**Figure 5.** Risk scenario

1. When postulating relevant risk scenarios, the risk context is taken into account, including the following factors detailed in Chapter 1:

the proposed dealings

the proposed limits including the extent and scale of the proposed dealings

the proposed controls to restrict the spread and persistence of the GMOs and

the characteristics of the parent organism(s).

* + 1. Risk source

1. The sources of potential harms can be intended novel GM traits associated with one or more introduced genetic elements, or unintended effects/traits arising from the use of gene technology.
2. As discussed in Chapter 1, the GM canola and Indian mustard lines have been modified by the introduction of up to seven genes involved in fatty acid biosynthesis. These introduced genes will be considered further as a source of potential harm.
3. The GM canola and Indian mustard lines may also contain the introduced *pat* gene which confers tolerance to glufosinate herbicide and was used as a selectable marker. The *pat* gene and its products have been extensively characterised and assessed as posing negligible risk to human and animal health or to the environment by the Regulator, as well as by other regulatory agencies in Australia and overseas (CERA, 2011). Commercial GM canola lines containing the *pat* gene have been assessed to pose negligible risks to human health and the environment in the RARMPs for DIR 021/2002 (OGTR, 2003), DIR 108 (OGTR, 2011) and DIR 155 (OGTR, 2018a). In addition, a herbicide tolerance trait has no effect except in an environment where the plant is exposed to the relevant herbicide, and it is unlikely that the proposed GM canola and Indian mustard lines would be exposed to glufosinate herbicide in the field. This is because the applicant does not propose to use glufosinate during the field trial, glufosinate is not used to control volunteer canola (AOF, 2019), and although glufosinate is registered for weed control in summer fallows, it is more expensive and infrequently used compared to the alternative knockdown herbicides glyphosate and paraquat (Walsh, 2021). For these reasons, the *pat* gene will not be further considered as a source of potential harm.
4. The introduced genes are controlled by regulatory sequences. These were originally derived from plants, plant viruses and a bacterium (Table 3). Regulatory sequences are naturally present in all plants, and the introduced elements are expected to operate in similar ways to endogenous elements. These sequences are DNA that is not expressed as a protein, so exposure is to the DNA only and dietary DNA has no toxicity (Society of Toxicology, 2003). Hence, potential harms from the regulatory sequences will not be further assessed for this application.
5. Genetic modifications involving introduction of genes have the potential to cause unintended effects in several ways. These include insertional effects such as interruptions, deletions, duplications or rearrangements of the genome, which can lead to altered expression of endogenous genes. There could also be increased metabolic burden due to expression of the introduced proteins, novel traits arising out of interactions with non-target proteins and secondary effects arising from altered substrate or product levels in biochemical pathways. However, these types of effects also occur spontaneously and in plants generated by conventional breeding. Accepted conventional breeding techniques such as hybridisation, mutagenesis and somaclonal variation can have a much larger impact on the plant genome than genetic engineering (Schnell et al., 2015). Plants generated by conventional breeding have a long history of use, and there are no documented cases where conventional breeding has resulted in the production of a novel toxin or allergen in a crop (Steiner et al., 2013). Therefore, the potential for the processes of genetic modification to result in unintended effects will not be considered further.
   * 1. Causal pathway
6. The following factors are taken into account when postulating plausible causal pathways to potential harm:

routes of exposure to the GMOs, the introduced gene(s) and gene product(s)

potential exposure to the introduced gene(s) and gene product(s) from other sources in the environment

the environment at the site(s) of release

agronomic management practices for the GMOs

spread and persistence of the GMOs (e.g. reproductive characteristics, dispersal pathways and establishment potential)

tolerance to abiotic conditions (e.g. climate, soil and rainfall patterns)

tolerance to biotic stressors (e.g. pest, pathogens and weeds)

tolerance to cultivation management practices

gene transfer to sexually compatible organisms

gene transfer by horizontal gene transfer (HGT)

unauthorised activities.

1. Although all of these factors are taken into account, some are not included in risk scenarios because they have been considered in previous RARMPs.
2. The potential for horizontal gene transfer (HGT) from GMOs to species that are not sexually compatible, and any possible adverse outcomes, have been reviewed in the literature (Keese 2008) and assessed in many previous RARMPs. HGT was most recently considered in the RARMP for DIR 108 (OGTR, 2011). HGT events rarely occur and the wild-type gene sequences are already present in the environment and available for transfer via demonstrated natural mechanisms. Therefore, no substantive risk was identified in previous assessments and HGT will not be further considered for this application.
3. The potential for unauthorised activities to lead to an adverse outcome has been considered in many previous RARMPs, most recently in the RARMP for DIR 117 (OGTR, 2013b). In previous assessments of unauthorised activities, no substantive risk was identified. The Act provides for substantial penalties for unauthorised dealings with GMOs or non-compliance with licence conditions, and also requires the Regulator to have regard to the suitability of an applicant to hold a licence prior to the issuing of the licence. These legislative provisions are considered sufficient to minimise risks from unauthorised activities. Therefore, unauthorised activities will not be considered further.
   * 1. Potential harm
4. Potential harms from GM plants are based on those used to assess risk from weeds (Virtue, 2008; Keese et al., 2014) including:

harm to the health of people or desirable organisms, including toxicity/allergenicity

reduced biodiversity through harm to other organisms or ecosystems

reduced establishment or yield of desirable plants

reduced products or services from the land use

restricted movement of people, animals, vehicles, machinery and/or water

reduced quality of the biotic environment (e.g. providing food or shelter for pests or pathogens) or abiotic environment (e.g. negative effects on fire regimes, nutrient levels, soil salinity, soil stability or soil water table).

1. Judgements of what is considered harm depend on the management objectives of the land where the GM plant may be present. A plant species may have different weed risk potential in different land uses such as dryland cropping or nature conservation.
   * 1. Postulated risk scenarios
2. Three risk scenarios were postulated and screened to identify any substantive risks. These scenarios are summarised in Table 4 and examined in detail in Sections 2.4.1 – 2.4.3.
3. In the context of the activities proposed by the applicant and considering both the short and long term, none of the risk scenarios gave rise to any substantive risks.

**Table 4.** Summary of risk scenarios from the proposed dealings with GM canola and Indian mustard

| **Risk scenario** | **Risk source** | **Causal pathway** | **Potential harm** | **Substantive risk?** | **Reason** |
| --- | --- | --- | --- | --- | --- |
| 1 | Introduced genes for altered oil content | Cultivation of GM canola and Indian mustard at trial sites  🡇  Exposure of people and desirable animals to products of the introduced genes | Increased toxicity or allergenicity for people  OR  increased toxicity to desirable animals | No | * The GM canola and Indian mustard would not be used as commercial human food or animal feed * The limits and controls of the field trial would restrict exposure of people and desirable animals to the GM plants * The proteins encoded by the introduced genes are not expected to be toxic or allergenic * Oil containing ω-3 LC-PUFAs is not expected to be toxic * Oxidation products of ω-3 LC‑PUFAs could be toxic, but people will not consume these products, and animals are highly unlikely to consume toxic levels due to low oxidation rates over the expected lifetime of seeds |
| 2 | Introduced genes for altered oil content | Cultivation of GM canola and Indian mustard at trial sites  🡇  Dispersal of GM seed outside trial limits  🡇  Establishment of populations of volunteer GM plants expressing the introduced genes in the environment | Increased toxicity or allergenicity for people  OR  increased toxicity to desirable animals  OR  reduced establishment or yield of desirable plants | No | * The controls of the field trial would minimise dispersal or persistence of GM seeds * GM canola and Indian mustard are susceptible to standard weed management measures * As discussed in Risk Scenario 1, the genetic modifications are not expected to cause increased toxicity or allergenicity * Canola and Indian mustard have limited ability to compete with other plants and the genetic modifications are not expected to increase their competitiveness |
| 3 | Introduced genes for altered oil content | Cultivation of GM canola and Indian mustard at trial sites  🡇  Pollen from GM plants dispersed outside the trial sites  🡇  Outcrossing with sexually compatible plants  🡇  Establishment of populations of hybrid GM plants expressing the introduced genes in the environment | Increased toxicity or allergenicity for people  OR  increased toxicity to desirable animals  OR  reduced establishment or yield of desirable plants | No | * The controls of the field trial would minimise pollen flow to sexually compatible plants outside the trial sites * As discussed in Risk Scenario 1, the genetic modifications are not expected to cause increased toxicity or allergenicity * As discussed in Risk Scenario 2, the genetic modifications are not expected to increase ability to compete with other plants |

* + - 1. Risk scenario 1

|  |  |
| --- | --- |
| *Risk source* | Introduced genes for altered oil content |
| *Causal pathway* | 🡇  Cultivation of GM canola and Indian mustard at trial sites  🡇  Exposure of people and desirable animals to products of the introduced genes  🡇 |
| *Potential harm* | Increased toxicity or allergenicity for people  OR  Increased toxicity to other desirable organisms |

**Risk source**

1. The source of potential harm for this postulated risk scenario is the introduced genes for altered oil content in GM canola and Indian mustard plants.

**Causal pathway**

1. The GM canola and Indian mustard would be grown at the trial sites. As the introduced genes for altered oil content are controlled by seed-specific promoters, the encoded proteins would be produced in the GM seeds. The seed oil is expected to be enriched in target ω-3 LC-PUFAs and may also be enriched in fatty acids that are intermediates in the biosynthesis pathway of the target ω-3 LC-PUFAs. People and desirable animals could be exposed to GM seeds containing the introduced proteins and seed oil enriched in ω‑3 LC-PUFAs.
2. The GM canola and Indian mustard would not be used for commercial human food. Only authorised and trained trial staff would be permitted to deal with the GM plants and their seeds. Therefore, there is little potential for the public to be exposed to GM seeds grown at the trial sites.
3. Trial staff would handle the GM seeds and plant material produced by processing of the GM seeds. Workers could be exposed to the introduced proteins and seed oil enriched in ω‑3 LC-PUFAs by dermal contact and inhalation.
4. The applicant proposes human sensory trials of GM plant material to test the feel, smell, taste and appearance of the seed oil or food products containing the oil. These tests would not involve consumption of the oil. Canola oil is highly refined and does not contain detectable amounts of protein. People participating in sensory trials could be exposed to seed oil enriched in ω-3 LC-PUFAs by dermal contact, contact with mucous membranes and inhalation.
5. The GM canola and Indian mustard would not be used for commercial animal feed and livestock would not be permitted to graze the trial sites. Therefore, livestock are not expected to be exposed to GM seeds grown at the trial sites.
6. Desirable wild animals, such as native mammals and birds, could enter the trial sites and consume GM seeds. The limited size and duration of the field trial would restrict the number of desirable wild animals exposed to GM seeds grown at the trial sites.
7. The applicant proposes animal feeding trials with GM plant material, which may include trials with rodents, chickens and farmed fish species. The experimental animals would be exposed to the introduced proteins and/or seed oil enriched in ω-3 LC-PUFAs.

