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**Supporting document 1 (at Approval)**

Risk and technical assessment report – Application A1186

Soy leghemoglobin in meat analogue products

# Executive summary

A genetically modified (GM) strain of the yeast *Pichia pastoris* has been developed for the production of soy leghemoglobin. Leghemoglobin is a globin protein, containing an iron-bound haem B prosthetic group and is typically expressed in the root nodules of leguminous plants.

The applicant’s Soy leghemoglobin, in a liquid preparation branded ‘LegH Prep’[[1]](#footnote-2), will be added to meat analogue products at levels of not more than 0.8% (w/w, in raw product) to replicate the nutritional source of iron, flavour and aroma of myoglobin, a protein found in the muscle tissue of animals. Soy leghemoglobin does have additional functions that align with the definition of a food additive (i.e. flavouring and aroma). FSANZ considered permissons for soy leghemoglobin in three separate Code standards/schedules (as a GM food, as a food additive, and as a nutritive substance) but decided that consideration as a food additive was not justified. Regulation as a nutritive substance allowed FSANZ to include permissions for it as a source of iron in Schedule S17—3, and review data and information on the bioavailability of iron in the ingredient.

The production of soy leghemoglobin is achieved through the expression of a leghemoglobin gene from soybean (*Glycine max*) as well as overexpression of several genes from the *Pichia* host. The overexpression of host genes support the production of the leghemoglobin protein. The soy leghemoglobin preparation (the Preparation) produced by the production strain MXY0541, is a soluble component of a fermentation culture that has undergone mechanical lysis, heat inactivation and partial purification. The Preparation will contain novel soy leghemoglobin and also proteins and genetic material from the *Pichia* host.

In conducting the risk assessment of the soy leghemoglobin and the Preparation, a number of criteria have been addressed, including the safety of the *P. pastoris* host strain, novel proteins, toxicity of the Preparation and a nutritional and dietary intake assessment of soy leghemoglobin. The safety assessment of the source organism and novel proteins concluded there were no public health and safety concerns. The source organism is a well characterised yeast with a recognised safe history of use for the production of food enzymes. It is neither pathogenic nor toxigenic.

The novel soy leghemoglobin was shown to be equivalent to that expressed in soybean and was shown to be expressed as a holoprotein[[2]](#footnote-3). Analyses of the potential allergenicity or toxicity of the soy leghemoglobin and the *Pichia* proteins did not identify any significant similarities to known allergens or toxins. The proteins were shown to be susceptible to pepsin digestion and were denatured at standard cooking temperatures and in acidic conditions that mimic the stomach environment. The shelf life and specifications of the Preparation are also appropriate for addition to meat analogue products.

The applicant submitted *in vitro* genotoxicity studies in bacterial and mammalian cells and an oral toxicity study in rats. These studies are intended to confirm the outcome of the compositional and bioinformatic analysis conducted as a part of the safety assessment. No hazard was identified in the submitted studies. The Preparation was not genotoxic *in vitro* and did not cause adverse effects in short-term toxicity studies in rats. The no observed adverse effect level (NOAEL) of freeze-dried Preparation in a 28-day dietary toxicity study in rats was 1536 mg/kg bw/day, the highest dose tested. This dose corresponds to 1421 mg/kg bw/day total organic solids (TOS).

Mean and P90 estimated dietary intakes of the Preparation at the maximum proposed use level were 20 – 60 mg/kg bw/day TOS and 45 – 124 mg/kg bw/day TOS, respectively. Mean and P90 estimated dietary intakes of the Preparation at the likely use level were 11 – 32 mg/kg bw/day TOS and 24 – 68 mg/kg bw/day TOS, respectively. The estimated intakes of the Preparation for both scenarios are considered to be conservative and over-estimate the intake as it is unlikely that consumers will eat meat analogue products containing soy leghemoglobin in the same amounts or with the same frequency they currently consume minced meat and poultry products, and vegetarian meat alternatives (particularly over a long period of time).

The margins of exposure (MOEs) between the NOAEL of 1421 mg/kg bw/day TOS in the rat oral toxicity study and estimated dietary intakes at the maximum proposed use level ranged between 20 – 70 for mean intakes and between 10 – 30 at the 90th percentile. At likely use levels, MOEs for mean and P90 estimated dietary intakes ranged between 40 – 130 and 20 – 60, respectively. These MOEs are not considered to be of concern given that: a sufficient body of knowledge exists on the safety of the source organism, the soy leghemoglobin and *Pichia* proteins will be digested like other other dietary proteins and do not share any significant similarities to known allergens or toxins; and the conservative nature of the dietary intake assessment which is likely to overestimate intakes over a long period of time.

The nutrition assessment concluded that haem iron from soy leghemoglobin is expected to have similar bioavailability to haem iron from mammalian haem proteins (e.g. myoglobin present in muscle tissue) based on the available evidence. Soy leghemoglobin has similar structural and physicochemical properties to animal myoglobins, and soy leghemoglobin is completely digested by pepsin thus making the haem group freely available for absorption. However, in the absence of *in vivo* studies, a quantitative comparison of haem iron bioavailability from soy legehemoglobin and other haem proteins is not possible. The absence of meat proteins in the proposed meat analogue products may decrease the bioavailability of haem iron from soy leghemoglobin. As iron absorption is regulated tightly by the body, and meat analogue products have higher total iron content due to higher content of non-haem iron relative to comparison meat products, any decrease in haem iron bioavailability should not result in a nutritional disadvantage to consumers in Australia and New Zealand.

The estimated intakes of iron (with the additional iron contribution from soy leghemoglobin) for all population age/sex groups assessed for both the Australian and New Zealand populations are below the ULs for iron. The estimated iron intakes in FSANZ’s assessment, for both the *maximum proposed use level* and *likely use level* scenarios for soy leghemoglobin, are considered to be conservative and result in an overestimation of actual iron intakes. It is unlikely that consumption of meat analogue products containing soy leghemoglobin would pose a risk of iron exceedances to the Australian and New Zealand populations, including at maximum use levels up to 0.8%.

As of March 2020, the applicant advised they had sold approximately 100,000,000 quarter-pound (113 g) servings of meat analogue products containing the Preparation. Its post-marketing surveillance has identified one complaint per 600,000 servings based on the current formulation (released on the market in the US in early 2019), but none of these complaints has been confirmed as an adverse event due to consumption of these products.

The assessment of soy leghemoglobin and the soy leghemoglobin preparation concluded there are no public health and safety concerns associated with its use in meat analogue products at the proposed levels.

Table of contents

[Executive summary i](#_Toc58847769)

[Index of Figures 2](#_Toc58847770)

[Index of Tables 2](#_Toc58847771)

[Abbreviations 2](#_Toc58847772)

[1 Introduction 4](#_Toc58847773)

[2 Risk assessment of soy leghemoglobin and the soy leghemoglobin preparation 5](#_Toc58847774)

[2.1 Technical assessment 5](#_Toc58847775)

[2.2 History of use 7](#_Toc58847776)

[2.3 Characterisation of the novel proteins in the soy leghemoglobin preparation 9](#_Toc58847777)

[2.4 Toxicological assessment of soy leghemoglobin preparation 13](#_Toc58847778)

[2.5 Nutritional impact 18](#_Toc58847779)

[2.6 Dietary intake assessment 21](#_Toc58847780)

[2.7 Manufacturing process 29](#_Toc58847781)

[3 Risk assessment summary and conclusion 34](#_Toc58847782)

[4 References 35](#_Toc58847783)

[Appendix 1: FSANZ Approach to the dietary intake assessment 42](#_Toc58847784)

[A1.1 Food consumption data used 42](#_Toc58847785)

[A1.2 Concentration data used 46](#_Toc58847786)

[A1.3 Assumptions and limitations of the dietary intake assessment 46](#_Toc58847787)

[References 47](#_Toc58847788)

[Appendix 2: Estimated dietary intake of iron with additional contribution from soy leghemoglobin 48](#_Toc58847789)

[References 51](#_Toc58847790)

# Index of Figures

|  | Title | Page |
| --- | --- | --- |
| Figure 1 | Lineage of the *Pichia pastoris* host organism Bg11 | 4 |
| Figure 2 | Tryptic peptide map of the leghaemoglobin in LegH | 6 |
| Figure 3 | Schematic Overview of the Manufacturing Process for the soy leghemoglobin preparation | 26 |
| Figure 4 | The DAS2 expression cassette | 30 |

# Index of Tables

|  | Title | Page |
| --- | --- | --- |
| Table 1 | Specifications and batch analyses for the the soy leghemoglobin preparation | 2 |
| Table 2 | Haematology values for which statistically significant changes were observed | 12 |
| Table 3 | Clinical chemistry values for which statistically significant changes were observed | 12 |
| Table 4 | Population groups used in the dietary intake assessment for the soy leghemoglobin preparation | 19 |
| Table 5 | Population groups used in the dietary intake assessment for iron | 19 |
| Table 6 | Estimated dietary intake of the soy leghemoglobin preparation for Australia and New Zealand | 21 |
| Table 7 | Amount of iron in meat analogue products containing soy leghemoglobin compared to minced meat and poultry products and other vegetarian meat alternatives | 22 |
| Table 8 | Estimated dietary intake of iron (mg/day) for the Australian population with additional contribution from soy leghemoglobin | 23 |
| Table 9 | Estimated dietary intake of iron (mg/day) for the New Zealand population with additional contribution from soy leghemoglobin | 24 |
| Table A1.1 | Food groups used to derive consumption amounts for meat analogue products containing soy leghemoglobin | 42 |
| Table A1.2 | Mean and 90th percentile consumption amounts for minced meat and poultry products and vegetarian meat alternatives for the Australian and New Zealand populations | 43 |
| Table A1.3 | Mean and 90th percentile consumption amounts of minced meat and poultry products and vegetarian meat alternatives by age group and sex | 43 |
| Table A2.1 | Current mean iron intake of meat eaters and non-meat eaters | 46 |
| Table A2.2 | Estimated dietary intake of iron (mg/day) for the Australian population with additional contribution from soy leghemoglobin | 47 |
| Table A2.3 | Estimated intake of iron (mg/day) for the New Zealand population with additional contribution from soy leghemoglobin | 48 |

# Abbreviations

|  |  |
| --- | --- |
| AOAC | Association of Official Agricultural Chemists |
| AOX1 | alcohol oxidase 1 gene |
| AOX2 | alcohol oxidase 2 gene |
| bw | body weight |
| cGMP | current good manufacturing practice |
| CSIRO | Commonwealth Scientific and Industrial Research Organisation |
| Da | dalton |
| DAS2 | dihydroxyacetone synthase 2 gene |
| DNA | deoxyribonucleic acid |
| EFSA | European Food Safety Authority |
| EHEC | enterohemorrhagic *Escherichia coli* |
| EPA | Environmental Protection Agency (USA) |
| FDA | Food and Drug Administration (USA) |
| FSANZ | Food Standards Australia New Zealand |
| g | gram |
| GLP | good laboratory practice |
| GM | genetically modified |
| GRAS | generally recognised as safe (US FDA) |
| GRN | GRAS notification number (US FDA) |
| kg | kilogram |
| L | litre |
| LOD | limit of detection |
| mg | milligram |
| m | minute |
| mL | millilitre |
| mmol | millimolar |
| MOE | margin of exposure |
| NHMRC | National Health and Medical Research Council |
| NOAEL | no observed adverse effect level |
| NRV | nutrient reference value |
| NRRL | Northern Regional Research Laboratory (US Department of Agriculture) |
| N-terminal | amino-terminal of a protein |
| OECD | Organisation for Economic Co-operation and Development |
| PAGE | polyacrylamide gel electrophoresis |
| PCR | polymerase chain reaction |
| ppm | parts per million |
| QPS | qualified presumption of safety |
| SDS | sodium dodecyl sulfate |
| Spp. | species |
| TOS | total organic solids |
| U | units |
| μg | microgram |
| μL | microliter |
| UL | upper level of intake |
| UPLC | Ultra performance liquid chromatography |
| USA | United States of America |
| UTR | untranslated region |
| UV | ultraviolet |
| w/w | weight for weight |

# 1 Introduction

FSANZ received an application from Impossible Foods Inc. to permit the use of soy leghemoglobin in meat analogue products. Soy leghemoglobin is a component of a cell lysate preparation from a genetically modified (GM) yeast, *Pichia pastoris*. The yeast has been modified to express the leghemoglobin gene from soybean *(Glycine max*) and other host proteins that support the expression of leghemoglobin. Leghemoglobin is a globin protein, containing an iron-bound haem B prosthetic group and is typically expressed in the root nodules of leguminous plants. The soy leghemoglobin preparation (the Preparation) [[3]](#footnote-4) also contains proteins and genomic DNA from the *Pichia* production strain.