**Potential harm**

1. As discussed in Chapter 1, Section 4.2, none of the introduced proteins are expected to be toxic or allergenic.
2. As discussed in Chapter 1, Section 4.3, the target ω-3 LC-PUFAs and fatty acid intermediates are not expected to be toxic.
3. As discussed in Chapter 1, Section 4.3, ω-3 LC-PUFAs are highly susceptible to oxidation, and the oxidation products can have toxic effects if consumed. There is little published information regarding the level of toxicity of ω-3 LC-PUFA oxidation products. When pregnant rats were fed heavily oxidised fish oil as a large component of their diet for the entire period of pregnancy, this caused increased newborn mortality but did not increase mortality of the mothers (Albert et al., 2016). Therefore, if GM seed oil were heavily oxidised, this could increase mortality of neonate animals whose mothers consumed the GM seed. It is uncertain whether toxic effects could occur in other animals feeding on the GM seed, especially animals that may be particularly sensitive to the oxidation products.
4. A study of a GM canola line containing all of the introduced genes for altered oil content tested the stability of seed oil during six months of seed storage after harvest, and did not detect any changes in the levels of ω-3 LC-PUFAs (Petrie et al., 2020). This suggests that oxidation of ω-3 LC-PUFAs in seeds occurs at a very slow rate under storage conditions, so GM seeds stored for planting or experimentation would not contain heavily oxidised oil. Oxidation may occur more rapidly in seeds that are lost during harvest and remain on the soil surface, as the oxidation rate of ω-3 LC-PUFAs is increased by exposure to air and light (Arab-Tehrany et al., 2012; Albert et al., 2013; Miyashita, 2019). However, seeds on the soil surface are very susceptible to predation and would probably be consumed before much oxidation could occur. For example, in a Canadian study where canola seeds were left on the soil surface, 42 ̶ 77% of seeds were consumed by invertebrate seed predators within a week (Kulkarni et al., 2017). Therefore, desirable animals are highly unlikely to be exposed to toxic levels of ω-3 LC-PUFA oxidation products through consumption of GM seeds. The timeframes for oxidation of ω-3 LC-PUFAs in seeds under different conditions are an area of uncertainty for this risk assessment.
5. As discussed in Chapter 1, Section 6.1, GM canola and Indian mustard lines containing all of the introduced genes for altered oil content have been previously approved by the Regulator for field trials and commercial release. To date, no adverse effects have been reported from these releases.

**Conclusion**

1. Risk scenario 1 is not identified as a substantive risk because the limits and controls of the field trial would restrict exposure of people and desirable animals to the GM plants, the introduced proteins are not expected to be toxic or allergenic, and oil containing ω-3 LC-PUFAs is not expected to be toxic. Although oxidation products of ω-3 LC PUFAs could be toxic, people will not consume these products and animals are highly unlikely to consume toxic levels. Therefore, this risk could not be considered greater than negligible and does not warrant further detailed assessment.
   * + 1. Risk scenario 2

|  |  |
| --- | --- |
| *Risk source* | Introduced genes for altered oil content |
| *Causal pathway* | 🡇  Cultivation of GM canola and Indian mustard at trial sites  🡇  Dispersal of GM seed outside trial limits  🡇  Establishment of populations of volunteer GM plants expressing the introduced genes in the environment  🡇 |
| *Potential harm* | Increased toxicity or allergenicity for people  OR  Increased toxicity to desirable animals  OR  Reduced establishment or yield of desirable plants |

**Risk source**

1. The source of potential harm for this postulated risk scenario is the introduced genes for altered oil content in GM canola and Indian mustard plants.

**Causal pathway**

1. The GM canola and Indian mustard would be grown at the trial sites. GM seeds could be physically dispersed outside the trial sites by human activity, animal activity, wind or water. GM seeds could also persist on trial sites after completion of the trial. These GM seeds could grow in the environment and establish populations of volunteer GM plants.
2. Viable GM canola and Indian mustard seeds could be dispersed outside the trial sites by human activity, such as transport of seeds and movement of agricultural machinery. To minimise dispersal of GM seeds by human activity, the applicant proposes to clean all equipment used with the GM plants after use, and to transport all GM seed in accordance with the Regulator’s guidelines for the transport of GMOs.
3. GM seeds could be dispersed outside the trial sites by animal activity. Canola and Indian mustard seeds have no specific adaptions, such as burrs or hooks, for dispersal by animals (OGTR, 2017). Dispersal of viable canola seed via endozoochory (consumption and excretion of seed) by birds only occurs at very low levels (Twigg et al., 2008; Woodgate et al., 2011). Canola and Indian mustard seeds could be transported short distances by hoarding animals, such as ants and mice. The applicant proposes that monitoring zones around trial sites would be inspected for volunteers.
4. Canola and Indian mustard seeds lack specialised structures that would assist their dispersal by wind (OGTR, 2017). However, the GM canola may be windrowed prior to harvesting, and under strong wind conditions plant material could disperse outside trial sites. The applicant proposes that monitoring zones around trial sites would be inspected for volunteers.
5. GM canola and Indian mustard seeds could be dispersed by water during flooding or heavy runoff, although seeds are unlikely to remain viable after prolonged exposure to water (OGTR, 2017). The applicant proposes to locate the trial sites at least 50 m from waterways to minimise the potential for seed dispersal during flooding.
6. During harvest of the GM canola and Indian mustard, a small percentage of the GM seeds are expected to be lost and to remain on the trial sites. Viable canola and Indian mustard seeds can persist in the seedbank for several years (OGTR, 2017). It is unlikely that the genetic modifications for altered seed oil composition would affect seed persistence. A field study of seedbank persistence in a GM canola line with altered oil content found no difference between seedbank persistence of the GM line and the non-GM control at the completion of the trial, which was 14-19 months after seed burial (Linder and Schmitt, 1995). To minimise persistence of residual GM seeds on the trial sites, the applicant proposes to promote seed germination by light post-harvest tillage and irrigation. During a post‑harvest monitoring period, the applicant would regularly inspect the trial sites and destroy any GM volunteers, until volunteers cease to emerge.
7. The suitability of the proposed controls to manage GM seed dispersal and persistence is discussed in detail in Chapter 3, Section 3.1.
8. If GM canola and Indian mustard seeds were dispersed outside trial limits, it is unlikely that they would establish ongoing volunteer populations. Even in environments without active weed management, volunteer canola populations along transportation routes rely on recurrent spillages to persist (Yoshimura et al., 2006) and volunteer canola dispersed into natural areas was reported to rapidly become extinct (Busi and Powles, 2016). The genetic modifications for altered seed oil composition are not expected to affect the ability of volunteers to survive in the environment. A GM canola line containing the introduced genes had no changes in agronomic traits compared to the non-GM control (Petrie et al., 2020).
9. In agricultural areas of Australia where canola and Indian mustard are grown, volunteer populations are controlled by weed management measures. Effective methods for control of canola volunteers include grazing, mowing, cultivation and application of a range of knockdown or selective herbicides (AOF, 2019). The genetic modifications for altered seed oil composition would not affect the susceptibility of GM volunteers to standard weed management measures.

**Potential harm**

1. As discussed in risk scenario 1, it is not expected that the GM canola and Indian mustard would have increased toxicity or allergenicity for people or increased toxicity to desirable animals.
2. Populations of volunteer GM canola and Indian mustard could reduce establishment or yield of desirable plants. GM volunteers could directly compete with agricultural crops, pastures or native vegetation. GM volunteers could also reduce yield of commercial canola and Indian mustard crops by providing a reservoir for pathogens, such as the important fungal diseases blackleg and stem rot (see Chapter 1, Section 5.2).
3. Canola is considered a less competitive crop species than wheat or barley (GRDC, 2011), which are the main crops grown in eastern Australia (ABARES, 2021). Indian mustard has a similar phenotype to canola, although it may be slightly more competitive due to greater seedling vigour (GRDC, 2015, 2017). All domesticated crop plant species are expected to be poor competitors with pasture species or established native vegetation. Therefore, canola and Indian mustard volunteers have limited ability to compete with desirable plants. The genetic modifications for altered seed oil composition are not expected to increase the competitiveness of GM plants. The biological purpose of plant seed oil is to provide an energy source for germination and seedling establishment, and highly unsaturated oil provides less energy per carbon atom than saturated or monounsaturated oil (Sanyal and Decocq, 2016).
4. Blackleg and stem rot diseases affect vegetative parts of canola and Indian mustard plants rather than seeds (GRDC, 2015, 2017). The genetic modifications for altered seed oil are not expected to increase the ability of GM plants to act as reservoirs for these pathogens.

**Conclusion**

1. Risk scenario 2 is not identified as a substantive risk because the controls of the field trial would minimise dispersal or persistence of GM seeds, GM canola and Indian mustard are susceptible to standard weed management measures, the genetic modifications are not expected to increase toxicity or allergenicity, and the genetic modifications are not expected to increase competitiveness with other plants. Therefore, this risk could not be considered greater than negligible and does not warrant further detailed assessment.
   * + 1. Risk scenario 3

|  |  |
| --- | --- |
| *Risk source* | Introduced genes for altered oil content |
| *Causal pathway* | 🡇  Cultivation of GM canola and Indian mustard at trial sites  🡇  Pollen from GM plants dispersed outside the trial sites  🡇  Outcrossing with sexually compatible plants  🡇  Establishment of populations of hybrid GM plants expressing the introduced genes in the environment  🡇 |
| *Potential harm* | Increased toxicity or allergenicity for people  OR  Increased toxicity to desirable animals  OR  Reduced establishment or yield of desirable plants |

**Risk source**

1. The source of potential harm for this postulated risk scenario is the introduced genes for altered oil content in GM canola and Indian mustard plants.