The purpose of adding soy leghemoglobin to meat analogue products is to replicate the nutrition (source of iron), flavour and aroma of myoglobin, an oxygen transporting haem protein found in the muscle tissue of animals (Ordway and Garry 2004). Though soy leghemoglobin does have additional functions that align with the definition of a food additive (i.e. flavouring and aroma), FSANZ considered three permissions in the Code for the same ingredient was excessive. Regulation as a nutritive substance allowed FSANZ to include permissions for it as a source of Iron in S17—3, and review data and information on the bioavailability of iron in the ingredient.

To prepare the Preparation for use in meat analogue products, *Pichia* cells expressing soy leghemoglobin undergo a fermentation process followed by mechanical lysis, with further processing to minimise the level of impurities and to ensure absence of viable *Pichia* cells. The partially purified preparation is then blended with stabilisers. This preparation contains ≤ 14% total protein, of which ≥ 65% is leghemoglobin, with the remainder of the protein consisting of native *Pichia* proteins.

The Preparation will not be sold in Australia or New Zealand as an individual product, rather it will be an ingredient in meat analogue products manufactured by the applicant at levels of not more than 0.8% soy leghemoglobin (w/w). The applicant currently manufactures both the Preparation and their meat analogue products outside of Australia and New Zealand, therefore these products will be imported as frozen, packaged products. In the future, the applicant may establish a co-manufacturing agreement in Australia and New Zealand to produce Impossible meat analogue products. Where possible, locally sourced ingredients would be used with the exception of the Preparation, which would continue to be manufactured in an Impossible Foods production facility located outside of Australia and New Zealand. Its use would be permitted in Impossible meat analogue products in the co-manufacturers’ foods for sale such as ready-meals and burgers etc.

As of March 2020, the applicant advised they had sold approximately 100,000,000 quarter-pound 113g servings. Meat analogue products containing the Preparation are permitted for sale in retail and catering outlets in the US, Canada, Singapore, Hong Kong and Macau.

# 2 Risk assessment of soy leghemoglobin and the soy leghemoglobin preparation

## 2.1 Technical assessment

### 2.1.1 Identity of the source organism and product

The Preparation containing the soy leghemoglobin protein is produced from a GM *P. pastoris* strain MXY0541. This strain contains several copies of the leghemoglobin gene from soybean (*G. max*) as well as extra copies of endogenous genes to overexpress a transcription factor Mxr1 and the eight enzymes involved in the synthesis of the prosthetic group haem B. A full characterisation of the production strain MXY0541 is provided in [Section 2.7.3](#_2.7.3_Characterisation_of). Some of the data provided to FSANZ for the risk assessment analyses was obtained from a predecessor of MXY0541, designated MXY0291. The major differences between these two strains is the copy number of the leghemoglobin gene (MXY0291 contains fewer copies) and MXY0541 contains extra DNA sequences associated with one of the haem-synthesis enzyme genes. Further information about MXY0291 can be found in the US FDA notification GRN 737 (US FDA 2018).

### 2.1.2 Technological purpose and justification

The Preparation, containing the soy leghemoglobin is to be added to meat analogue products produced by the applicant. The meat analogue products are intended for consumption by the general population (aged 2 years and older). There are multiple purposes for adding soy leghemoglobin to meat analogue products, including to provide the nutrition (i.e*.* source of iron), flavour, and aroma of myoglobin.

### 2.1.3 The soy leghemoglobin preparation specifications

The specifications for the Preparation are outlined in Table 1, which shows the combined minimum and maximum results for strains MXY0291, MXY0541 and the individual average results for each strain. The Preparation contains up to 9% soy leghemoglobin, which has a minimum protein purity of 65%. Acceptable limits for heavy metal and microbiological contaminants have been established. Additionally, no viable *P. pastoris* cells are present, which is in accordance with the recommendations for safety evaluation by the International Food Biotechnology Committee (IFBC, 1990). The inability to detect viable *Pichia* provides evidence of effective process controls for the removal and/or inactivation of the production organism. Details of the methods of analysis that were internally developed and validated by Impossible Foods for the evaluation of the specification parameters (i.e. content and purity of the soy leghemoglobin protein) were provided as confidential information. All other parameters are assessed by recognised and validated methods of analysis.

*Table 1:* Specifications and batch analyses for the soy leghemoglobin preparation product

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| *Parameter* | *Specification (%)* | *Min (%)1*  | *Max (%)1* | *Avg (%) MXY02912* | *Avg (%) MXY05413* |
| Total protein (w/w)4 | ≤14 | Average of 12.73% |
| Soy leghemoglobin (w/w) | ≤ 9 | 4.50 | 7.20 | 6.62 | 5.85 |
| Soy Leghemoglobin Purity (%) | ≥ 65 | 68.00 | 86.00 | 80.20 | 76.00 |
| Fat (w/w) | ≤ 2 | 0.03 | 1.70 | 0.05 | 0.80 |
| Carbohydrates (w/w) | ≤ 6 | 0.99 | 4.10 | 1.82 | 2.83 |
| Ash (w/w)  | ≤ 4 | 0.09 | 3.60 | 2.11 | 1.82 |
| Solids (w/w)5, 6 | ≤ 26 | 12.55 | 20.30 | 15.61 | 19.08 |
| Moisture (w/w) | ≤ 90 | 79.7 | 87.45 | 84.37 | 80.92 |
|  |  ***Range*** |
| Lead (ppm) | < 0.4 | <0.01 |
| Arsenic (ppm) | < 0.05 | <0.01 - 0.04 |
| Mercury (ppm)  | < 0.05 | <0.005 |
| Cadmium (ppm)  | < 0.2 | <0.001 - 0.003 |
| *Escherichia coli* EHEC (inc O157:H7) | Absent by test | Absent |
| *Salmonella* spp.  | Absent by test | Absent |
| *Listeria monocytogenes*  | Absent by test | Absent |

1. the results are combined data from MXY0291 & MXY0541 batch analyses

2. the number of batches analysed = 5

3. the number of batches analysed = 6

4. total protein is averaged from 3 batches

5. solids = Sum of total protein, fat, carbohydrate and ash

6. total organic solids (TOS) content = 22% (100% – (% Ash + % Water + % Diluent))

### 2.1.4 Product Stability

Stability of the Preparation prepared from nine batches of MXY0541, stored at the recommended -20°C were measured via an ultra-performance liquid chromatography assay developed and validated by Impossible Foods. Three parameters were measured: percent soy leghemoglobin, percent purity of soy leghemoglobin of total protein and pH. All data was compared to observed values on the day of production. Measurements were taken at variable times for up to 24 months. The results showed no significant changes in protein stability. The results are comparable to those obtained with the Preparation from MXY0291.

Stability of the soy leghemoglobin was also determined in the meat analogue product stored for up to 9 days at 4°C and 6 months at -20°C. There was minimal loss of soy leghemoglobin at day 5 and up to 10-15% loss at day 9. Similarly, after storage at -20°C for 6 months, there was up to 15% loss of soy leghemoglobin.

FSANZ considers the stability (shelf-life) of meat analogue products containing soy leghemoglobin would be determined by the same factors that determine shelf life in minced (ground[[4]](#footnote-5)) meat. Frozen product can be affected by factors that are not microbiological such as: meat colour and appearance; rancidity caused by chemical oxidation of fats at low temperature, and changes in texture caused by extended enzymatic activity; product drying during storage, e.g. freezer burn; or flavour and odour changes caused from protein denaturation or non-enzymatic browning reactions. Similarly, spoilage that can occur in the chilled product can be physical, chemical or microbiological. The two principal spoilage mechanisms that affect the chilled shelf-life of meat are microbial growth and oxidation of myoglobin causing browning of lipids causing rancidity.

Generally, meat is considered to be past its shelf-life when it is no longer acceptable to the consumer as a result of key attribute deterioration such as colour, flavour, texture, aroma or nutrient content. Another determinant of shelf-life would also be when a food-safety issue arises, where the food product may make consumers unwell (CSIRO 2006).

FSANZ conclude that the mechanisms inducing spoilage of meat would similarly affect meat analogue products containing soy leghemoglobin and therefore have no concerns around the stability information provided by the applicant.

### 2.1.5 Conclusion

Analyses of the data demonstrated that the Preparation containing the soy leghemoglobin is stable in chilled and frozen forms and within the final meat analogue product. FSANZ concludes that, in the form and prescribed amounts, soy leghemoglobin is technologically justified.

## 2.2 History of use

***Soy leghemoglobin***

Leghemoglobin is a haem-containing protein, with the highest levels of expression found in the roots of leguminous plants, more specifically in root nodules infected with nitrogen-fixing bacteria known as rhizobia (Kubo 1939; Appleby and Bergersen 1980; Appleby 1984). The edible portion of a legume is the seed or seed pod not the roots, thus leghemoglobin at the levels found in the root does not have a history of use in food.

While roots are not commonly used as food, plants do express a range of haemoglobins within the phytoglobin family (Hill et al. 2016) that share a high degree of structural similarity. These phytoglobins are expressed in leaf, stem and rapidly growing tissue in all higher plants (Garrocho-Villegas et al. 2007; Smagghe et al. 2009; Hill 2012). Expression of these phytoglobins is dependent on growth stage and reported levels are low (1-100 nM) compared to that found for leghemoglobin in root tissue (1-3 mM) (Appleby 1984, Arechaga-Ocampo et al. 2001; Lira-Ruan et al. 2001; Ross et al. 2001). The level of soy leghemoglobin present in the meat analogue products falls with the range found in plants and is equivalent to the reported levels of myoglobin found in beef (Cho et al. 2014; Texas A&M Institute 2019).

The role of leghemoglobin is to regulate oxygen levels in the root nodules, providing enough oxygen for cells undergoing cellular respiration while maintaining a moderately hypoxic environment to ensure nitrogen-fixation can occur (Appleby and Bergersen 1980; Ott et al. 2005). This oxygen transporting role is similar to the animal myoglobin, which is a haem protein that regulates oxygen levels in muscle tissue and is therefore present in meat products. Beyond function, leghemoglobin shares a common evolutionary origin as well as similar secondary and tertiary structure to myoglobin (Vainshtein et al. 1975; Arredondo-Peter et al. 1998). There is no evidence that myoglobin or other commonly ingested haem proteins are either allergenic or toxic to humans and myoglobin is considered to have a long history of safe use in food.

The soy leghemoglobin within the Preparation is produced by a GM strain of *P. pastoris* containing the leghemoglobin gene *LBC2* from soybean (*G. max*), encoding the leghemoglobin C2 protein ([P02236](https://www.uniprot.org/uniprot/P02236)[[5]](#footnote-6)). The introduced gene encodes a protein of 145 amino acids with a theoretical monoisotopic mass of 15515.08 Da ([ExPASy Compute MW tool](https://web.expasy.org/compute_pi/)[[6]](#footnote-7)). When bound to haem, the holoprotein will have an expected mass of 16,131 Da, noting that the haem component has a mass of 616 Da (Li et al. 1993). The characterisation and assessment of this novel protein is presented in [Sections 2.3.1](#_2.3.1_Characterisation_of), [2.3.2](#_2.3.2_Safety_of) and [2.3.3](#_2.3.3_Safety_of).

***Pichia pastoris***

*Pichia pastoris* is a cosmopolitan organism, commonly found on rotting wood. Since this organism was first characterised over 40 years ago, it has been chosen as a preferred organism for the production of heterologous proteins. Genomic sequencing analyses have led to the reclassification of *P. pastoris* into two distinct species: *Komagataella phaffii* and *K. pastoris* (Yamada et al. 1995; Kurtzman 2009; De Schutter et al. 2009; Sturmberger et al*.* 2016). The strain used to produce soy leghemoglobin is from a lineage derived from an original isolate NRRL Y-11430 (Figure 1), which has been reclassified as *K. phaffi*.

**NRRL Y-11430**

**X-33**

**GS115**

**Parent strain Bg11**

Figure 1: Lineage of the Pichia pastoris host organism Bg11. NRRL Y-11430 is the one of the original isolates from which many of the commercialised P. pastoris strains such as GS115 (Cregg et al. 1985), were derived for the production of food and industrial chemicals.

Both *Komagataella* organisms have been classified as Biosafety Level 1 organisms based on the [United States Public Health Service Guidelines](https://www.cdc.gov/biosafety/publications/bmbl5/index.htm) (US PHS). They have also been granted qualified presumption of safety (QPS) status by the European Food Safety Authority (EFSA) for the production of enzymes to be added to food (EFSA 2017; 2018). The restriction of QPS status to the production of enzymes is a reflection of the body of knowledge relating predominantly to use of *K. phaffi* in protein expression systems and as a model organism (EFSA 2018).

*P. pastoris* belongs to the *Saccharomycetaceae* family of budding yeast and is related to *Saccharomyces cerevisiae*, commonly known as baker’s and brewer’s yeast and *S. cerevisiae* var. *boulardi*, a common organism marketed as a probiotic yeast. *S. cerevisiae* and *S. cerevisiae* var. *boulardii* both have a long history of safe use in food. While there is limited evidence that *P. pastoris* has been consumed by humans, this organism does have a long history of safe use for the production of pharmaceuticals and industrial chemicals, including a number of enzyme processing aids approved by EFSA, US FDA and FSANZ (Spohner et al. 2015). Furthermore, a search of the literature did not identify any potential safety concerns associated with *P. pastoris*, *K. phaffi* or *K. pastoris* and no reports of adverse effects from products produced from *P. pastoris* strains were identified.

While *P. pastoris* may not have a history of use in food, a strain identified as X-33 has been investigated as a feed supplement in chicken and mice. X-33 is a mutation revertant of GS115, which is the parental strain from which Bg11 is derived ([Figure 1](#Fig1)). In these animal studies, no adverse effects were reported from the ingestion of viable *P. pastoris* X-33 (Gil de los Santos et al. 2012; Franca et al. 2015; Gil de los Santos et al. 2018). These data support the evidence that *P. pastoris* X-33 does not produce toxins or other compounds that might cause adverse effects.

A comparison of the three strains NRRL Y-11430, GS115 and X-33 ([Figure 1](#Fig1)) at the genome, transcriptome and proteome level have identified the differences between these highly related strains (Brady et al. 2019; Braun-Galleani et al. 2019). The presence of toxin encoding genes was not reported for any of these strains. However NRRL Y-11430 does contain two linear cytoplasmic plasmids, commonly known as killer plasmids (Banerjee and Verma 2000; Sturmberger et al, 2017). These plasmids have been characterised in a range of *Pichia* species and encode proteins used to protect the organism from pathogens and competitors. While NRRL Y-11430 does contain these plasmids, the host strain Bg11 used for the production of soy leghemoglobin does not contain these plasmids.

The most common *Pichia* proteins present in the Preparation have been identified and characterised and the assessment is presented in [Section 2.3.4](#_2.3.4__Safety).

The microbiological assessement undertaken by FSANZ did not identify any public health and safety concerns associated with the consumption of the Preparation containing *P. pastoris*.

## 2.3 Characterisation of the novel proteins in the soy leghemoglobin preparation

In considering the safety of novel proteins it is important to consider that a large and diverse range of proteins are ingested as part of the normal human diet without any adverse effects. Only a small number of dietary proteins have the potential to impair health (Delaney et al. 2008). As proteins perform a wide variety of functions, different possible effects are considered during the safety assessment including potential toxic, anti-nutrient or allergenic effects.

### 2.3.1 Biochemical analyses of the soy leghemoglobin in the soy leghemoglobin preparation

The soy leghemoglobin component of the Preparation was isolated by UPLC and peptide mapping was performed using mass spectrometry on tryptic-digested peptides. The results confirmed that the amino acid sequence was as expected ([Figure 2](#Fig2)). The analysis showed absence of the N-terminal methionine (M) and C-terminal alanine (A) and phenylalanine (F). The absence of the N-terminal M is a common post-translational modification. The absence of the C-terminal AF is due to the small size of the peptide (2 amino acids) resulting from the tryptic digest, that would not be highly visible in the spectroscopic analysis. Therefore, coverage of the peptide mapping was 99%. Mass spectrometry of the undigested protein determined that the mass of the full *Pichia* expressed protein was 15384.04 Da.

 1 -GAFTEKQEA LVSSSFEAFK ANIPQYSVVF YTSILEKAPA AKDLFSFLSN GVDPSNPKLT

 61 GHAEKLFGLV RDSAGQLKAN GTVVADAALG SIHAQKAITD PQFVVVKEAL LKTIKEAVGD

 121 KWSDELSSAW EVAYDELAAA IKKAF

*Figure 2: Tryptic peptide map of the leghemoglobin in the soy leghemoglobin preparation. Trypsin is an enzyme that digests proteins after lysine (K) residues. These residues are shown in red. The mapping provided sequence coverage of 99%.*

As a control for the identification of the protein and mass determination, the three main isoforms of leghemoglobin were isolated from the root nodules of soybean. After purification by column chromatography, the mass spectrum for the three isoforms were identified and the mass of the leghemoglobin C2 protein expressed in soybean was 15384.04 Da.

The observed masses for the *Pichia* and plant-extracted leghemoglobins are the same but differ to the [expected mass](#MWofLegH) (15515.08 Da). Taking into consideration the missing N-terminal M in the expressed protein, an adjusted theoretical mass was calculated2 as 15384.04 Da, which agrees with that observed. This data also indicates that the leghemoglobin expressed in *Pichia* has not undergone post-translational modification, as the observed mass is equivalent to the expected mass.

Mass spectrometry also identified a major peak with a mass of 616.1752 Da, that matches the [expected peak](#MWofHaem) for haem B. This major peak was co-associated with other isotopic forms, ranging from 614-619 Da. Further analysis using UV-visual spectroscopy showed that haem dissociated from the soy leghemoglobin when the protein was denatured.

### 2.3.2 Safety of the expressed leghemoglobin protein – potential allergenicity

Leghemoglobin is normally expressed in the part of the soybean plant (root nodules) that is not typically consumed by humans therefore it is a new protein to the diet. As discussed in [Section 2.2](#_2.2_History_of), leghemoglobin has structural similarity to myoglobin, a commonly ingested protein that is not associated with allergic reactions in humans.

To assess potential allergenicity, consideration must be given to whether the new protein is one which certain individuals may already be sensitive as well as whether a protein new to the diet is likely to induce allergic reactions in some individuals (Codex, 2009). There are presently no reliable animal models for the assessment of allergenicity, and no single test that can be applied to a protein to predict whether it is likely to be allergenic to humans. Instead, the assessment follows a step-wise, weight of evidence approach that considers:

1. source of the new protein
2. similarity of the new protein amino acid sequence to known allergens
3. structural properties:
	1. susceptibility to enzymatic degradation
	2. thermal stability
	3. acid stability

If the new protein is from a source known to be allergenic, consideration may also be given to specific serum screening using sera obtained from individuals with a clinically validated allergy to the source of the protein as well as oral challenge studies with allergic individuals.

***Source of the new protein***

The soy leghemoglobin component of the Preparation is sourced from the soybean plant *G. max*. which is known to be allergenic to humans. Certain products derived from soybean require a mandatory [allergen warning label](https://www.foodstandards.gov.au/consumer/foodallergies/Pages/Allergen-labelling.aspx)[[7]](#footnote-8) in Australia and New Zealand, due to soy being one of the eight common food allergen sources. Some soy products (e.g. refined soybean oil) are exempt from allergen labelling because they have been processed in a way that makes them safe for soy allergic individuals.

The examination of soy allergic individuals have identified allergen specific immunoglobulin E (IgE) antibodies that bind to proteins found in soybean (Ogawa et al. 1993; Holzhauser et al. 2009; Ito et al. 2011). To date, there have been 42 proteins identified in the seed that react to IgE antibodies ([AllergenOnline database](http://www.allergenonline.org/databasebrowse.shtml)[[8]](#footnote-9)) but only a portion of these proteins have been confirmed as clinically relevant to humans (Selb et al. 2017).

***Similarity of the leghemoglobin amino acid sequence to known allergens***

The applicant provided results of *in silico* analyses comparing the *Pichia*-expressed soy leghemoglobin sequence ([Figure 2](#Fig2)) to known allergenic proteins listed in the Food Allergy Research and Resource Program (FARRP) dataset, which is available through [AllergenOnline](http://www.allergenonline.org)[[9]](#footnote-10) (University of Nebraska). At the date of the search, there were 1,956 sequences in the allergen database. This broad ranging database includes all the known allergens from legumes, including soybean and peanut. The methodology followed the approach recommended in the Codex guidelines for foods derived from modern biotechnology (Codex 2009).

The results from the analyses identified no similarity between the leghemoglobin protein and known allergenic proteins. The results from these analyses have been published by Jin et al. (2018).

***Structural properties***

To assess the susceptibility to enzymatic degradation, *Pichia*-produced soy leghemoglobin (test protein) from production strain MXY0541, was incubated with pepsin at a concentration of 10.6U enzyme/μg protein. The incubation was performed at 37°C for 0-60 m, in a simulated gastric fluid system at pH 2.0 (Thomas et al. 2004; Ofori-Anti et al. 2008). Controls included a no enzyme control (test protein only) and a no protein control (enzyme only), incubated for 0 and 60 m. The extent of digestion was visualised by Coomassie staining of the protein run on a tris-glycine SDS-PAGE gel. An LOD was not determined for this test system but samples from the digestion mixes were analysed by liquid chromatography combined with mass spectrometry and the resulting peptides identified.

The results from the pepsin digestions showed that by 2 m there was no visible soy leghemoglobin remaining in the digestion mix. There was no loss of band intensity in the no enzyme control incubated for 60 m therefore the loss of protein in the reaction mix indicated that leghemoglobin was being fully digested by pepsin. Additional evidence was provided through the analysis of the resulting peptides. A total of 49 different peptide fragments were identified across 3 digestion samples, ranging from 5-16 amino acids in length. The majority of the fragments were between 8-11 amino acids. Analysis of the peptide data confirmed that by 2 m soy leghemoglobin is fully digested by pepsin in a simulated gastric fluid assay. These data support the results obtained for the soy leghemoglobin from the MXY0291 production strain (Jin et al*.* 2018).

In order to assess thermal stability, the applicant provided results from a melt curve analysis. The test sample was the leghemoglobin in the Preparation. Control samples were horse myoglobin and chicken egg lysozyme. The data shows that leghemoglobin denatures at 64°C. This indicates that at standard grilling and cooking temperatures, the leghemoglobin protein will be fully denatured.