**Causal pathway**

1. The GM canola and Indian mustard would be grown at the trial sites. Pollen from the GM plants could be transported out of the trial sites by wind or insect vectors and fertilise sexually compatible plants. Hybrid seeds containing the introduced genes could be harvested by farmers and planted as a crop or could grow as volunteers.
2. It should be noted that vertical gene flow *per se* is not considered an adverse outcome, but may be a link in a chain of events that may lead to an adverse outcome.
3. Canola and Indian mustard are primarily self-pollinating, but approximately 30% of seeds are produced by cross pollination. Outcrossing decreases rapidly with distance, with the majority of cross-pollination occurring over distances less than 10 m (OGTR, 2017). The introduced genes for altered oil content are only expressed in seeds and are not expected to affect the pollen dispersal characteristics of the GM canola and Indian mustard.
4. The GM canola and Indian mustard could outcross with nearby canola or Indian mustard crops or volunteers, if there is synchronicity of flowering. As discussed in Chapter 1, Section 5.4, canola or Indian mustard can also occasionally hybridise with the related horticultural crops *B. oleracea* and *B. rapa* and the related weeds *H. incana*, *R. raphanistrum* and *S. arvensis.*
5. The applicant has proposed control measures to minimise pollen flow from GM plants growing on the trial sites to sexually compatible plants outside the trial sites (Chapter 1, Section 2.2). During flowering of the GM plants, each planting area would be (a) covered by an insect proof tent and surrounded by a monitoring zone and isolation zone, or (b) surrounded by a pollen trap, monitoring zone and isolation zone, or (c) surrounded by a monitoring zone and a large isolation zone. In addition, any GM volunteers growing on the trial sites after harvest would be destroyed prior to flowering.
6. The suitability of the proposed controls to manage pollen flow is discussed in detail in Chapter 3, Section 3.1.
7. If pollen from GM plants fertilised plants in a commercial canola or Indian mustard crop, the farmer could harvest some of the crop as planting seed and plant hybrid GM seeds in a crop in the next growing season. Hybrid GM plants could then enter commercial human food and animal feed. However, even in the complete absence of measures to restrict pollen flow, outcrossing rates between neighbouring commercial canola fields are less than 0.1% under Australian conditions (Rieger et al., 2002). Therefore, the planting seed described in this risk pathway could only contain a very low proportion of hybrid GM seed, so people and desirable animals could only be exposed to very low levels of the hybrid GMOs.
8. If pollen from GM plants fertilised sexually compatible plants growing as crops, volunteers or weeds, the hybrid GM seeds could grow as volunteers. Populations of hybrid GM volunteers could be consumed by desirable animals or could reduce the establishment or yield of desirable plants.

**Potential harm**

1. As discussed in risk scenario 1, the GM canola and Indian mustard are not expected to have greater toxicity or allergenicity for people or greater toxicity to desirable animals than non-GM canola or Indian mustard. Similarly, in hybrids between the GM plants and sexually compatible plants, the genetic modifications would not increase toxicity or allergenicity.
2. As discussed in risk scenario 2, the GM canola and Indian mustard are not expected to have greater competitiveness than non-GM canola or Indian mustard. Similarly, in hybrids between the GM plants and sexually compatible plants, the genetic modifications would not increase ability to compete with other plants.

**Conclusion**

1. Risk scenario 3 is not identified as a substantive risk because the controls of the field trial would minimise pollen flow to sexually compatible plants outside the trial sites, the genetic modifications are not expected to cause increased toxicity or allergenicity, and the genetic modifications are not expected to increase the ability to compete with other plants. Therefore, this risk could not be considered greater than negligible and does not warrant further detailed assessment.
   1. Uncertainty
2. Uncertainty is an intrinsic property of risk analysis and is present in all aspects of risk analysis. This is discussed in detail in the Regulator’s [Risk Analysis Framework](https://www.ogtr.gov.au/resources/publications/risk-analysis-framework-2013) document.
3. Uncertainty is addressed by approaches such as balance of evidence, conservative assumptions, and applying risk management measures that reduce the potential for risk scenarios involving uncertainty to lead to harm. If there is residual uncertainty that is important to estimating the level of risk, the Regulator will take this uncertainty into account in making decisions.
4. As field trials of GMOs are designed to gather data, there are generally data gaps when assessing the risks of a field trial application. However, field trial applications are required to be limited and controlled. Even if there is uncertainty about the characteristics of a GMO, limits and controls restrict exposure to the GMO, and thus decrease the likelihood of harm.
5. For DIR 188, uncertainty is noted particularly in relation to the potential for toxicity of ω-3 LC-PUFA oxidation products in seeds of the GM canola and Indian mustard.
6. Additional data, including information to address these uncertainties, may be required to assess possible future applications with reduced limits and controls, such as a larger scale trial or the commercial release of these GMOs.
7. Chapter 3, Section 4, discusses information that may be required for future release.
   1. Risk evaluation
8. Risk is evaluated against the objective of protecting the health and safety of people and the environment to determine the level of concern and, subsequently, the need for controls to mitigate or reduce risk. Risk evaluation may also aid consideration of whether the proposed dealings should be authorised, need further assessment, or require collection of additional information.
9. Factors used to determine which risks need treatment may include:

risk criteria

level of risk

uncertainty associated with risk characterisation

interactions between substantive risks.

1. Three risk scenarios were postulated whereby the proposed dealings might give rise to harm to people or the environment. In the context of the limits and controls proposed by the applicant, and considering both the short and long term, none of these scenarios were identified as substantive risks. The principal reasons for these conclusions are summarised in Table 4 and include:

the GM plants would not be used as commercial human food or animal feed

limits on the size and duration of the proposed release

controls proposed by the applicant to restrict the spread and persistence of the GM canola and Indian mustard plants and their genetic material (see Chapter 3 for their suitability)

the products of the introduced genes are not expected to be toxic or allergenic

GM canola and Indian mustard volunteers could be controlled by standard weed management measures.

1. Therefore, risks to the health and safety of people, or the environment, from the proposed release of the GM canola and Indian mustard plants into the environment are considered to be negligible. The *Risk Analysis Framework* (OGTR, 2013a) which guides the risk assessment and risk management process, defines negligible risks as risks of no discernible concern with no present need to invoke actions for mitigation. Therefore, no additional controls are required to treat these negligible risks. The Regulator considers that the dealings involved in this proposed release do not pose a significant risk to either people or the environment[[1]](#footnote-1).
2. Risk management plan
   1. Background
3. Risk management is used to protect the health and safety of people and to protect the environment by controlling or mitigating risk. The risk management plan addresses risks evaluated as requiring treatment and considers limits and controls proposed by the applicant, as well as general risk management measures. The risk management plan informs the Regulator’s decision-making process and is given effect through licence conditions.
4. Under Section 56 of the Act, the Regulator must not issue a licence unless satisfied that any risks posed by the dealings proposed to be authorised by the licence are able to be managed in a way that protects the health and safety of people and the environment.
5. All licences are subject to three conditions prescribed in the Act. Section 63 of the Act requires that each licence holder inform relevant people of their obligations under the licence. The other statutory conditions allow the Regulator to maintain oversight of licensed dealings: Section 64 requires the licence holder to provide access to premises to OGTR inspectors and Section 65 requires the licence holder to report any information about risks or unintended effects of the dealing to the Regulator on becoming aware of them. Matters related to the ongoing suitability of the licence holder must also be reported to the Regulator.
6. The licence is also subject to any conditions imposed by the Regulator. Examples of the matters to which conditions may relate are listed in Section 62 of the Act. Licence conditions can be imposed to limit and control the scope of the dealings and to manage risk to people or the environment. In addition, the Regulator has extensive powers to monitor compliance with licence conditions under Section 152 of the Act.
   1. Risk treatment measures for substantive risks
7. The risk assessment of risk scenarios listed in Chapter 2 concluded that there are negligible risks to people or the environment from the proposed field trial of GM canola and Indian mustard. These risk scenarios were considered in the context of the scale of the proposed release (Chapter 1, Section 2.1), the proposed control measures (Chapter 1, Section 2.2), and the receiving environment (Chapter 1, Section 5), and considering both the short and the long term. The risk evaluation concluded that no specific risk treatment measures are required to treat these negligible risks. Limits and controls proposed by the applicant and other general risk management measures are discussed below.
   1. General risk management
8. The limits and controls proposed in the application were important in establishing the context for the risk assessment and in reaching the conclusion that the risks posed to people and the environment are negligible. Therefore, to maintain the risk context, licence conditions have been proposed to limit the release to the proposed size, location and duration, and to restrict the spread and persistence of the GMOs and their genetic material in the environment. The conditions are discussed and summarised in this Chapter and listed in detail in the draft licence.
   * 1. Draft licence conditions to limit and control the release
9. Sections 2.1 and 2.2 of Chapter 1 provide details of the limits and controls proposed by Nuseed in their application. Many of these are discussed in the three risk scenarios considered for the proposed release in Chapter 2. The appropriateness of these limits and controls is considered further in the following sections.
   * + 1. Consideration of limits proposed by Nuseed
10. The applicant proposes that the field trial would take place at up to twenty sites per year. Ten of these sites would have a maximum planting area of 10 ha and ten sites would have a maximum planting area of 5 ha, so the total trial area would be up to 150 ha/year. Sites would be selected from 96 local government areas in NSW, Victoria and Queensland. The duration of the field trial would be five years, from November 2022 to December 2027. The limited size and duration of the trial would restrict the potential exposure of people and desirable animals to the GMOs (Risk Scenario 1).
11. The applicant proposes that only trained and authorised staff would be permitted to deal with the GMOs. These limits would restrict the number of people exposed to the GMOs (Risk Scenario 1).
    * + 1. Consideration of proposed controls regarding exposure to the GMOs
12. The applicant proposes that GM plant material would not be used in commercial human food or animal feed. The applicant proposes to use GM plant material in animal feeding trials and possibly human taste testing trials (as part of broader sensory trials). The draft licence requires that GM plant material must not be used as food for humans or feed for animals, except for use in specified animal feeding experiments or taste testing experiments. Animal feeding experiments must be approved by an Animal Ethics Committee operating under the Australian Code for the Care and Use of Animals for Scientific Purposes and taste testing experiments must be approved by a Human Research Ethics Committee in accordance with the National Statement on Ethical Conduct in Human Research. These conditions would restrict the exposure of people and desirable animals to the GMOs (Risk Scenario 1).
    * + 1. Consideration of proposed controls regarding pollen flow from the GMOs
13. The applicant proposes three different options to control pollen flow from the trial sites while the GMOs are flowering.
14. The first option to control pollen flow is to surround the planting area with a 50 m monitoring zone and a 1 km isolation zone. The GMOs would not be planted at a trial site if any plants that are sexually compatible with canola or Indian mustard were being grown in the monitoring or isolation zones. The monitoring zone would be inspected at least once every 35 days from 14 days prior to flowering of the GMOs until the GMOs are harvested, to ensure that it is free from any sexually compatible plants. The isolation zone would be inspected at least once every 35 days from 14 days prior to flowering of the GMOs until the GMOs complete flowering, to ensure that it is free from intentionally planted sexually compatible plants. This option was proposed for previous GM canola field trials and was considered an effective means of restricting pollen flow from canola (e.g. DIR 164 (OGTR, 2018b)).
15. The pattern of pollen movement for *B. juncea* is similar to *B. napus* (Salisbury, 2006). The Canadian Regulations and Procedures for Pedigreed Seed Crop Production (CSGA, 2022) require that foundation production of male sterile *B. juncea* or *B. napus* seed must be separated from other *B. juncea* or *B. napus* plants by an 800 m isolation distance, of which the first 50 m must be practically free from related plants, and the remaining distance must be reasonably free from related plants. Therefore, the proposed 50 m monitoring zone and 1 km isolation zone, which are more stringent than these Canadian requirements, are considered effective measures to restrict pollen flow from Indian mustard.
16. The second option to control pollen flow is to surround the planting area with a 15 m pollen trap of non‑GM canola plants, a 50 m monitoring zone and a 400 m isolation zone. The pollen trap would be managed to flower at the same time as the GMOs. Pollen trap plants may provide sufficient forage for incoming pollinating insects that they do not visit the GM plants, and any insects that reach the GM plants are expected to deposit most GM pollen on pollen trap plants while exiting the trial site. Pollen trap plants may also absorb some pollen dispersed by wind. Therefore, the use of a pollen trap justifies reducing the isolation zone from 1 km to 400 m. As a non-GM pollen trap or buffer zone can also serve the same function as an unplanted monitoring zone (Hüsken and Dietz-Pfeilstetter, 2007), it is considered unnecessary to have both a pollen trap and a full-sized 50 m monitoring zone. The draft licence permits the applicant to use a 15 m pollen trap combined with a 35 m monitoring zone.
17. The third option to control pollen flow is to cover the planting area with an insect proof tent, and to surround the planting area with a 10 m monitoring zone and a 400 m isolation zone. The tents would be in place from at least seven days before flowering until the GMOs complete flowering, and would be inspected for damage fortnightly and after any extreme weather event. The tents are expected to prevent all insect‑mediated pollen flow and to greatly reduce wind-mediated pollen flow. Therefore, the use of an insect-proof tent justifies a reduced monitoring zone and isolation zone.
18. The proposed measures to control pollen flow would minimise outcrossing between the GMOs grown on the trial sites and sexually compatible plants growing outside the trial sites (risk scenario 3).
19. After harvest of the trial sites, the applicant proposes to monitor the sites for volunteers (see Section 3.1.5). The applicant would inspect at least once every 35 days, in order to find and destroy volunteers before they flower. These post-harvest inspections were proposed for previous GM canola field trials and were considered an effective means of restricting pollen flow from GM canola volunteers to plants outside the trial sites (e.g. DIR 164 (OGTR, 2018b)).
20. However, studies from Australia and North America have reported that canola-quality *B. juncea* varieties often begin flowering earlier than comparable canola varieties (Gunasekera et al., 2006; Gan et al., 2007; Riar, 2015; Hunter et al., 2017). Canola-quality *B. juncea* varieties planted in spring were reported to begin flowering 44-46 days after sowing (Gan et al., 2007; Hunter et al., 2017). This suggests that GM Indian mustard volunteers germinating in spring or summer could flower around 35-40 days after emergence. Therefore, the draft licence requires post-harvest inspections at least once every 30 days for any trial sites that grew Indian mustard and at least once every 35 days for any trial sites that grew only canola. These post-harvest monitoring requirements would minimise outcrossing between GM volunteers and sexually compatible plants growing outside the trial sites (risk scenario 3).
    * + 1. Consideration of proposed controls regarding dispersal of the GMOs
21. The applicant proposes to treat any non-GM canola and Indian mustard plants grown in planting areas or pollen traps like the GMOs. These non-GM plants may be mingled with or fertilised by the GM plants and it is therefore necessary to handle the non-GM plants in the same way as the GMOs to manage the dispersal or persistence of GM seed.
22. The applicant proposes that the GM canola and Indian mustard would be harvested separately from other crops, to avoid inadvertent seed mixing. Any equipment used with the GMOs would be cleaned as soon as practicable and before use for any other purpose, to avoid movement of viable plant material together with equipment. The applicant would contain the GM seeds during transport and storage in accordance with the Regulator’s [Guidelines for the Transport, Storage and Disposal of GMOs](https://www.ogtr.gov.au/resources/publications/guidelines-transport-storage-and-disposal-gmos). These measures would minimise human-mediated dispersal of GM seeds (risk scenario 2)
23. The applicant proposes to locate trial sites at least 50 m away from waterways. In addition, the draft licence requires that trial sites must not be located in flood-prone areas, and that any extreme weather events must be reported to the Regulator. These measures would minimise dispersal of GM seeds by flooding (risk scenario 2).
24. GM canola and Indian mustard seeds could be dispersed short distances from the trial sites during sowing, windrowing or harvest activities, by pod shattering, by seed-hoarding behaviours of animals such as ants or rodents, or by strong winds or runoff after heavy rain. As described in Section 3.1.3, the planting areas would be surrounded by monitoring zones that are inspected while the GMOs are growing, so any volunteers growing from dispersed GM seeds during this period would be detected and destroyed. The applicant also proposes to inspect the monitoring zones after harvest to destroy any volunteers growing from dispersed GM seeds. The size of the monitoring zones is 10 m, 35 m or 50 m, depending on the measures used to control pollen flow. As the short-distance seed dispersal mechanisms listed above are unlikely to transport seeds further than 10 m from the trial sites, the draft licence only requires post-harvest inspections of the innermost 10 m of each monitoring zone.
25. The draft licence includes additional conditions to manage short-distance dispersal of GM seeds. These include taking measures to minimise dispersal of windrowed GMOs by wind or rain, requiring the trial site to be cleaned within 14 days after harvest by a method that removes GM seeds from the soil surface, and requiring post-harvest inspections of any area used to clean equipment or any other area where GMOs are known to have dispersed. This combination of controls would minimise short-distance dispersal of GM seeds leading to establishment of volunteer populations outside the trial sites (risk scenario 2).
    * + 1. Consideration of proposed controls regarding persistence of the GMOs
26. After harvest of each trial site, the applicant proposes to destroy GMOs not required for further evaluation or future trials. This would involve both cleaning the trial site within 14 days after harvest in a manner that destroys any surviving GM plants, and destroying any harvested GM seed that is not required for experimentation or future planting. The applicant’s proposed methods for destruction of GMOs were approved for previous canola field trials (e.g. DIR 164 (OGTR, 2018b)). In addition, uprooting of plants and crushing or grinding of seeds are considered to be effective methods of destruction and have been included as options in the draft licence.
27. To deal with the case of failed crops that are not harvested, draft licence conditions require that GMOs must be harvested or destroyed within eight months after planting, and that if all GMOs in a planting area have been destroyed, then the area is considered to have been harvested and cleaned.
28. The applicant proposes to monitor trial sites after harvest and destroy any volunteers that emerge. The areas that would be monitored are the planting area, the pollen trap, and other areas where GM seed may have dispersed, as discussed in Section 3.1.4. The frequency of inspections of the trial sites are discussed in Section 3.1.3. The proposed duration of monitoring is at least 24 months, and until the site is free of volunteer canola or Indian mustard plants for at least 12 months. This monitoring duration was proposed for previous GM canola field trials and was considered effective for managing persistence of canola seed (e.g. DIR 164 (OGTR, 2018b)).
29. In minimum-tillage Australian farms, the canola seedbank is reported to decline rapidly, and no viable seed was recovered from the seedbank by 2.5 years after canola harvest (Baker and Preston, 2008). Similarly, OGTR monitoring data for nine GM canola trial sites planted in 2015 found that in most sites no canola volunteers emerged more than 1 year after harvest and no volunteers emerged at any site more than 2.5 years after harvest. However, OGTR monitoring data for three GM Indian mustard trial sites found that in two of the sites Indian mustard volunteers continued to emerge for about 5 years after harvest. Although these post-harvest trial sites were maintained in conditions conducive to germination of volunteers, there were three periods of 9-14 months where no volunteers were detected prior to reappearance of Indian mustard volunteers. This data suggests that Indian mustard seeds have greater dormancy in the seedbank than canola seeds. Therefore, for any trial site where Indian mustard is grown, the draft licence requires a monitoring duration of at least 36 months and until the site is free of volunteer plants for at least 18 months. If only canola is grown on a trial site, the required monitoring duration is at least 24 months and until the site is free of volunteer canola plants for at least 12 months.
30. The applicant proposes at least two post-harvest tillages of the trial sites to encourage seed germination. Tillage depth would be no greater than 5 cm, to avoid deep burial of seed that could induce dormancy. The first tillage would occur within 60 days after harvest and the final tillage would occur during the volunteer-free period prior to sign-off. To ensure that the final tillage produces conditions that are conducive to germination of volunteers, the draft licence requires this tillage to be followed by specified levels of rainfall or irrigation that provide sufficient moisture to the seedbank.
31. During the post-harvest monitoring period for each trial site, the applicant proposes to only plant crops permitted on GM brassica trial sites by the Regulator’s [Policy on Post-Harvest Crops](https://www.ogtr.gov.au/resources/publications/policy-post-harvest-crops). This will help to maintain the area in a manner appropriate to allow identification of volunteers.
32. The combination of control measures described in this section would minimise the persistence of GM seeds leading to establishment of GM volunteer populations in the environment (risk scenario 2).
    * + 1. Summary of draft licence conditions to be implemented to limit and control the release
33. A number of licence conditions are proposed to limit and control the release, based on the above considerations. These include requirements to:

limit the duration of the release to the period from November 2022 to December 2027

limit the size of the release to a maximum of twenty sites per year, with ten trial sites of up to 10 ha and ten trial sites of up to 5 ha

limit the location of the release to nominated local government areas in NSW, Victoria and Queensland

not allow GM plant material to be used in human food or animal feed, except for specified animal feeding experiments or taste testing experiments

control pollen flow from the trial sites using one of the following options:

1. surround the planting area with a monitoring zone of 50 m and an isolation zone of a further 950 m
2. surround the planting area with a pollen trap of 15 m, a monitoring zone of 35 m and an isolation zone of a further 350 m
3. cover the planting area with an insect proof tent, and surround the planting area with a monitoring zone of 10 m and an isolation zone of a further 390 m

treat any non-GM canola or Indian mustard grown in planting areas or pollen traps like the GMOs

harvest the GM canola and Indian mustard separately from other crops

clean equipment used with the GMOs before use for any other purpose

transport and store the GMOs in accordance with the Regulator’s guidelines

locate trial sites at least 50 m from any natural waterways

destroy all GMOs not required for further evaluation or future trials

conduct post-harvest monitoring of the planting area and other areas where GM seeds may have been dispersed and destroy any volunteers that emerge

on sites where any Indian mustard has been grown, monitor at least once every 30 days for at least 36 months after harvest and until the site is free of volunteers for at least 18 consecutive months

on sites where only canola has been grown, monitor at least once every 35 days for at least 24 months after harvest and until the site is free of volunteers for at least 12 consecutive months

conduct post-harvest tillage and irrigation of trial sites to encourage seed germination.