The applicant also provided results examining acid stability by looking at the dissociation of haem by UV-visual spectroscopy. Examination of the Soret region at pH 7 and pH 2, showed the absorbance decreased at the acidic pH. This reduced absorbance demonstrates the release of the haem prosthetic group due to denaturation of the protein. These data indicate that at an acidic pH that would be found in the stomach, soy leghemoglobin is susceptible to acidic denaturation.

***Specific serum screening***

Serum screening is an *in vitro* test that allows examination of cross-reactivity of the soy-allergen IgE antibodies from an allergic patient to the leghemoglobin protein. This test requires collection of sera from a range of patients, which is an invasive technique. FSANZ does not consider there would be any additional value in undertaking serum screening given the findings from the weight of evidence approach outlined above. These studies, showing soy leghemoglobin shares no amino acid sequence similarity to known allergens and is readily digested by pepsin, indicates a low likelihood of cross-reactivity between the protein and IgE antibodies of allergic individuals. Moreover, there is an absence of evidence linking haem-proteins with allergenic reactions in humans.

### 2.3.3 Safety of the expressed soy leghemoglobin protein – potential toxicity

To assess potential toxicity, a weight of evidence approach is employed (Codex, 2009) that considers:

1. similarity of the new protein amino acid sequence to known toxins
2. information on the stability of the protein to enzyme degradation
3. animal toxicity studies if the protein shares significant similarity to known toxins or resistance to proteolysis or if it is not similar to proteins that have previously been consumed safely in food

***Similarity of the soy leghemoglobin amino acid sequence to known toxins***

The full length sequence for soy leghemoglobin ([Figure 2](#Fig2)) was compared to sequences in the NCBI-Entrez database designated as “toxin” or “toxic” . No results were returned when using the keyword “toxic” and an *E* score set to less than 10. Two proteins were returned using the keyword “toxin”, with sequence identity of 35% and an *E* score of less than 0.1 however, the alignments of these proteins to leghemoglobin was short (31%). Subsequent analysis identified that the homologous proteins are from toxin-producing bacteria but that the proteins themselves are not toxins. It was concluded that the soy leghemoglobin protein does not share similarity to known protein toxins (Jin et al. 2018).

***Stability of the protein to enzyme degradation***

An analysis of the stability of the protein to enzyme degradation has already been provided in the [*Potential Allergenicity*](#Proteolysis) discussion ([Section 2.3.2](#_2.3.2_Safety_of)).

***Animal Toxicity studies***

A range of toxicity assessments were performed by the Applicant and the analyses are provided in [Section 2.4](#_2.4_Toxicological_assessment).

### 2.3.4 Safety of the native *Pichia* proteins present in the soy leghemoglobin preparation

The Preparation that is the focus of this study, exists as a partially purified cell lysate containing both the novel soy leghemoglobin and native *Pichia* proteins. A comparison of different batches of LegH from MXY0291 and MXY0541, using Coomassie stained protein gels, found several *Pichia* proteins with similar molecular masses that were consistently present. FSANZ notes that not all preparations contained the same proteins, nor proteins at the same levels. This is expected because the proteins that are present in cells would be dependent on the cell cycle stage of each cell, at the time the fermentation run was completed.

The identity of the *Pichia* proteins consistently present in the preparation was determined by mass spectrometry. The amino acid sequences of these proteins were then subjected to bioinformatic analyses, looking for similarity to known toxins and allergens. The data indicates there are no similarities between the proteins identified and known toxins or allergens. These data pertaining to the *Pichia* proteins identified in the MXY0291 preparations were also included in the publication by Jin et al. (2018).

### 2.3.5 Conclusion

Biochemical analysis of the novel soy leghemoglobin showed the protein expressed was equivalent to that found in the host plant. The molecular mass of the protein was as expected, it was present as a holoprotein and had not undergone post-translational modification, such as glycosylation. The protein was also shown to be denatured when exposed to heat and acidic conditions.

No safety concerns were identified regarding the potential allergenicity or toxicity of soy leghemoglobin or the *Pichia* proteins. There were no significant similarities between the protein and known allergens or toxins. All the novel proteins were also shown to be susceptible to pepsin digestion and were denatured at standard cooking temperatures and in acidic conditions that mimic the stomach environment.

## 2.4 Toxicological assessment of soy leghemoglobin preparation

The applicant submitted reports of dietary toxicity studies in rats and *in vitro* genotoxicity studies. The Preparation used in these studies was produced using the initial MXY0291 production strain. This test item is considered to be relevant to the assessment of the Preparation produced by the MXY0541 strain because the expressed soy leghemoglobin protein is equivalent in both strains, several of the Pichia proteins expressed in the Preparation from MXY0541 are also present in the Preparation from MXY0291, and the composition of the Preparation from each strain meets the same specifications.

The Joint Food and Agriculture Organization (FAO)/World Health Organisation (WHO) Expert Committee on Food Additives (JECFA) has previously considered the safety of an enzyme preparation produced by a genetically modified form of *P. pastoris*, phospholipase C, in 2008 (WHO 2009). JECFA noted that *P. pastoris* is a methylotropic yeast which is not known to be associated with disease in humans or animals. The enzyme preparation was found to be free of antimicrobial activity and mycotoxins, was not genotoxic *in vitro* or *in vivo*, and did not cause adverse effects in a 13-week oral toxicity study in rats at doses up to 2000 mg/kg bw/day (1672 mg/kg bw/day on a TOS basis).

### 2.4.1 Toxicity studies in animals

The applicant submitted reports of three repeated dose dietary toxicity studies with the Preparation conducted in rats. The results of these studies were also reported in the publication by Fraser et al. (2018). The Preparation test article used in all three studies was freeze-dried to enable an increased concentration of soy leghemoglobin in the feed, and to facilitate homogenous mixing with the diet.

##### 14-day dietary toxicity/palatability study in rats (Product Safety Labs 2016a) Regulatory status: Non-GLP; conducted in accordance with OECD TG 407 (2008) and US FDA Redbook 2000, IV. C. 4. a. (2007a)

CRL Sprague Dawley CD® IGS rats (6/sex/group), aged 7 – 8 weeks, were administered freeze-dried Preparation (47.6% soy leghemoglobin, Lot no. PP-PGM2-16-081-302) in the diet at concentrations targeting soy leghemoglobin doses of 0, 125, 250 or 500 mg/kg bw/day for 14 days. Stability of the test substance and homogeneity in the diet were evaluated and confirmed. Clinical signs were monitored daily and weekly detailed observations were performed. Body weight and food consumption were monitored at intervals throughout the study. At study termination blood samples were collected for haematology and animals were killed and subjected to a gross necropsy. The liver and spleen were weighed and the liver, spleen and femur bone marrow were subjected to histopathological examination.

No mortalities occurred during the course of the study. In-life clinical signs were limited to discolouration of the urine in 6/6 males and 5/6 females administered the high dose on study day 5 only. This finding was not considered to be treatment related or of toxicological significance in the absence of a correlation with clinical haematology. Mean weekly body weights and body weight gains in all treated groups were similar to those of controls. Food consumption and food efficiency were similar in all groups, with the exception of significantly lower food consumption between study days 0 – 14 in females given the low and mid-doses. Absolute and relative (to body weight) weights of the liver and spleen were similar to those of controls. Numbers of white blood cells, neutrophils and lymphocytes in males were >25% lower than those of controls, but no dose response was observed at the lower doses and the study authors considered there were no changes in haematology parameters attributable to administration of the Preparation. No treatment related macroscopic or microscopic changes were observed in the liver, spleen and bone marrow. It was concluded that doses of at least 500 mg/kg bw/day soy leghemoglobin would be tolerated by rats in studies of longer duration.

##### 28-day dietary toxicity study in rats (Product Safety Labs 2017a) Regulatory status: GLP; conducted in compliance with OECD TG 407 (2008) and US FDA Redbook 2000, IV. C. 4. a. (2007a)

CRL Sprague Dawley CD® IGS rats (10/sex/group), aged 7 – 8 weeks, were administered freeze-dried Preparation (48.82% soy leghemoglobin; Lot no. PP-PGM2-16/088-301; Total Organic Solids [TOS] content 92.5%) via the diet at concentrations targeting doses of 0, 512, 1024 or 1536 mg/kg bw/day Preparation for 28 days. These doses corresponded to 0, 474, 947 and 1421 mg/kg bw/day on a TOS basis. Stability of the test substance and homogeneity in the diet was evaluated and confirmed. Body weight and food consumption were monitored at intervals throughout the study. Clinical signs were monitored daily, with detailed observations made weekly. Ophthalmologic evaluations were performed in all animals before study initiation and again on study day 23. Blood and urine samples were collected for haematology, blood coagulation, clinical chemistry and urinalysis during the final week of the study. At study termination, animals were killed and subjected to gross necropsy. Organs and tissues were collected and weighed. Histopathological examination was performed on organs and tissues from animals in the control and high dose groups. Based on observations in the uterus, female reproductive organs from animals in all four groups underwent histopathology. Apeer review of the histopathology of the female reproductive organs was also performed.

No mortalities were observed, and there were no clinical signs attributable to administration of the test item. Mean body weights, body weight gains, food consumption and food efficiency from study days 0 – 28 were similar in all groups. A transient significant reduction in mean daily body weight gain (p < 0.01) and mean food efficiency (p < 0.01) was found between days 7 – 14 in females administered the Preparation at 512 mg/kg bw/day.Significant increases in food consumption (p < 0.05-0.01) in males given 1024 mg/kg bw/day and 1536 mg/kg bw/day were observed on days 7 – 14 and 7 -10, respectively. These findings were not considered to be of toxicological relevance given that overall body weight gains and food efficiency over the course of the study were similar to controls and there was no dose-response. Absolute and relative organ weights in all treated groups were similar to controls, with the exception of decreased absolute and relative (to body weight and brain weight) uterus weights in treated females, which were statistically significant (p < 0.05-0.01) in the low and high dose groups. No treatment-related changes in ophthalmology, haematology, clinical chemistry or urinalysis parameters were observed. A small number of statistically significant changes in haematology ([Table 2](#Table2)) and clinical chemistry ([Table 3](#Table3)) values were observed, but these were not considered to be treatment-related as they were of a small magnitude, did not show a dose-response and were only seen in one sex. Changes in coagulation parameters were limited to a slight increase in activated partial thromboplastin time in males given 1024 and 1536 mg/kg bw/day compared with controls (24.9, 23.9 and 20.2 seconds, respectively, p < 0.05). This change was not considered adverse as the increase did not show a dose-response, the magnitude was slight and there were no correlated pathological or clinical changes.