3.2 Other risk management considerations

1. All DIR licences issued by the Regulator contain a number of conditions that relate to general risk management. These include conditions relating to:

applicant suitability

contingency plans

identification of the persons or classes of persons covered by the licence

reporting requirements

access for the purpose of monitoring for compliance.

3.2.1 Applicant suitability

1. In making a decision whether or not to issue a licence, the Regulator must have regard to the suitability of the applicant to hold a licence. Under Section 58 of the Act, matters that the Regulator must take into account, for either an individual applicant or a body corporate, include:

any relevant convictions of the applicant

any revocation or suspension of a relevant licence or permit held by the applicant under a law of the Commonwealth, a State or a foreign country

the capacity of the applicant to meet the conditions of the licence.

1. If a licence were issued, the conditions would include a requirement for the licence holder to inform the Regulator of any information that would affect their suitability.
2. In addition, any applicant organisation must have access to a properly constituted Institutional Biosafety Committee and be an accredited organisation under the Act.

3.2.2 Contingency plan

1. If a licence were issued, Nuseed would be required to submit a contingency plan to the Regulator before planting the GMOs. This plan would detail measures to be undertaken in the event of any unintended presence of the GM canola and Indian mustard outside permitted areas.
2. Before planting the GMOs, Nuseed would also be required to provide the Regulator with a method to reliably and uniquely detect the GMOs or the presence of the genetic modifications in a recipient organism.

3.2.3 Identification of the persons or classes of persons covered by the licence

1. If a licence were issued, the persons covered by the licence would be the licence holder and employees, agents or contractors of the licence holder and other persons who are, or have been, engaged or otherwise authorised by the licence holder to undertake any activity in connection with the dealings authorised by the licence. Prior to growing the GMOs, Nuseed would be required to provide a list of people and organisations that will be covered by the licence, or the function or position where names are not known at the time.

3.2.4 Reporting requirements

1. If issued, the licence would require the licence holder to immediately report any of the following to the Regulator:

any additional information regarding risks to the health and safety of people or the environment associated with the trial

any contraventions of the licence by persons covered by the licence

any unintended effects of the trial.

1. A number of written notices would also be required under the licence to assist the Regulator in designing and implementing a monitoring program for all licensed dealings. The notices would include:

expected and actual dates of planting

details of areas planted to the GMOs

expected dates of flowering

expected and actual dates of harvest, method of harvest and dates of cleaning after harvest

details of inspection activities.

3.2.5 Monitoring for compliance

1. The Act stipulates, as a condition of every licence, that a person who is authorised by the licence to deal with a GMO, and who is required to comply with a condition of the licence, must allow inspectors and other persons authorised by the Regulator to enter premises where a dealing is being undertaken for the purpose of monitoring or auditing the dealing. Post-release monitoring continues until the Regulator is satisfied that all the GMOs resulting from the authorised dealings have been removed from the release sites.
2. If monitoring activities identify changes in the risks associated with the authorised dealings, the Regulator may also vary licence conditions, or if necessary, suspend or cancel the licence.
3. In cases of non-compliance with licence conditions, the Regulator may instigate an investigation to determine the nature and extent of non-compliance. The Act provides for criminal sanctions of large fines and/or imprisonment for failing to abide by the legislation, conditions of the licence or directions from the Regulator, especially where significant damage to health and safety of people or the environment could result.
   1. Issues to be addressed for future releases
4. Additional information has been identified that may be required to assess an application for a commercial release of these GM canola and Indian mustard lines, or to justify a reduction in limits and controls. The identified information is additional biochemical characterisation of the GM seeds with respect to levels of potentially toxic ω-3 LC-PUFA oxidation products and how these levels change over time.
   1. Conclusions of the consultation RARMP
5. The RARMP concludes that the proposed limited and controlled release of GM canola and Indian mustard poses negligible risks to the health and safety of people or the environment as a result of gene technology, and that these negligible risks do not require specific risk treatment measures.
6. If a licence were issued, conditions would be imposed to limit the release to the proposed size, location and duration, and to restrict the spread and persistence of the GMOs and their genetic material in the environment, as these were important considerations in establishing the context for assessing the risks.
7. Proposed licence conditions
   1. Interpretations and definitions
8. In this licence:
   * + - 1. unless defined otherwise, words and phrases used in this licence have the same meaning as they do in the Act and the Regulations;
         2. words denoting a gender include any other gender;
         3. words in the singular include the plural and words in the plural include the singular;
         4. words denoting persons include a partnership and a body whether corporate or otherwise;
         5. references to any statute or other legislation (whether primary or subordinate) are a reference to a statute or other legislation of the Commonwealth of Australia as amended or replaced from time to time and equivalent provisions, if any, in corresponding State law, unless the contrary intention appears;
         6. where any word or phrase is given a defined meaning, any other part of speech or other grammatical form in respect of that word has a corresponding meaning;
         7. specific conditions prevail over general conditions to the extent of any inconsistency.
9. In this licence:

**‘Act’** means the *Gene Technology Act 2000* (Commonwealth) or the corresponding State law under which this licence is issued.

**‘Canola’** means plants of the species *Brassica napus* L.

**‘Clean’** means, as the case requires:

* + - * 1. in relation to Equipment or a facility, remove and/or Destroy the GMOs; or
        2. in relation to an area of land specified in this licence as requiring Cleaning:
    1. Destroy canola and Indian mustard plants, if present, to the reasonable satisfaction of the Regulator, and
    2. remove canola and Indian mustard seeds from the soil surface to the reasonable satisfaction of the Regulator.

*Note: The intent of removing seeds from the soil surface is to minimise seed dispersal. One method of removing seeds from the soil surface is Tillage, which moves seeds to under the soil. Tillage must be in accordance with condition 38.*

**‘Contingency Plan’** means a written plan detailing measures to be taken in the event of the unintended presence of the GMOs outside an area that must be inspected. A Contingency Plan must include procedures to:

* + - * 1. ensure the Regulator is notified immediately if the licence holder becomes aware of the event; and
        2. recover and/or Destroy the GMOs to the reasonable satisfaction of the Regulator; and
        3. inspect for and Destroy any Volunteers that may exist as a result of the event to the reasonable satisfaction of the Regulator.

**‘Destroy’** (or **‘Destruction’**) means, as the case requires, kill by one or more of the following methods:

* 1. uprooting;
  2. cutting and shredding/mulching;
  3. Tillage, but only in accordance with condition 38;
  4. treatment with herbicide;
  5. burning/incineration;
  6. autoclaving;
  7. crushing or grinding of seed;
  8. burial, but only in accordance with condition 39;
  9. a method approved in writing by the Regulator.

*Note: ‘As the case requires’ has the effect that, depending on the circumstances, one or more of these techniques may not be appropriate. For example, treatment with herbicide would not successfully kill GM seeds.*

**‘Equipment’** includes, but is not limited to, seeders, harvesters, storage equipment, transport equipment (e.g. bags, containers, trucks), clothing, footwear and tools.

**‘Extreme Weather’** includes, but is not limited to, fires, flooding, cyclones or torrential rain, that could disperse GMOs or affect the licence holder’s ability to comply with licence conditions.

**‘Flowering’** is taken to begin when any plant of the class of plants referred to in a particular condition first has an open flower, and is taken to end when all plants in the class of plants no longer have flowers.

**‘GM’** means genetically modified.

**‘GMOs’** means the genetically modified organisms that are the subject of the dealings authorised by this licence. GMOs include live plants and viable seed.

**‘Indian mustard’** means plants of the species *Brassica juncea* (L.) Czern. & Coss.

**‘Insect-proof’** means sufficient to prevent the entry of insects that commonly pollinate canola and Indian mustard flowers.

**‘Isolation Zone’** means an area of land extending outwards from the outer edge of the Monitoring Zone, as shown in Figure 1.

**‘Logbook’** means a written or electronic record containing information required to be collected and maintained by this licence and which is able to be presented to the Regulator on request.

**‘Monitoring Zone’** means an area of land extending outwards from the outer edge of the Planting Area, or the outer edge of a Pollen Trap if a Pollen Trap is employed, as shown in Figure 1.

**‘OGTR’** means the Office of the Gene Technology Regulator.

**‘Personal Information’** means information or an opinion about an identified individual, or an individual who is reasonably identifiable:

* + - * 1. whether the information or opinion is true or not; and
        2. whether the information or opinion is recorded in a material form or not.

**‘Planting Area’** means an area of land where the GMOs and non-GM canola and Indian mustard are intentionally planted and grown pursuant to this licence, but does not include the Pollen Trap.

**‘Plant Material’** means any part of the GM or non-GM canola and Indian mustard plants grown at a Planting Area or Pollen Trap, whether viable or not, or any product of these plants.

**‘Pollen Trap’** means an area of land extending outwards at least 15 metres from the outer edge of a Planting Area, where only Pollen Trap Plants are grown, as shown in Figure 1.

**‘Pollen Trap Plants’** means non-GM canola grown in a Pollen Trap.

**‘Regulations’** means the Gene Technology Regulations 2001 (Commonwealth) or the corresponding State law under which this licence is issued.

**‘Regulator’** means the Gene Technology Regulator.

**‘Related Species’** means plants of the species *Brassica napus*, *B. rapa*, *B. juncea*, *B. oleracea*, *Hirschfeldia incana*, *Raphanus raphanistrum* or *Sinapis arvensis*, but does not include plants intentionally grown in the Planting Area or Pollen Trap in accordance with licence conditions.

**‘Sign off’** means a notice in writing from the Regulator, in respect of an area, that post-Cleaning obligations no longer apply to that area.

**‘Tillage’** means the use of any technique to disturb the soil.

*Note: Tillage must be in accordance with condition 38.*

**‘Volunteers’** means GM or non-GM canola or Indian mustard plants, which have not been intentionally grown.

950 m Isolation Zone

50 m Monitoring Zone

00

Planting Area

**c**

390 m Isolation Zone

10 m Monitoring Zone

350 m Isolation Zone

15 m Pollen Trap

Planting Area

00

Planting Area covered with Insect-proof tent

35 m Monitoring Zone

**a**

**b**

**Figure 1. Diagrams (not to scale) showing the relationships between Planting Area, Pollen Trap, Monitoring Zone and Isolation Zone.** Site layout (a) with Insect-proof tent, (b) without Insect‑proof tent and with Pollen Trap, and (c) without Insect-proof tent or Pollen Trap. Monitoring and Isolation Zones must be kept free of Related Species.