*Table 2:* Haematology values for which statistically significant changes were observed

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| *Parameter* | *0 mg/kg bw/day* | *512 mg/kg bw/day* | *1024 mg/kg bw/day* | *1536 mg/kg bw/day* |
| Females |
| Red blood cells (x106/µL) | 7.59 ± 0.24 | 8.01 ± 0.38\* | 7.86 ± 0.24 | 7.63 ± 0.3 |
| Haemoglobin (g/dL) | 15.3 ± 0.5 | 16.2 ± 0.5\* | 15.7 ± 0.4 | 15.5 ± 0.6 |
| Haematocrit (%) | 43.6 ± 1.2 | 45.9 ± 1.2 | 44.7 ± 1.3 | 44.0 ± 1.7 |
| Absolute basophil count (x103/µL) | 0.04 ± 0.01 | 0.07 ± 0.03\* | 0.06 ± 0.03 | 0.05 ± 0.04 |
| Absolute Reticulocyte count (x103/µL) | 205.8 ± 33.9 | 182.4 ± 32.9 | 169.1 ± 30.9\* | 184.2 ± 33.7 |

\*p < 0.05, Dunnett’s test

*Table 3:* Clinical chemistry values for which statistically significant changes were observed

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| *Parameter* | *0 mg/kg bw/day* | *512 mg/kg bw/day* | *1024 mg/kg bw/day* | *1536 mg/kg bw/day* |
| Males |
| Albumin (g/dL) | 3.1 ± 0.1 | 3.2 ± 0.1 | 3.3 ± 0.1\* | 3.2 ± 0.1 |
| Potassium (mmol/L) | 5.03 ± 0.25 | 5.19 ± 0.26 | 5.55 ± 0.61\* | 5.10 ± 0.25 |
| Females |
| Alkaline phosphatase (U/L) | 137 ± 16 | 107 ± 19\* | 121 ± 29 | 108 ± 25\* |
| Glucose (mg/dL) | 118 ± 15 | 103 ± 10\* | 104 ± 10\* | 110 ± 14 |
| Globulin (g/dL) | 2.9 ± 0.1 | 3.1 ± 0.2 | 3.1 ± 0.2\* | 3.0 ± 0.1 |
| Calcium (mg/dL) | 10.5 ± 0.3 | 10.9 ± 0.3\* | 11.0 ± 0.3\* | 10.7 ± 0.4 |
| Chloride (mmol/L) | 102.6 ± 1.2 | 101.3 ± 1.4\* | 101.1 ± 1.0\* | 102.1 ± 1.1 |

\*p < 0.05, Dunnett’s test

No macroscopic or microscopic findings related to administration of the test substance were observed. The presence of fluid-filled uteri, typically associated with the proestrus/estrus stage of the oestrous cycle, was decreased in treated female rats compared with controls. The incidence of animals in metestrus was not significantly different between groups. Histopathological examination of the female reproductive organs found a decreased incidence of dilated uterine lumens in low and high dose rats compared with controls. Uteri were dilated in 4/10 control females, which was consistent with estrus. There were no females with dilated uterine lumens in the low and high dose groups, and 2/10 in the mid-dose group, which correlated with lower incidences of animals in the proestrus/estrus stage of the estrus cycle compared with controls. Rats given the low or high dose tended to be in the metestrus stage of the oestrous cycle, which correlated with the lower uterine weights observed in these groups. This was considered to be an unusual distribution, but the presence of both new and old corpora lutea in females from all groups indicated that females were cycling normally and there were no treatment-related effects on the estrus cycle. The pathology peer review of reproductive organs from all female animals concurred with this conclusion.

The no observed adverse effect level (NOAEL) in this study was 1536 mg/kg bw/day Preparation, the highest dose tested. Based on a TOS content of 92.5%, this corresponds to 1421 mg/kg bw/day TOS.

##### Investigative 28-day dietary study in female rats with a 14-day pre-dosing oestrous cycle determination (Product Safety Labs 2017b) Regulatory status: Non-GLP; conducted in accordance with OECD TG 407 (2008) and US FDA Redbook 2000, IV. C. 4. a. (2007a)

Female CRL Sprague Dawley CD® IGS rats (15/group), aged 7 – 8 weeks, were examined for the potential effects of the Preparation on the oestrous cycle and reproductive organ histopathology. Prior to dosing, the oestrous cycle was determined in all rats for 14 days (study days 0-13) by vaginal lavage. On study day 14, animals were administered diets containing target doses of 0, 512, 1024 or 1536 mg/kg bw/day of the Preparation (48.82% soy leghemoglobin; Lot no. PP-PGM2-16/088-301) for 28 days. These doses corresponded to 0, 474, 947 and 1421 mg/kg bw/day on a TOS basis. Homogenous distribution of the test substance in the diet was confirmed. Clinical signs were monitored daily, with detailed clinical observations made weekly during the test substance exposure period. Body weight, body weight gain, food consumption and food efficiency were monitored periodically throughout the study. Oestrous cycling was evaluated by vaginal lavage on study days 29-42 and at termination on day 43. At the end of the study, animals were killed and the uterus and ovaries (with oviducts) from all animals were weighed. Oestrous cycle stage at study termination was also determined in all animals by blinded evaluation of the anterior portion of the vagina, the cervix, the uterine bifurcation and horns, the oviducts and the ovary. Reproductive organs and tissues (ovaries, oviducts, uterus, cervix and anterior-most portion of the vagina) from the control and high dose groups were evaluated microscopically.

No mortalities occurred during the course of the study, and no treatment-related clinical signs were observed. Body weight, body weight gain, food consumption and food efficiency were similar to controls in all treated groups. The mean number of oestrous cycles for rats in the treatment groups were similar to controls both prior to and during test substance administration. No test substance-related changes in organ weights were observed. Microscopic analysis of the oestrous cycle at study termination found that all animals were cycling normally with the exception of one animal given the low dose, which appeared to have a prolonged estrus. This finding was considered to be incidental because of the lack of similar findings in animals at the higher dose levels. No treatment-related macroscopic or microscopic changes were observed in the organs and tissues examined.

It was concluded that under the conditions of this study administration of the Preparation at doses up to 1536 mg/kg bw/day did not affect the oestrous cycle of female Sprague Dawley rats. Based on a TOS content of 92.5%, this corresponds to 1421 mg/kg bw/day TOS.

### 2.4.2 Genotoxicity studies

Two *in vitro* genotoxicity studies were submitted, a bacterial reverse mutation assay (Ames test) and a chromosomal aberration test in human lymphocytes. These studies used the standard liquid formulation of the Preparation. The results of these studies were also included in the publication by Fraser *et al.* (2018).

##### Bacterial reverse mutation assay (Product Safety Labs 2016b) Regulatory status: GLP; conducted in compliance with OECD TG 471 (1997), US FDA Redbook 2000, IV. C. 1. a. (2007b) and Commission Regulation (EC) No 440/2008 B.13/14

The test article for this study was the Preparation (Batch No. PP-PGM2-16-015-101; purity 6.74% soy leghemoglobin[[10]](#footnote-11)) and the vehicle control was water. Test systems for this assay were *Salmonella typhimurium* strains TA1535, TA1537, TA98, TA100, and *Escherichia coli* strain WP2 *uvr*A. For assays conducted without S9 mix for metabolic activation, positive control articles were sodium azide for TA 1535 and TA 100, ICR 191 Acridine for TA157, daunomycin for TA98 and methyl methane sulfonate for WP2 uvrA. For assays conducted with addition of S9 mix for metabolic activation, 2-aminoanthracene in DMSO was used as the positive control for all test strains. The main test was conducted in triplicate with and without metabolic activation using the plate incorporation method, and results were confirmed by a second test using the pre-incubation method. Concentrations of the Preparation in both tests were 23.384, 74, 233.84, 740, 2338.4, 7400, 23,384 and 74,000 µg/plate, corresponding to soy leghemoglobin concentrations of 1.58, 5.0, 15.8, 50, 158, 500, 1580 and 5000 µg/plate.

For the plate incorporation test, test solution (100 µL), 500 µL S9 mix or substitution buffer, 100 µL bacterial solution and 2.0 mL of overlay agar were mixed and poured on minimal agar plates. After the agar set, plates were incubated for approximately 65 hours at 37°C and then assessed for numbers of revertant colonies and effects on the growth of the bacterial lawn. For the confirmatory pre-incubation test, the same volumes of test solution, S9 mix or substitution buffer and bacterial solution were incubated under agitation for approximately 30 minutes at 37ºC prior to mixing with the overlay agar, pouring onto the plates and proceeding as described for the plate incorporation test.

The test article showed no evidence of precipitation or toxicity, and there was no increase in revertant colonies compared with vehicle controls, with or without S9 mix in either test. A substantial increase in revertant colonies was observed in the presence of the positive controls, confirming the validity of the test system.

It was concluded that the Preparation was not mutagenic under the conditions of the study.

##### Chromosomal aberration assay in cultured human lymphocytes (Eurofins 2017) Regulatory status: GLP; conducted in compliance with OECD TG 473 (2014) and Commission Regulation (EC) No 440/2008 B.10

The test article for this study was the Preparation (Batch No. PP-PGM2-16-015-101; purity 6.74% soy leghemoglobin8) and the vehicle control was cell culture medium. The test system comprised human peripheral blood lymphocytes collected from a healthy non-smoking donor with no known recent exposure to genotoxic chemicals and radiation. Lymphocytes were pre-cultured in the presence of the mitogen phytohaemagglutinin for 48 hours prior to the start of exposure. Following a preliminary cytotoxicity test two experiments were conducted. In experiment I cells were exposed to the test substance in the presence or absence of metabolic activation (S9 mix) for 4 hours, then washed and cultured in fresh medium for a further 20 hours. In experiment II cells were exposed to the test substance for 24 hours in the absence of S9. Cells were cultured in duplicate. Parallel cultures of the negative controls and highest concentrations were treated in the presence of BrdU in order to determine the proliferation index. Positive controls were cyclophosphamide and ethylmethanesulfonate in the presence and absence of metabolic activation, respectively. Two hours prior to harvest, Colcemid® was added to the cultures to arrest cells in metaphase. At harvest cells were stained and 300 metaphases per concentration were evaluated for the presence of chromosomal aberrations.

Concentrations were reported as the concentration of soy leghemoglobin. In experiment I without S9, mitotic indices lower than 70% of control were observed at 1000 µg/mL (69%), 2500 µg/mL (56%) and 5000 µg/mL (54%). In experiment I with S9, no reductions in mitotic index below 70% of control were observed. In experiment II, mitotic indices below 70% of control were seen at 500 µg/mL (69%), 1000 µg/mL (53%), 2000 µg/mL (26%), 3000 µg/mL (13%), 4000 µg/mL (38%) and 5000 µg/mL (42%). Concentrations evaluated for chromosomal aberrations were 500, 1000, 2500 and 5000 µg/mL in experiment I with and without S9. Concentrations of 100, 200, 500 and 1000 µg/mL were evaluated in experiment II, due to the substantial cytotoxicity at concentrations ≥ 2000 µg/mL. No precipitation was observed in experiment I. Precipitation was observed at concentrations ≥ 500 µg/mL in experiment II, but this did not affect the evaluation of aberration rates.

No significant increases in the percentage of cells with chromosomal aberrations compared with controls were observed. In experiment I, with and without S9, some concentrations had aberration rates slightly higher than the laboratory’s historical negative control ranges, but these were not considered to be relevant due to the negative control also being slightly higher than the historical negative control range and/or a lack of a dose-response. No significant increases in the number of polyploid cells were observed. A distinct increase in the percentage of structurally aberrant cells was observed in all the positive control cultures, confirming the validity of the assay.

It was concluded that the Preparation was not clastogenic in human lymphocytes.

### 2.4.3 Conclusion

The Preparation was not genotoxic *in vitro*, and did not cause adverse effects in short-term toxicity studies in rats. The NOAEL in a 28-day repeated dose dietary toxicity study in rats was the highest dose tested, 1536 mg/kg bw/day or 1421 mg/kg bw/day on a TOS basis.

## 2.5 Nutritional impact

### 2.5.1 Introduction

Dietary iron is an essential micronutrient that is required for a range of cell functions. Iron can be found in several proteins including haemoglobin which transports oxygen for cell respiration; myoglobin which acts as an oxygen reserve in muscle cells; cytochromes which are involved in metabolism; transferrin, an iron transport protein; and ferritin and haemosiderin that store iron in the body. Iron can exist in two interchangeable oxidation states that allow it to reversibly bind to ligands such as oxygen, nitrogen and sulphur.

The iron content of the body is highly conserved, with only a small amount being lost each day in the absence of bleeding (Bothwell et al. 1979). The estimated average requirement (EAR) for iron is 6 mg/day for adult males, 8 mg/day for women to age 50 and 5 mg/day thereafter, and 4–8 mg/day for children aged 1–18 years. The EAR during pregnancy is 23 mg/day (age 14–18 years) and 22 mg/day (age 19–50 years) (NHMRC and MoH 2006).

The two main forms of dietary iron are haem and non-haem iron. Iron from animal tissue including meat, fish and poultry may be haem or non-haem, with haem iron contributing approximately 55–70% of the total iron content (Lombardi-Boccia et al. 2002). Most iron from plant sources, including cereals and legumes, is non-haem iron. Although plants do contain small quantities of haem iron they are not considered to be a major source of haem iron in the diet.

### 2.5.2 Bioavailability of iron from soy leghemoglobin

As discussed previously (Section 2.2 and 2.3.2) soy leghemoglobin is structurally similar to myoglobin; is completely digested by pepsin; and the haem moiety of leghemoglobin is released at pH 2, which is within the pH range in the stomach. The applicant provided data indicating that soy leghemoglobin and equine myoglobin denature at 64°C and 69°C, respectively. Published studies have reported that most of the myoglobin (approximately 85%) in beef samples denatures at 70°C (Trout 1989; Hunt et al. 1999). Therefore soy leghemoglobin has similar thermal stability properties to bovine myoglobin.

Several factors contribute to the bioavailability of iron including other dietary constituents of a meal, the form of iron (e.g haem or non-haem), physiological iron status, exercise, and age. Iron absorption is regulated by the body, increasing in individuals that are iron deficient and decreasing in cases of iron overload.

The bioavailability of non-haem iron can be decreased by dietary factors such as phytates from grains and legumes, tannins from spices, fruits, vegetables or tea, calcium, and phosphates (Hurrell and Egli 2010), with phytate being the primary inhibitor (Hallberg et al. 1987). The insolubility of complexes formed between iron, phytate and proteins is considered to be the major reason for phytate inhibition of iron absorption (Champagne 1988). However, the presence of meat, fish, poultry or ascorbic acid (vitamin C) enhances the bioavailability of non-haem iron and have been found to counteract the effect of inhibitors such as phytates and tannins (Lopez and Martos 2004).

In general, haem iron is more bioavailable than non-haem iron. It has been estimated that 15–25% of haem iron is absorbed compared to 5–12% of non-haem iron (Hallberg 1983), and that although haem iron contributes 10–15% of the total dietary iron intake of omnivores, it represents at least 40% of the total absorbed iron (Björn-Rasmussen et al.1974).

Similar to non-haem iron, other dietary components can affect the absorption of haem iron. For example, calcium has been shown to inhibit haem iron absorption at intake levels greater than 40 mg (Hallberg 1998), while the presence of meat enhances haem iron absorption (Carpenter and Mahony 1992). The reason for the improved bioavailability of both haem and non-haem iron due to the presence of meat may be due to the binding of peptides formed from digestion of meat which prevent the formation of insoluble iron aggregates that are poorly absorbed (Lopez and Martos 2004). The interaction of haem with peptides produced from the proteolytic digestion of globin is also thought to prevent the formation of insoluble haem polymers (Hooda et al. 2014).

The applicant provided a published study to support the bioequivalence of haem iron from soy leghemoglobin and bovine haemoglobin (Proulx and Reddy 2006). This in vitro study used Caco-2 cells to compare the iron bioavailability of crude and purified soy leghemoglobin compared to bovine haemoglobin. Caco-2 cells are a human intestinal adenocarcinoma cell line that exhibit enterocyte-like properties and have been previously used for non-haem iron bioavailability studies (Au and Reddy 2000; Yun et al*.* 2004).

Experiments were undertaken with and without a food matrix. The change in ferritin concentration (“ferritin response”) was used as an index of iron bioavailability (Glahn et al. 1998). In the food matrix experiment, corn tortillas were prepared using unfortified tortilla flour and either soy root nodule (SRN) extract, partially purified soy leghemoglobin (LHbA), bovine haemoglobin (BHb) or ferrous sulphate. Tortillas were cooked and lyophilised. Gastric and intestinal digestion was then simulated for all samples. The non-food matrix test solutions contained either SRN, purified soy leghemoglobin (LHbD), BHb or ferrous sulphate. Caco-2 cells were incubated with test solutions for 24 h, lysed and ferritin concentration was calculated as percentages compared to ferrous sulphate.

The results of the non-food matrix experiment found that the ferritin response for BHb was approximately 2-fold higher than all other samples. No statistically significant difference was observed between the ferritin responses for SRN, LHbD and ferrous sulphate (P > 0.05).

For the food matrix experiment, the LHbA and BHb enriched tortillas exhibited similar ferritin responses that were 27 ± 6% and 33 ± 10% higher, respectively, than the ferrous sulphate response (P < 0.05). The ferritin response for SRN was 19% lower than for ferrous sulphate, although it was not statistically significant (P > 0.05). Heat treatment had no effect on the ferritin response with either LHbA or BHb.

The results of this *in vitro* study indicate similar bioavailability of soy leghemoglobin and bovine haemoglobin, however no *in vivo* studies of bioavailability were provided in the application. FSANZ searched Pubmed using the search terms “((leghemoglobin or leghaemoglobin) and (bioavailability or absorption))” and no additional publications on the bioavailability of haem iron from soy leghemoglobin were located.

***Bioavailability of haem iron from soy leghemoglobin in an Impossible Foods meat analogue product***

As previously discussed, soy leghemoglobin is a haem protein with structural similarities to animal myoglobins. In general haem iron is more bioavailable than non-haem iron, however it is usually consumed as part of a meal containing meat and the meat component has been shown to enhance the bioavailability of haem iron.

The evidence is conflicting on the effect of plant proteins similar to those that would be present in the proposed Impossible Foods meat analogue products on haem iron bioavailability and there is limited research in this area. Some studies have found that partial substitution of meat with soy protein increased the bioavailability of haem iron (Lynch et al. 1985) while others have reported that the absorption of haem iron from meat-free meals is half that of haem in a meal with meat when protein content is constant (Hallberg et al*.* 1979). A more recent study found that the addition of cereal, pea or lentil protein to haem iron did not affect iron absorption but soy protein had an inhibitory effect. However the relevance of this study is unclear as the haem iron used in the study was dissociated from globin and pure haem iron has low solubility at low gastric pH (Vaghefi et al. 2002; Weinborn et al. 2015).

### 2.5.3 Conclusion

Approximately 15–25% of haem iron in the diet is absorbed compared to 5–12% of non-haem iron. Iron absorption is regulated by the body, increasing in individuals that are iron deficient and decreasing in cases of iron overload.

Haem iron from soy leghemoglobin is expected to have similar bioavailability to haem iron from mammalian haem proteins (e.g. myoglobin present in muscle tissue). Data provided in the application indicates that soy leghemoglobin has similar structural and physicochemical properties to animal myoglobins. Soy leghemoglobin is completely digested by pepsin thus making the haem group freely available for absorption. An *in vitro* study found similar bioavailability of haem iron from soy leghemoglobin and bovine haemoglobin. However, in the absence of *in vivo* studies, a quantitative comparison of haem iron bioavailability from soy legehemoglobin and other haem proteins is not possible.

The absence of meat proteins in the proposed meat analogue product may decrease haem iron bioavailability. However, because the proposed meat analogue products have a higher total iron content relative to comparison meat products, FSANZ considers that meat analogue products containing soy leghemoglobin do not present a nutritional disadvantage to consumers in Australia and New Zealand.

## 2.6 Dietary intake assessment

### 2.6.1 Approach to estimating intakes of soy leghemoglobin preparation and iron

Dietary intake assessments require concentration data for the chemical of interest in food, and consumption data for those foods collected through national nutrition surveys.

Intakes were estimated in two ways. Firstly using the maximum concentration of 0.8% soy leghemoglobin requested by the applicant, and secondly using a more refined estimate based on the likely use levels of soy leghemoglobin (0.45% for beef analogue products and 0.25% for pork analogue products). Additionally, as soy leghemoglobin is a source of iron, potential increases in population iron intakes from consumption of meat analogue products containing soy leghemoglobin have also been estimated.

The data used in this assessment include:

* food consumption data from the available Australian and New Zealand national nutrition surveys (Australian Bureau of Statistics [ABS], 2014; Ministry of Health [MoH] 2005; MoH 2012)
* published (baseline) iron intakes derived from the Australian and New Zealand national nutrition surveys (ABS 2015a; ABS 2015b; MoH 2003; University of Otago and MoH 2011)
* iron concentrations in foods from the food composition datasets from the Australian and New Zealand national nutrition surveys (FSANZ 2016a; MoH 2005; MoH 2012).

Further details of FSANZ’s approach to conducting the dietary intake assessment are at [Appendix 1](#_Appendix_1:_FSANZ). This includes details of the:

* food consumption data used in the assessment ([A1.1](#_A1.1_Food_consumption))
* concentration data used in the assessment ([A1.2](#_A1.2_Concentration_data))
* assumptions and limitations of the dietary intake assessment ([A1.3](#_A1.3_Assumptions_and)).

A detailed discussion of the FSANZ methodology and approach to conducting dietary intake assessments is set out in *Principles and Practices of Dietary Exposure Assessments for Food Regulatory Purposes* (FSANZ, 2009).

### 2.6.2 Estimated intakes of soy leghemoglobin preparation and iron

The applicant has requested permission to add soy leghemoglobin in the form of the Preparation to meat analogue products at levels of not more that 0.8% soy leghemoglobin. The applicant has stated that the Preparation will be added to a meat analogue product at levels needed to recreate the nutrition (source of iron), flavour and aroma of meat, for example 0.45% soy leghemoglobin for beef analogue products and 0.25% soy leghemoglobin for pork analogue products.

There are no existing permissions for use of the Preparation and soy leghemoglobin in foods on the market in Australia and New Zealand, therefore the estimated intakes of the Preparation are based only on intake from meat analogue products containing soy leghemoglobin at the levels proposed in the application. As there are existing sources of iron currently in the food supply, the estimated intakes for iron take into account the baseline intakes of iron from existing sources (including foods, beverages and supplements), as well as the amount that could be contributed by foods containing soy leghemoglobin at the levels proposed in the application.

It was assumed for the dietary intake assessment that consumers may choose to eat meat analogue products containing soy leghemoglobin in the same amounts they currently consume minced meat and poultry products, and vegetarian meat alternatives. The foods included in each of these food groups, which have been used to derive the consumption amounts for the assessment, are detailed in [Appendix 1: Table A1.1](#TableA1_1). The mean and 90th percentile consumption amounts of these food groups, based on the most recent national nutrition survey data for Australia and New Zealand, are listed in [Appendix 1](#_Appendix_1:_FSANZ) ([Table A1.2](#TableA1_2) and [Table A1.3](#TableA1_3)).

FSANZ conducted the dietary intake assessment of the Preparation and iron using two scenarios:

* ‘*Maximum proposed use level’*: all meat analogue products consumed contain the maximum proposed concentration of soy leghemoglobin at 0.8%.
* ‘*Likely use level’*: beef, lamb and unspecified meat analogue products contain 0.45% soy leghemoglobin, and pork and poultry analogue products contain 0.25% soy leghemoglobin.

The estimated intakes of the Preparation and iron based on the *‘likely use level’* scenario more accurately reflect estimates of dietary intake over a long period of time or lifetime, reflecting chronic dietary exposure.

The population groups assessed are listed in Table 4 and Table 5. In regards to the Preparation, the toxicological assessment did not identify any population sub-groups for which there were specific safety considerations. Therefore the dietary intake assessment for the Preparation was conducted for the general Australian and New Zealand populations based on the dietary survey data available. The population groups used for the iron intake assessment align with the baseline published iron intake data available for Australia and New Zealand. For Australia, the population groups used for the dietary intake assessment are the same as the Nutrient Reference Value (NRV) [[11]](#footnote-12) age groups. The New Zealand survey age groups and the NRV age groups do not match because results from the New Zealand surveys were not reported exactly according to the NRV age group cut offs.