* 1. General conditions and obligations

1. This licence does not authorise dealings with the GMOs that are otherwise prohibited as a result of the operation of State legislation recognising an area as designated for the purpose of preserving the identity of GM crops, non-GM crops, or both GM crops and non-GM crops, for marketing purposes.
2. This licence remains in force until it is suspended, cancelled or surrendered. No dealings with the GMOs are authorised during any period of suspension.

*Note: Although this licence has no expiry date, the period when GMOs may be grown is restricted in accordance with Condition 18.*

1. The licence holder is Nuseed Pty Ltd.
2. The persons covered by this licence are the licence holder and employees, agents or contractors of the licence holder and other persons who are, or have been, engaged or otherwise authorised by the licence holder to undertake any activity in connection with the dealings authorised by this licence.
3. The GMOs with which dealings are authorised by this licence are those listed at **Attachment** **A**.
4. The dealings authorised by the licence are to:
5. conduct experiments with the GMOs;
6. breed the GMOs;
7. propagate the GMOs;
8. use the GMOs in the course of manufacture of a thing that is not the GMOs;
9. grow the GMOs;
10. import the GMOs;
11. transport the GMOs;
12. dispose of the GMOs;

and the possession, supply or use of the GMOs in the course of any of these dealings.

1. This licence does not apply to dealings with the GMOs conducted as a Notifiable Low Risk Dealing (NLRD) or pursuant to another authorisation under the Act.

*Note: Dealings conducted as an NLRD must be assessed by an Institutional Biosafety Committee (IBC) before commencement and must comply with the requirements of the Regulations.*

##### *General obligations of the licence holder*

1. The licence holder must, at all times, remain an accredited organisation in accordance with the Act and must comply with its instrument of accreditation.
2. The licence holder must be able to access and control all Planting Areas, Pollen Traps, Monitoring Zones, Isolation Zones and approved facilities to the extent necessary to comply with this licence.

*Note: Arrangements to access and control these areas must be notified to the Regulator as part of each planting notification (Condition 47(a)).*

1. The licence holder must inform any person covered by this licence, to whom a particular condition of the licence applies, of the following:
2. the particular condition, including any variations of it;
3. the cancellation or suspension of the licence;
4. the surrender of the licence.
5. The licence holder must not permit a person covered by this licence to conduct any dealing with the GMOs unless:
6. the person has been informed of any applicable licence conditions, including any variation of them; and
7. the licence holder has obtained from the person a signed and dated statement that the person:
8. has been informed by the licence holder of the licence conditions including any variation of them; and
9. has understood and agreed to be bound by the licence conditions, or variation.
10. The licence holder must inform the persons covered by this licence that any Personal Information relevant to the administration and/or enforcement of the licence may be released to the Regulator.

##### *General obligations of persons covered by the licence*

1. If a person is authorised by this licence to deal with the GMOs and a particular condition of the licence applies to the dealing by the person, the person must allow the Regulator, or a person authorised by the Regulator, to enter premises where the dealing is being undertaken, for the purposes of auditing or monitoring the dealing.

*Note: Under the Act, the definition of premises includes a building, area of land or vehicle.*

* 1. Limits and control measures

3.1 Limits on the release

*The following licence conditions impose limits on where and when the GMOs may be grown.*

1. The only plants that may be intentionally grown at a Planting Area are:
2. the GMOs covered by this licence; and
3. non-GM canola and Indian mustard; and
4. plants approved in writing by the Regulator.
5. Non-GM canola and Indian mustard plants grown in a Planting Area must be handled as if they were the GMOs.
6. Planting and growing of the GMOs may only occur within the following limits:

**Area and period**

| **Period** | **Number of Planting Areas per year** | **Maximum size of each Planting Area** |
| --- | --- | --- |
| 5 years (November 2022 to December 2027) | 20 | 5 ha for 10 Planting Areas  10 ha for 10 Planting Areas |

**Local government areas in which Planting Areas may be located**

| **New South Wales** | **Victoria** | **Queensland** |
| --- | --- | --- |
| Albury City Council | Ararat Rural City Council | Goondiwindi Regional Council |
| Balranald Shire Council | Ballarat City Council | Lockyer Valley Regional Council |
| Berrigan Shire Council | Benalla Rural City Council | Somerset Regional Council |
| Bland Shire Council | Buloke Shire Council | Southern Downs Regional Council |
| Blayney Shire Council | Campaspe Shire Council | Toowoomba Regional Council |
| Cabonne Shire Council | Central Goldfields Shire Council | Western Downs Regional Council |
| Carrathool Shire Council | Colac-Otway Shire Council |  |
| Coolamon Shire Council | Corangamite Shire Council |  |
| Coonamble Shire Council | Gannawarra Shire Council |  |
| Cootamundra- Gundagai Regional Council | Glenelg Shire Council |  |
| Cowra Shire Council | Golden Plains Shire Council |  |
| Dubbo Regional Council | Greater Bendigo City Council |  |
| Edward River Council | Greater Geelong City Council |  |
| Federation Council | Greater Shepparton City Council |  |
| Forbes Shire Council | Hepburn Shire Council |  |
| Gilgandra Shire Council | Hindmarsh Shire Council |  |
| Greater Hume Shire Council | Horsham Rural City Council |  |
| Griffith City Council | Indigo Shire Council |  |
| Gunnedah Shire Council | Latrobe City Council |  |
| Gwydir Shire Council | Loddon Shire Council |  |
| Hay Shire Council | Macedon Ranges Shire Council |  |
| Hilltops Council | Melton Shire Council |  |
| Junee Shire Council | Mildura Rural City Council |  |
| Lachlan Shire Council | Mitchell Shire Council |  |
| Leeton Shire Council | Moira Shire Council |  |
| Liverpool Plains Shire Council | Moorabool Shire Council |  |
| Lockhart Shire Council | Mount Alexander Shire Council |  |
| Mid-Western Regional Council | Moyne Shire Council |  |
| Moree Plains Shire Council | Murrindindi Shire Council |  |
| Murray River Council | Northern Grampians Shire Council |  |
| Murrumbidgee Council | Pyrenees Shire Council |  |
| Muswellbrook Shire Council | South Gippsland Shire Council |  |
| Narrabri Shire Council | Southern Grampians Shire Council |  |
| Narrandera Shire Council | Strathbogie Shire Council |  |
| Narromine Shire Council | Surf Coast Shire Council |  |
| Orange City Council | Swan Hill Rural City Council |  |
| Parkes Shire Council | Towong Shire Council |  |
| Snowy Valleys Council | Wangaratta Rural City Council |  |
| Tamworth Regional Council | Warrnambool City Council |  |
| Temora Shire Council | Wellington Shire Council |  |
| Upper Hunter Shire Council | West Wimmera Shire Council |  |
| Wagga Wagga City Council | Wodonga City Council |  |
| Walgett Shire Council | Wyndham City Council |  |
| Warren Shire Council | Yarriambiack Shire Council |  |
| Warrumbungle Shire Council |  |  |
| Weddin Shire Council |  |  |

3.2 Control measures

*The following licence conditions restrict the spread or persistence of the GMOs and their genetic material in the environment.*

##### *Restrictions on GMOs in food or feed*

1. Subject to conditions 20 and 21, Plant Material must not be used, sold or otherwise disposed of for any purpose which would involve or result in its use as food for humans or feed for animals.
2. Non-viable products derived from the GMOs may be fed to rodents, chickens or farmed fish species for experimental purposes, subject to those experiments being approved by an Animal Ethics Committee operating under the Australian Code for the Care and Use of Animals for Scientific Purposes.
3. Oil from the GMOs may be used in taste testing experiments, subject to those experiments being under oversight by a Human Research Ethics Committee, which is required to review and approve the research proposals in accordance with the National Statement on Ethical Conduct in Human Research.

##### *Conditions to restrict pollen flow*

1. For each Planting Area, one of the following measures to limit gene flow must be adopted:
   1. cover all GMOs with Insect-proof tents from at least seven days prior to Flowering and until all GMOs have completed Flowering, and surround the Planting Area with a Monitoring Zone of at least 10 metres, and surround the Monitoring Zone with an Isolation Zone of at least 390 metres (as shown in Figure 1a); or
   2. surround the Planting Area with a Pollen Trap of at least 15 metres, and surround the Pollen Trap with a Monitoring Zone of at least 35 metres, and surround the Monitoring Zone with an Isolation Zone of at least 350 metres (as shown in Figure 1b); or
   3. surround the Planting Area with a Monitoring Zone of at least 50 metres, and surround the Monitoring Zone with an Isolation Zone of at least 950 metres (as shown in Figure 1c).
2. If a Pollen Trap is used in accordance with condition 22, Pollen Trap Plants must:
   1. have a reasonably dense and vigorous growth; and
   2. be Flowering at the same time as the GMOs; and
   3. form a continuous barrier at least 15 metres wide around the Planting Area while the GMOs are Flowering, although one path of up to 3 metres in width is allowed in order to access the Planting Area; and
   4. be handled as if they were the GMOs.
3. The Monitoring Zone must be maintained in a manner appropriate to allow the identification and Destruction of Related Species while the GMOs are growing in the Planting Area.

*Note: Measures to achieve this could include maintaining the area free of vegetation and/or keeping vegetation mown. Condition 48 requires details of current land use and recent land management practices to be recorded upon inspection of the Monitoring Zone.*

1. The GMOs must not be planted in a Planting Area if any Related Species are being grown at the same time in the Monitoring or Isolation Zones.

*Note: Refer to Condition 11 regarding access and control of areas.*

1. While the GMOs are growing in a Planting Area, associated areas and Insect-proof tents must be inspected by people trained to recognise plants of Related Species, and actions must be taken as follows:

| **Area** | **Period of inspection** | **Inspection frequency** | **Inspect for** | **Action** |
| --- | --- | --- | --- | --- |
| Planting Area, Pollen Trap (if applicable) and Monitoring Zone | **From** 14 days prior to the expected commencement of Flowering of any GMOs\*  **until** all GMOs have been harvested or Destroyed | At least once every 35 days | Related Species | Destroy before Flowering or prevent from Flowering |
| Insect-proof tents | While tents are in place | At least once every 14 days and after any Extreme Weather event | Damage that may render tents not Insect-proof | Repair any damage or replace if repair not possible |
| Isolation Zone | **From** 14 days prior to the expected commencement of Flowering of any GMOs\*  **until** all GMOs in the Planting Area have finished Flowering | At least once every 35 days | Intentionally planted Related Species | Destroy before Flowering or prevent from Flowering or Destroy the GMOs in the Planting Area |

*\*Condition 47(a) requires the licence holder to provide information to the Regulator on the expected Flowering period, however the inspection period should be based on the observed development of the GMOs, so that inspections commence prior to Flowering of any GMOs.*

*Note: Details of any inspection activity must be recorded in a Logbook (Condition 48) and reported to the Regulator (Condition 47).*

##### *Conditions to restrict seed dispersal*

1. Equipment used in connection with the GMOs must be Cleaned as soon as practicable after use with the GMOs and before use for any other purpose.
2. Planting Areas and Pollen Traps must be at least 50 metres away from any permanent natural watercourses or man‑made drainage features that flow into natural watercourses.