*Table 4:* Population groups used in the dietary intake assessment for the soy leghemoglobin preparation

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| ***Country***  | ***Nutrition survey***  | ***Age group***  | ***No. respondents (Day 1 only)***  | ***No. respondents (Day 1 and 2)***  |
| **Australia**  | 2011-12 NNPAS  | 2 years and above | Not used | 7735  |
| **New Zealand**  | 2002 NZ CNS  | 5 – 14 years  | 3275  | n/a  |
| 2008 NZ ANS | 15 years and above  | 4721  | n/a  |

*Table 5:* Population groups used in the dietary intake assessment for iron

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| ***Country***  | ***Survey***  | ***Population surveyed***  | ***Age groups analysed*** | ***Sex groups analysed*** |
| **Australia**  | 2011–12 NNPAS  | 2 years and above  | 2–3 years 4–8 years 9–13 years 14–18 years 19–30 years 31–50 years 51–70 years 71 years and over  | Male and Female |
| **New Zealand** | 2002 NZ CNS | 5-14 years | 5-6 years 7-10 years 11-14 years  | Male and Female |
| 2008 NZ ANS | 15 years and above | 15–18 years 19–30 years 31–50 years 51–70 years 71 years and over | Male and Female |

For the dietary intake assessment of iron, FSANZ first compared the baseline intakes of iron from food and beverages for meat-eaters with that of non-meat eaters in order to determine whether non-meat eaters should be considered as a separate population group ([Appendix 2: Table A2.1](#TableA2_1)). For the following reasons it was determined that it was not necessary to consider non-meat eaters as a separate population group to meat-eaters:

* the baseline mean iron intake of non-meat eaters was below that of meat eaters, and the purpose of estimating iron intakes was to consider potential increases in population iron intakes and whether these were above safe levels;
* the two scenarios used for the assessment are conservative and considered protective for estimating the iron intakes for meat eaters and non-meat eaters alike.

#### Estimated intake of the soy leghemoglobin preparation

Dietary intake assessment results were calculated based on ‘consumers only’, that is, those people in the national nutrition survey who reported consuming minced meat and poultry products, and/or vegetarian meat alternatives. Population statistics (mean and 90th percentile) were derived from each individual’s intake of soy leghemoglobin then converted to the corresponding amount of the Preparation, based on the commercial Preparation containing up to 9% soy leghemoglobin, and having a TOS content of 22%. The TOS encompasses the protein component and other organic material derived from the protein source and manufacturing process.

The estimated intake of the Preparation for the Australian and New Zealand population is shown in Table 6.

*Table 6:* Estimated dietary intake of the soy leghemoglobin preparation for Australia and New Zealand

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| ***Country***  | ***Population (consumers only)*** | ***Mean intake*** ***(g/day TOS)*** | ***P90 intake******(g/day TOS)*** | ***Mean intake******(mg/kg bw/day TOS)*** | ***P90 intake******(mg/kg bw/day TOS)*** |
|  | ***Maximum proposed use level*** | ***Likely use level*** | ***Maximum proposed use level*** | ***Likely use level*** | ***Maximum proposed use level*** | ***Likely use level*** | ***Maximum proposed use level*** | ***Likely use level*** |
| **Australia**\* | 2 years and above | 1.2 | 0.7 | 2.7 | 1.5 | 20.4 | 11.0 | 45.4 | 24.3 |
| **New Zealand^** | 5-14 years | 2.2 | 1.1 | 4.3 | 2.4 | 59.9 | 31.8 | 123.6 | 67.5 |
| 15 years and above | 2.3 | 1.2 | 5.0 | 2.8 | 29.3 | 15.5 | 65.1 | 35.3 |

\* 2011-12 Australian National Nutrition and Physical Activity Survey. Based on consumption data from consumers with two days of data only.

**^** 2002 New Zealand National Children’s Nutrition Survey and the 2008–09 New Zealand Adult Nutrition Survey. Based on day 1 consumption data for consumers only.

#### Iron content of meat analogue products containing soy leghemoglobin

Soy leghemoglobin is a source of iron which, if approved, will be added as an ingredient in meat analogue products that may already contain naturally occurring iron from other ingredients. The total amount of iron in a representative meat analogue product (i.e. the Impossible Burger) containing the likely use level of 0.45% soy leghemoglobin was provided in Table D.3-1, Section D of the application. In this same table the applicant compared the average amount of iron in the Impossible Burger to the typical amount of iron in raw minced meat and unprepared veggie or soy burgers, based on composition data from the USDA Food Composition Databases. Table 7 shows the amount of iron in the Impossible Burger and in minced meat and poultry products and other vegetarian meat alternatives, based on Australian and New Zealand food composition data (FSANZ 2016a; New Zealand Institute for Plant & Food Research Limited [PFR] and MoH 2019). As the amount of haem and non-haem iron contributing to the total iron content is not available in the food composition databases for Australia and New Zealand only total iron content is included in the table. The bioavailability of haem and non-haem iron is discussed within this document in section [2.5 Nutritional impact](#_2.5_Nutritional_impact).

*Table 7:* Amount of iron in meat analogue products containing soy leghemoglobin compared to minced meat and poultry products and other vegetarian meat alternatives

|  |  |  |
| --- | --- | --- |
| ***Country*** | ***Food*** | ***Iron content*** ***(mg/100g)*** |
| **Not applicable** | Impossible Burger with 0.45% soy leghemoglobin (raw form)\* | 3.7 |
| Impossible Burger with 0.8% soy leghemoglobin (raw form)\*\* | 4.9 |
| **Australia^** | Beef, mince, >10% fat, raw | 1.45 |
| Lamb, mince, raw | 1.77 |
| Pork, mince, raw | 0.93 |
| Chicken, mince, raw | 0.7 |
| Sausage, vegetarian style, added Fe, Zn and vitamin B12, raw | 3.5 |
| Sausage, vegetarian style, unfortified, raw |  2.5 |
| Tofu (soy bean curd), firm, as purchased | 2.9 |
| **New Zealand∑** | [Beef mince, standard, 10-20% fat, raw](https://www.foodcomposition.co.nz/search/food/M1233) | 1.96 |
| [Lamb, mince, standard, raw](https://www.foodcomposition.co.nz/search/food/M1102) | 1.04 |
| [Pork, mince, raw](https://www.foodcomposition.co.nz/search/food/M537) | 2.05 |
| [Chicken, mince, raw, premium, Tegal](https://www.foodcomposition.co.nz/search/food/M1152) | 0.32 |
| [Tofu, soy bean curd, regular, firm, raw](https://www.foodcomposition.co.nz/search/food/X1151) | 2.00 |
| Tempeh | 2.70 |

\*The expected amount of iron from soy leghemoglobin at use level 0.45% (see [A1.2](#_A1.2_Concentration_data)) in addition to the average amount of non-haem iron in 100 g of the Impossible Burger (2.1 mg) calculated from the non-haem iron content reported in Table E.1.1-1, Section E of the application.

\*\* The expected amount of iron from soy leghemoglobin at the proposed maximum level of 0.8% soy leghemoglobin (see [A1.2](#_A1.2_Concentration_data)) in addition to the average amount of non-haem iron in 100 g of the Impossible Burger calculated from the non-haem iron content reported in Table E.1.1-1, Section E of the application.

^ AUSNUT 2011-13 Food Nutrient Database (FSANZ 2016a).

∑ New Zealand Food Composition Database (PFR and MoH 2019).

#### Estimated intakes of iron with contribution from soy leghemoglobin

The estimated mean and high intakes of iron with the additional contribution from soy leghemoglobin are summarised in Table 8 (Australia) and Table 9 (New Zealand). The food composition datasets from the Australian and New Zealand national nutrition surveys do not separate the amount of haem and non-haem iron in food and beverages, therefore only total iron intakes were estimated.

The estimated amount of additional iron contributed by soy leghemoglobin (mean and 90th percentile) was derived from each individual’s intake of soy leghemoglobin then converted to the corresponding amount of iron, based on 0.35% of the mass of the protein being iron (see [Appendix 1: Section A1.2](#_A1.2_Concentration_data)). The range of meat analogue products that would contain soy leghemoglobin, should the permission to add it be approved is unknown, therefore it was assumed that the amount of iron contributed by ingredients other than the soy leghemoglobin would be within the range of iron in minced meat and poultry products, and vegetarian meat alternative products currently on the market in Australia and New Zealand. In reality there would be variation in the actual amount of iron contributed by other plant-based ingredients in products containing soy leghemoglobin, however, this was unable to be taken into account for this assessment.

Therefore to estimate the total iron intake, the contribution of iron from soy leghemoglobin was added to baseline mean and high usual intakes of iron from food and beverages (ABS 2015a; MoH 2003; University of Otago and MoH 2011) and an additional amount from dietary supplements (e.g. vitamin and mineral tablets) (ABS 2015b). For the high usual intakes of iron from food and beverages, the highest reported percentile from the published usual intakes of iron from food and beverages were used for each country: 95th percentile intakes for the Australian population and 90th percentile intakes for the New Zealand population. The sources of iron that were combined to derive the estimated total intakes are shown in full in [Appendix 2](#_Appendix_2:_Estimated) ([Table A2.2](#TableA2_2) and [Table A2.3](#TableA2_3)).

*Table 8:* Estimated dietary intake of iron (mg/day) for the Australian population with additional contribution from soy leghemoglobin

|  |  |  |  |
| --- | --- | --- | --- |
| *Age group (years)* | *UL\**  | *Mean total intake¥* | *High total intakeΩ* |
|  |  | ***Maximum proposed use level*** | ***Likely use level*** | ***Maximum proposed use level*** | ***Likely use level*** |
| *Males* |  |
| 2-3 | **20** | 12 | 12 | 17 | 16 |
| 4-8 | **40** | 13 | 13 | 19 | 18 |
| 9-13 | **40** | 18 | 17 | 25 | 23 |
| 14-18 | **45** | 20 | 18 | 28 | 26 |
| 19-30 | **45** | 23 | 22 | 32 | 30 |
| 31-50 | **45** | 22 | 21 | 31 | 29 |
| 51-70 | **45** | 18 | 17 | 27 | 25 |
| 71 and over | **45** | 20 | 19 | 28 | 26 |
| *Females* |  |
| 2-3 | **20** | 11 | 11 | 16 | 14 |
| 4-8 | **40** | 12 | 12 | 17 | 16 |
| 9-13 | **40** | 15 | 15 | 22 | 21 |
| 14-18 | **45** | 20 | 20 | 27 | 25 |
| 19-30 | **45** | 23 | 22 | 29 | 28 |
| 31-50 | **45** | 19 | 18 | 25 | 24 |
| 51-70 | **45** | 17 | 16 | 24 | 22 |
| 71 and over | **45** | 16 | 16 | 23 | 22 |

\*Upper Level of intake for iron based on the Nutrient Reference Values for Australia and New Zealand (NHMRC and MoH 2006).

¥Sum of mean intake from food and beverages, dietary supplements and soy leghemoglobin (ABS 2015a; ABS 2015b).

ΩSum of high intake from food and beverages, mean intake from dietary supplements and high intake from soy leghemoglobin (ABS 2015a; ABS 2015b).