*Note: This includes irrigation channels or storm water drains that flow into a natural watercourse.*

1. Planting Areas and Pollen Traps must not be located in flood prone areas.
2. If the GMOs are windrowed, the licence holder must take, or have taken, measures to minimise the likelihood of dispersal of the GMOs by wind or rain. Appropriate measures may include:
   1. ensuring high density planting and growth of the GMOs prior to windrowing; or
   2. cutting/windrowing to allow maximum stubble height; or
   3. use of windrow roller; or
   4. appropriate site selection.

*Note: Appropriate site selection includes avoidance of windy areas. Windrowing dates and details of measures used to minimise dispersal of GMOs must be reported to the Regulator (Condition 47(d)).*

##### *Conditions relating to harvesting*

1. GMOs must be harvested or Destroyed within eight months after planting.
2. If all GMOs in a PlantingArea have been Destroyed, then for the purposes of this licence:
   1. the GMOs are taken to have been harvested; and
   2. the Planting Area is taken to have been Cleaned.

*Note: Cleaning activities must be reported to the Regulator (Condition 47). Areas of land that have been Cleaned are subject to inspections (Condition 36).*

1. The GMOs must be harvested and threshed separately from any other crop.
2. Harvested GM seed not required for experimentation or future planting must be Destroyed as soon as practicable.

##### *Conditions to restrict persistence of GMOs on trial sites*

1. Areas of land used in connection with the GMOs must be Cleaned as follows:

| **Areas of land to be Cleaned** | **When** |
| --- | --- |
| * + 1. Planting Area     2. Pollen Trap, if used     3. 10 metres around each Planting Area, or around the Pollen Trap, if used (innermost 10 metres of Monitoring Zone) | Within 14 days after harvest of the GMOs |
| Any other area used to Clean any Equipment used in connection with the GMOs | As soon as practicable |
| Any other area where the GMOs have dispersed, e.g. during planting, growing, harvesting or Destruction | As soon as practicable |

*Note: Cleaning activities must be reported to the Regulator (Condition 47). Areas of land that have been Cleaned are subject to inspections (Condition 36).*

1. After Cleaning, areas of land must be inspected by people trained to recognise canola and Indian mustard. Inspections must cover the entirety of areas to be inspected. Actions must be taken as follows:

| **Area** | **Period of inspection** | **Inspection frequency** | **Inspect for** | **Action** |
| --- | --- | --- | --- | --- |
| Planting Area, Pollen Trap, innermost 10 metres of Monitoring Zone and other areas of land that were Cleaned in accordance with Condition 35 | From the day of Cleaning, until:   * + 1. the area is planted as a new Planting Area in accordance with condition 16; or     2. the Regulator has issued a Sign off for the area | * + 1. At least once every 30 days if any Indian mustard was grown on the Planting Area; or     2. at least once every 35 days if only canola was grown on the Planting Area | Volunteers | Destroy before Flowering |

*Note: Details of any inspection activity must be recorded in a Logbook (Condition 48) and reported to the Regulator (Condition 47).*

1. While post-Cleaning inspection requirements apply to an area:
   1. the area must be Tilled within 60 days of harvest of the GMOs at a Planting Area, unless otherwise approved in writing by the Regulator; and

*Note: If Tillage is used as a method of Cleaning, the Tillage done as Cleaning also meets the requirements for a Tillage within 60 days of harvest.*

* 1. within the 12 months prior to submission of a Sign off application, the area must be Tilled and then receive a watering event as described in **Attachment B**; and
  2. the area must be maintained in a manner appropriate to allow identification of Volunteers; and
  3. the area must not be used for grazing livestock; and
  4. no plants may be intentionally grown in the area unless:
     1. the area is planted as a new Planting Area in accordance with condition 16; or
     2. the plants are listed as post-harvest crops permitted for GM Brassica field trial sites in the OGTR Policy on Post Harvest Crops as current at the time of planting; or
     3. the plants are agreed to in writing by the Regulator.

*Note: The OGTR’s Policy on Post Harvest Crops can be found on the* [*OGTR website*.](https://www.ogtr.gov.au/resources/publications/policy-post-harvest-crops)

##### *Tillage*

1. Any Tillage of the Planting Area and the Pollen Trap must be to a depth no greater than five centimetres.

##### *Destruction by burial*

1. If Destruction of GMOs occurs by burial:
   1. the GMOs must be buried in a pit and covered by a layer of soil at least one metre in depth, the top of which is no higher than the surrounding soil surface; and
   2. seeds must be wet when buried to encourage decomposition; and
   3. the licence holder must take measures to ensure that the burial site is not disturbed for a period of at least two years from the date of burial.

*Note: If GMOs are dispersed on the soil surface during the process of burial, the burial site becomes an area of land that requires Cleaning under Condition 35, and is subject to post-Cleaning requirements.*

*Note: The date and location of burial, and measures used to ensure that the burial site is not disturbed, must be reported to the Regulator (Condition 47g).*

##### *Processing or experimentation with the GMOs*

1. Treatment, threshing or processing of GM seed or experimentation or analysis with the GMOs may only be undertaken within:
   1. a Planting Area before Cleaning; or
   2. a Pollen Trap before Cleaning; or
   3. the innermost 10 m of a Monitoring Zone before Cleaning; or
   4. a facility approved in writing by the Regulator.

*Note: This condition does not apply to dealings conducted as an NLRD (see Condition 9).*

1. Within a facility approved in writing by the Regulator in accordance with Condition 40, any area that is used for treatment, threshing, processing, experimentation or analysis of the GMOs must be Cleaned as soon as practicable and before use for any other purpose.

##### *Transport or storage of the GMOs*

1. Transport or storage of the GMOs must:
   1. only occur to the extent necessary to conduct the dealings permitted by this licence or other valid authorisation under the Act, or to the extent necessary to enable export of the GMOs; and
   2. be in accordance with the Regulator’s *Guidelines for the Transport, Storage and Disposal of GMOs* for PC2 GM plants as current at the time of transportation or storage; and
   3. comply with all other conditions of this licence.

*Note: Activities with the GMOs within a Planting Area prior to Cleaning are not regarded as transport or storage.*

*Note: Condition 13 requires signed statements for persons transporting the GMOs.*

*Note: This condition does not apply to dealings conducted as an NLRD (see Condition 9).*

1. Methods and procedures used to transport GMOs must be recorded, and must be provided to the Regulator, if requested.

*Note: The Contingency Plan must be implemented if the GMOs are detected outside areas under inspection (Condition 44).*

##### *Contingency plan*

1. If any unintentional presence of the GMOs is detected outside the areas requiring Cleaning, the Contingency Plan must be implemented.
   1. Sign off
2. The licence holder may make written application to the Regulator that planting restrictions and inspection requirements no longer apply to the Planting Area and other areas requiring Cleaning if:
   1. post-Cleaning inspection activities have been conducted on the area for at least 36 months if any Indian mustard was grown on the Planting Area, or at least 24 months if only canola was grown on the Planting Area; and
   2. conditions have been conducive for germination and detection of Volunteers; and
   3. prior to the Sign-off request, no Volunteers have been detected in the area for at least 18 months if any Indian mustard was grown on the Planting Area, or at least 12 months if only canola was grown on the Planting Area.

*Note: The licence requires two Tillages and a watering event prior to a Sign off application (Condition 37).*

*Note: The Regulator will take into account the management and inspection history for the Planting Area and other areas requiring Cleaning, including post-harvest crops planted (if any), Tillage, irrigation, rainfall, application of herbicide and occurrence of Volunteers, in deciding whether or not further inspections are required to manage persistence of the GMOs.*

* 1. Reporting and documentation

*The following licence conditions are imposed to demonstrate compliance with other conditions and facilitate monitoring of compliance by staff of the OGTR.*

1. General notifications must be sent to the Regulator as follows:

*Note: please send all correspondence related to the licence to* [*OGTR.M&C@health.gov.au*](mailto:OGTR.M&C@health.gov.au)*.*

|  |  |  |
| --- | --- | --- |
| **Notice** | **Content of notice** | **Timeframe** |
| 1. Changes to contact details | Changes to any of the contact details of the project supervisor that were notified in the licence application or subsequently | As soon as practicable |
| 1. Ongoing suitability to hold a licence | * + 1. any relevant conviction of the licence holder; or     2. any revocation or suspension of a licence or permit held by the licence holder under a law of the Australian Government, a State or a foreign country, being a law relating to the health and safety of people or the environment; or     3. any event or circumstances that would affect the capacity of the licence holder to meet the conditions of the licence; and | As soon as practicable after any of these events occur |
| * + 1. any information related to the licence holder's ongoing suitability to hold a licence, that is requested by the Regulator | Within the timeframe stipulated by the Regulator |
| 1. People covered by the licence | * + 1. names of all organisations and persons, or functions or positions of the persons, who will be covered by the licence, with a description of their responsibilities; and   *Note: Examples of functions or positions are ‘project supervisor’, ‘site manager’, ‘farm labourer’ etc*.   * + 1. detail of how the persons covered by the licence will be informed of licence conditions | At least 14 days prior to conducting any dealings with the GMOs (to be updated within 14 days if the notified details change) |
| 1. Testing methodology | A written methodology to reliably detect the genetic modifications described in this licence. The detection method/s must be capable of identifying each GM canola and Indian mustard line planted under this licence | At least 14 days prior to conducting any dealings with the GMOs (to be updated within 14 days if the notified details change) |
| 1. Contingency plan | A Contingency Plan to respond to inadvertent presence of the GMOs outside an area that must be inspected | At least 14 days prior to conducting any dealings with the GMOs (to be updated within 14 days if the notified details change) |
| 1. Training records | Copies of the signed and dated statements referred to in condition 13 if requested by the Regulator | Within the timeframe stipulated by the Regulator |
| 1. Additional information required by the Act | 1. additional information as to any risks to the health and safety of people, or to the environment, associated with the dealings authorised by the licence; or 2. any contraventions of the licence by a person covered by the licence; or 3. any unintended effects of the dealings authorised by the licence   *Note: The Act requires, for the purposes of the condition 46.g, that:*   * *the licence holder will be taken to have become aware of additional information of a kind mentioned in Condition 46.g if he or she was reckless as to whether such information existed; and* * *the licence holder will be taken to have become aware of contraventions, or unintended effects, of a kind mentioned in Condition 46.g, if he or she was reckless as to whether such contraventions had occurred, or such unintended effects existed*   *Note: Contraventions of the licence may occur through the action or inaction of a person.* | Without delay after becoming aware of any new information  *Note: An example of notification without delay is contact made within a day of a contravention of the licence via the OGTR free call phone number 1800 181 030, which provides emergency numbers for incidents that occur out of business hours. Notification without delay will allow the OGTR to conduct a risk assessment on the incident and attend the location, if required* |
| 1. Further details regarding additional information | Any further details requested by the Regulator in relation to information provided under condition 46.g | Within the timeframe stipulated by the Regulator |

1. Notifications relating to each trial site must be sent to the Regulator as follows:

*Note: please send all correspondence related to the licence to* [*OGTR.M&C@health.gov.au*](mailto:OGTR.M&C@health.gov.au)*.*

| **Notice** | **Content of notice** | **Timeframe** |
| --- | --- | --- |
| * 1. Intention to plant | * + 1. Details of the Planting Area including size, the local government area, GPS coordinates, a street address, a diagrammatical representation of the trial site (e.g. Google Maps) and any other descriptions     2. Whether an Insect-proof tent or Pollen Trap will be used     3. Details of how the licence holder will access and control the Planting Area and the associated Pollen Trap, Monitoring Zone and Isolation Zone, in accordance with condition 11   *Note: this should include a description of any contracts, agreements, or other enforceable arrangements.*   * + 1. Whether any Indian mustard will be planted in the Planting Area     2. Date on which the GMOs will be planted     3. Period when the GMOs are expected to Flower     4. Period when windrowing (if intended) is expected to commence     5. Period when harvesting is expected to commence     6. How all areas requiring post-Cleaning inspections are intended to be used until Sign off, including proposed post-harvest crops (if any)     7. Details of how inspection activities will be managed, including strategies for the detection and Destruction of Volunteers     8. History of how the trial site has been used for the previous two years | At least 7 days prior to each planting (to be updated as soon as practicable if the notified details change) |
| * 1. Planting | * + 1. Actual date(s) of planting the GMOs     2. Any changes to the details provided under part (a) of this condition | Within 7 days of any planting |
| * 1. Extreme Weather | Any Extreme Weather event that is expected to affect or has already affected an area where the GMOs are or may be present.  *Note: The Contingency Plan must be implemented if the GMOs are detected outside areas requiring Cleaning (Condition 44).* | As soon as practicable |
| * 1. Windrowing | Actual date(s) of windrowing and details of measures used to minimise dispersal of the GMOs during windrowing (Condition 30). | Within 7 days of commencement of windrowing |
| * 1. Harvest | Actual date(s) of harvesting the GMOs | Within 7 days of commencement of any harvesting |
| * 1. Cleaning | * + 1. Date(s) on which required Cleaning was performed on any areas of land     2. Method(s) of Cleaning | Within 7 days of completion of Cleaning |
| * 1. Destruction by burial | Date of burial, location of burial including GPS co‑ordinates, and details of measures used to ensure that the burial site will not be disturbed for the period required by Condition 39 | Within 7 days of burial of any GMOs |
| * 1. Inspection activities | Information recorded in a Logbook as per the inspection requirements (Conditions 26, 36 and 48). | Within 35 days of inspection |

*Note: Additional records must be provided to the Regulator, if requested, in accordance with condition 43.*

1. Details of any inspection activity must be recorded in a Logbook and must include:
   1. date of the inspections; and
   2. name of the person(s) conducting the inspections; and
   3. details of the experience, training or qualification that enables the person(s) to recognise canola, Indian mustard and/or Related Species, if not already recorded in the Logbook; and
   4. details of areas inspected including current land use (including any post-harvest crops) and recent management practices applied; and

*Note: management practices include Tillage events, spraying or maintenance measures used to facilitate inspections.*

* 1. details of the developmental stage of the GMOs while they are being grown; and
  2. details of any post-Cleaning rainfall events including measurements at or near the area, or any irrigation events; and
  3. details of any Volunteers and/or Related Species observed during inspections or during land-management activities, including number, developmental stage and approximate position of the Volunteers and/or Related Species within each area inspected†; and
  4. date(s) and method(s) of Destruction of or preventing Flowering of any Volunteers and/or Related Species, including destruction of Volunteers and/or Related Species during land-management activities; and
  5. details of any damage and any repairs to the Insect-proof tents, while Insect-proof tents are required.

*† Examples of acceptable ways to record the positional information for Volunteers and/or Related Species in the Logbook include:*

*- descriptive text*

*- marking on a diagram*

*- indicating grid references on a corresponding map/sketch.*

*Note: Details of inspection activities must be provided to the Regulator (Condition 47). The Regulator has developed a standardised proforma for recording inspection activities. This can be made available on request.*

## ATTACHMENT A

**DIR No: 188**

**Full Title:** Limited and controlled release of canola and Indian mustard genetically modified for altered oil content and herbicide tolerance

**Organisation Details**

Postal address: Nuseed Pty Ltd

103-105 Pipe Road

Laverton North, VIC 3026

Phone No:(03) 9282 1000

**IBC Details**

IBC Name: Nuseed Institutional Biosafety Committee

**GMO Description**

**GMOs covered by this licence**

Canola and Indian mustard plants genetically modified by introduction of only the genes and genetic elements listed below.

**Parent Organisms**

Common Names: Canola and Indian mustard

Scientific Names: *Brassica napus* L. and *Brassica juncea* (L.) Czern. & Coss.

**Modified traits**

Category: Composition – food (human nutrition)

Composition – animal nutrition

Herbicide tolerance

Description: Canola and Indian mustard plants have been genetically modified by introduction of up to seven genes involved in fatty acid biosynthesis. The GMOs are intended to produce ω-3 long-chain polyunsaturated fatty acid in seed oil. The GMOs may also contain a selectable marker gene that confers herbicide tolerance. The introduced genes are listed in Table 1 and the associated regulatory sequences are listed in Table 2.

**Purpose of the dealings with the GMO**

The purpose of the release is to evaluate the altered oil content trait under field conditions. The GM canola and Indian mustard are not permitted to be used for human food or animal feed except in specified animal feeding experiments and taste testing experiments.

**Table 1**. Introduced genes in the GM canola and Indian mustard

|  |  |  |
| --- | --- | --- |
| **Gene** | **Source organism** | **Description of encoded protein** |
| *Lackl-Δ12D* | *Lachancea kluyveri* | Fatty acid Δ12-desaturase |
| *Picpa-ω3D* | *Pichia pastoris* | Fatty acid ω-3 desaturase |
| *Micpu-Δ6D* | *Micromonas pusilla* | Fatty acid Δ6-desaturase |
| *Pyrco-Δ6E* | *Pyramimonas cordata* | Fatty acid Δ6-elongase |
| *Pavsa-Δ5D* | *Pavlova salina* | Fatty acid Δ5-desaturase |
| *Pyrco-Δ5E* | *Pyramimonas cordata* | Fatty acid Δ5-elongase |
| *Pavsa-Δ4D* | *Pavlova salina* | Fatty acid Δ4-desaturase |
| *pat* | *Streptomyces viridochromogenes* | Enzyme for glufosinate herbicide tolerance |

**Table 2**. Introduced regulatory sequences in the GM canola and Indian mustard

|  |  |  |
| --- | --- | --- |
| **Sequence** | **Source** | **Intended function** |
| PRO\_Arath-FAE1 | Promoter of *Arabidopsis thaliana* fatty acid elongase 1 | Seed specific promoter |
| PRO\_Brana-FP1 | Promoter of *Brassica napus* napin | Seed specific promoter |
| PRO\_Linus-Cnl1 | Promoter of *Linum usitatissimum* conlinin1 | Seed specific promoter |
| PRO\_Linus-Cnl2 | Promoter of *Linum usitatissimum* conlinin2 | Seed specific promoter |
| PRO\_35S×2 | Promoter of Cauliflower mosaic virus 35S RNA | Constitutive promoter |
| Tobacco mosaic virus 5' UTR leader | Enhancer from Tobacco mosaic virus 59 | Increase gene expression |
| MAR\_Nicta- RB7 | Rb7 matrix attachment region from *Nicotiana tabacum* | Increase gene expression |
| TER\_Agrtu-NOS | Terminator of *Agrobacterium tumefaciens* nopaline synthase | Terminator |
| TER\_Glyma-Lectin | Terminator of *Glycine max* lectin *Le1* | Terminator |
| TER\_Linus-Cnl1 | Terminator of *Linum usitatissimum* conlinin1 | Terminator |
| TER\_Linus-Cnl2 | Terminator of *Linum usitatissimum* conlinin2 | Terminator |

## ATTACHMENT B

A watering event is irrigation or natural rainfall that provides sufficient soil moisture to promote germination of canola and Indian mustard seeds on a trial site.

Examples of acceptable watering events are:

* At least 26 millimetres of rainfall over one day; or
* At least 28 millimetres of rainfall over two days; or
* At least 30 millimetres of rainfall over three days; or
* At least 32 millimetres of rainfall over four days; or
* Irrigation that provides equivalent levels of soil moisture to one of the examples of rainfall above.

Rainfall measurements must be taken on the site or within 3 km of the site. An irrigation or natural rainfall that matches one of the examples listed above, and occurs during the time period specified for a watering event in Condition 37 of the licence, is considered a valid watering event. The licence holder should keep records of the date/s and amount of water applied during the watering event, and provide this information when requesting Sign off of the relevant site.

If an irrigation or natural rainfall does not match one of the examples listed above, the licence holder may submit a request to the Regulator for it to be considered a watering event. The request should provide:

* evidence of amount of water applied, such as rainfall measurements on the site or within 3 km of the site, and
* evidence that resultant soil moisture is suitable for germination, such as photos of germinating plants on the site.

It is recommended that any requests that an irrigation or natural rainfall be considered a watering event be submitted at the time of the event, to minimise potential delays to Sign off of the site.

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1. As none of the proposed dealings are considered to pose a significant risk to people or the environment, Section 52(2)(d)(ii) of the Act mandates a minimum period of 30 days for consultation on the RARMP. [↑](#footnote-ref-1)