*Table 9:* Estimated dietary intake of iron (mg/day) for the New Zealand population with additional contribution from soy leghemoglobin

|  |  |  |  |
| --- | --- | --- | --- |
| *Age group (years)* | *UL\** | *Mean total intake ¥*  | *High total intake Ω*  |
|  |  | ***Maximum proposed use level*** | ***Likely use level*** | ***Maximum proposed use level*** | ***Likely use level*** |
| *Males* |  |  |
| 5-6 | **40** | 16 | 15 | 21 | 19 |
| 7-10 | **40** | 20 | 18 | 26 | 24 |
| 11-14 | **40-45\*\*** | 22 | 21 | 31 | 27 |
| 15-18 | **45** | 22 | 20 | 31 | 27 |
| 19-30 | **45** | 27 | 25 | 39 | 34 |
| 31-50 | **45** | 25 | 23 | 35 | 31 |
| 51-70 | **45** | 20 | 19 | 31 | 27 |
| 71 and over | **45** | 20 | 19 | 27 | 25 |
| *Females* |  |  |
| 5-6 | **40** | 14 | 13 | 18 | 16 |
| 7-10 | **40** | 18 | 16 | 23 | 21 |
| 11-14 | **40-45\*\*** | 23 | 22 | 30 | 28 |
| 15-18 | **45** | 23 | 21 | 29 | 26 |
| 19-30 | **45** | 25 | 24 | 31 | 28 |
| 31-50 | **45** | 21 | 20 | 28 | 25 |
| 51-70 | **45** | 18 | 17 | 24 | 21 |
| 71 and over | **45** | 18 | 17 | 24 | 22 |

\* Upper Level of intake for iron based on the Nutrient Reference Values for Australia and New Zealand (NHMRC and MoH 2006).

\*\*The UL for 9-13 years olds is 40 mg/day. The UL for 14 year olds is 45 mg/day.

¥Sum of mean intake from food and beverages, dietary supplements and soy leghemoglobin (ABS 2015b; MoH 2003; University of Otago and MoH 2011).

ΩSum of high intake from food and beverages, mean intake from dietary supplements and high intake from soy leghemoglobin (ABS 2015b; MoH 2003; University of Otago and MoH 2011).

### 2.6.3 Risk characterisation

No hazard was identified in a repeated dose oral toxicity study in rats at doses of up to 1421 mg/kg bw/day of the Preparation TOS, the highest dose tested. Mean and P90 estimated dietary intakes of the Preparation at the maximum proposed use level were 20 – 60 mg/kg bw/day TOS and 45 – 124 mg/kg bw/day TOS, respectively. Mean and P90 estimated dietary intakes of the Preparation at the likely use level were 11 – 32 mg/kg bw/day TOS and 24 – 68 mg/kg bw/day TOS, respectively. The estimated intakes of the Preparation for both scenarios are considered to be conservative and over-estimate consumption as it is unlikely that consumers will eat meat analogue products containing soy leghemoglobin in the same amounts or with the same frequency they currently consume minced meat and poultry products, and vegetarian meat alternatives (particularly over a long period of time).

The margins of exposure (MOEs) between the NOAEL of 1421 mg/kg bw/day TOS in the rat oral toxicity study and estimated dietary intakes at the maximum proposed use level ranged between 20 – 70 for mean intakes and between 10 – 30 at the 90th percentile. At likely use levels, MOEs for mean and P90 estimated dietary intakes ranged between 40 – 130 and 20 – 60, respectively. These MOEs are not considered to be of concern given that: a sufficient body of knowledge exists on the safety of the source organism (it is not pathogenic or toxigenic), the soy leghemoglobin and *Pichia* proteins will be digested like other dietary proteins and do not share any significant similarities to known allergens or toxins; and the conservative nature of the dietary intake assessment which is likely to overestimate intakes over a long period of time.

As shown in Table 7, the total amount of iron in a representative meat analogue product containing soy leghemoglobin at 0.8% is higher than the amount of iron in minced meat and poultry and other vegetarian meat alternatives on the market in Australia and New Zealand. The amount of iron in the Impossible Burger (3.7 mg/100 g) at the likely use level of 0.45% is similar to the amount of iron in vegetarian style sausages with added iron (3.5 mg/100 g).

The Upper Level of Intake (UL) is a NRV that defines the highest average daily nutrient intake level likely to pose no adverse health effects to almost all individuals in the general population. As intake increases above the UL, the potential risk of adverse effects increases (NHMRC and MoH 2006). At the current baseline levels of iron intake from food and beverages for the Australian and New Zealand populations there are no age/sex groups which exceed the UL for iron at the highest reported percentile of iron intake. With the addition of iron from soy leghemoglobin to baseline iron intakes from food and beverages, and from dietary supplements, the estimated high intake of iron within all population age/sex groups for both Australia and New Zealand were still below the respective ULs for both scenarios. The estimated iron intakes are considered to be conservative and an overestimation of actual iron intakes for the following reasons:

* While the proportion of Australians with an intake of iron from dietary supplements was less than 14% in the most recent survey (ABS 2015b), the mean intake of iron from supplements from this proportion of consumers was used to derive the estimated mean and high intakes for each age/sex group, and they are not likely to be consumed every day over a lifetime.
* It is unlikely that consumers will eat meat analogue products containing soy leghemoglobin in the same amounts they currently consume minced meat and poultry products, and other vegetarian meat alternatives, particularly over a long period of time or a lifetime.

As the estimated iron intakes fall below the UL for iron for each age/sex group, for both the *maximum proposed use level* and *likely use level* scenarios, it is unlikely that consumption of meat analogue products containing soy leghemoglobin would pose a risk of iron exceedances to the Australian and New Zealand populations.

## 2.7 Manufacturing process

### 2.7.1 Manufacturing process of the soy leghemoglobin preparation

The Preparation is manufactured in compliance with cGMP. A schematic overview of the manufacturing process is presented in Figure 3.



***Figure 3*** *Schematic overview of the manufacturing process for the soy leghemoglobin preparation*

The production strain MXY0541 is cultured to high biomass in a submerged fed-batch fermentation system. At the end of the fermentation run, the *Pichia* cells are lysed by mechanical shearing or high-pressure homogenisation. The insoluble and soluble fraction of the cells are separated by centrifugation, followed by several steps to remove impurities and to ensure no viable cells remain in the lysate. The lysate is then concentrated, mixed with stabilisers (for example sodium ascorbate and sodium chloride), heat treated and stored at -20°C.

FSANZ supports that the raw materials and processing aids employed in the production of the Preparation are standard ingredients used in the food/enzyme industry and are suitable for their intended use. Impossible Foods have demonstrated that they have in-process controls in place, under cGMP, to ensure the purity and consistency of the final preparation. the Preparation is well characterized and meets food-grade specifications, which include acceptable limits on compositional criteria and impurities such as heavy metals and microbiological contaminants.

### 2.7.2 Analytical methods for detection

Methods for measuring proximate levels in the the Preparation follow standard methodologies from the AOAC, although internal methods were developed for determining the percentage of and purity of the soy leghemoglobin and using these values. Details of the internal methods were provided as confidential information. Levels of metal impurities are measured following standard methodologies from the US EPA. Microbial presence was determined by standard methods from the AOAC.

### 2.7.3 Characterisation of the production strain MXY0541

### Safety of host and donor organisms

#### Host organism

The host organism is a commercial strain of *Pichia pastoris*, known as Bg11 A detailed discussion of the history of use of *Pichia* has been addressed in [Section 2.2](#_2.2_History_of).

#### Donor organisms

*Glycine max*

The gene for leghemoglobin was based on the leghemoglobin C2 protein sequence found in soybean (*Glycine max*).

Soy is known as one of the major food allergens, due to the presence of a range of proteins in the mature seed that have been associated with anti-nutrients and allergenicity (OECD 2012). Due to this allergenicity, many soy products require allergen labelling in Australia and New Zealand. Although soybean is known to express allergenic proteins, there is no evidence to suggest that soy leghemoglobin is an allergenic protein, especially as haem-containing proteins are found in common food sources, especially red meat. A more detailed assessment of allergenicity concerns are addressed in [Section 2.3.2](#_2.3.2_Safety_of).

*Pichia pastoris*

Several endogenous genes and regulatory elements were re-introduced into the host organism, *Pichia pastoris*. The genes include Mxr1 and the enzymes involved in the haem biosynthesis pathway. Mxr1, a transcription factor, drives expression of genes associated with the AOX1 promoter (pAOX1). The regulatory elements include the promoters and terminators from AOX1, AOX2 and DAS2, which are commonly used for the expression of recombinant proteins in *Pichia*. A detailed discussion of the history of use of *Pichia* has been addressed in [Section 2.2](#_2.2_History_of).

### Nature and characterisation of the genetic modification

Characterisation of the genetic modification is necessary to provide an understanding of the genetic material introduced into the host genome and helps to frame the subsequent parts of the safety assessment. This characterisation addresses three main aspects:

* A detailed description of the DNA sequences introduced to the host genome and the method used to transform the host
* a characterisation of the inserted DNA, including any rearrangements that may have occurred as a consequence of the transformation
* the genetic stability of the inserted DNA and any accompanying expressed traits.

### Expression of leghemoglobin from *Glycine max* in *Pichia pastoris*

In order to express the leghemoglobin C2 protein in *P. pastoris*, three expression cassettes were developed. Each expression cassette was designed to target three distinct loci in the *Pichia* genome using homologous recombination. Each expression cassette contained the leghemoglobin gene flanked by regulatory elements belonging to the targeted gene.

The gene for leghemoglobin was chemically synthesised based on the sequence from soybean (*G. max*). Chemical synthesis was used to enable codon use to be optimised for expression of the plant protein in yeast. The primary amino acid sequence of the expressed protein is the same as that found in soybean ([P02236](https://www.uniprot.org/uniprot/P02236)[[12]](#footnote-13)).

The expression cassettes were introduced separately into the host using electroporation of a linearised fragment containing just the expression cassette sequences. As there was no selection marker present within the leghemoglobin expression cassettes, co-transformation was performed using a plasmid containing an antibiotic resistance gene. After selection of transformants, plasmid curing was used to remove the antibiotic resistance plasmid.

To identify positive transformants, PCR analysis was used as an initial screening tool. When the final production strain MXY0541 was chosen, whole genome analysis using short-read DNA sequencing was performed using a Pac-Bio system. The production organism MXY0541 contains all three leghemoglobin transformation events. The production organism MXY0291 that was granted GRAS status in the US (US FDA 2018; GRN737) contains only one of the transformation events.

***Locus 1:*** FSANZ’s analysis of a range of data from MXY0541 indicates:

1. inserts containing the leghemoglobin gene expression cassette are present at the insertion locus. The inserts contain multiple tandem repeats of the expression cassette
2. the sequence for the leghemoglobin gene is as expected, with no mutations or rearrangements
3. the flanking regions on either side of the inserted gene contain the expected regulatory sequences.

***Locus 2:*** FSANZ’s analysis of the PCR and sequencing data from MXY0541 indicates:

1. several tandem repeats of the expression cassette at the insertion locus
2. the gene copy number cannot be fully determined. The actual copy number could not be completely delineated using whole genome sequencing because of the repetitive nature of the sequence in the tandem repeats and the short-read DNA sequencing employed
3. the insert has replaced the target gene of locus 2
4. the sequence located at the 5’ end of the gene contains the expected regulatory sequences.

FSANZ further identified that the sequence located at the 3’ end of the gene includes the terminator sequence of the target gene for locus 2 but this sequence is in the complement direction and may not encode a functional polyadenylation site. This could reduce the stability and transport of the mRNA transcript and thus translation efficiency, but would not impact safety.

FSANZ also identified base pair differences in some of the leghemoglobin gene inserts, where the sequence data was of high quality. These differences could result in amino acid changes and expression of truncated proteins. However, as already noted, it is unlikely these proteins would be translated because of the non-functional polyadenylation sequence. These changes would therefore not impact safety.

***Locus 3***: FSANZ’s analysis of the PCR and sequencing data from MXY0541 indicates:

1. there are several tandem repeats of the expression cassette
2. the sequence for leghemoglobin gene is as expected, with no mutations or rearrangements
3. the sequence located at the 3’ end of the gene contains the expected regulatory sequences
4. the target gene at locus 3 has not been disrupted.

FSANZ further identified that the sequence located at the 5’ end of the leghemoglobin gene did not contain the expected promoter for the target gene. Instead, this region contained the promoter sequence from another gene located upstream of the locus in the *Pichia* genome and included a partial sequence of the neighbouring protein coding gene and terminator sequence. This would explain why the target gene at locus 3 was not disrupted, as the flanking sequences in this expression cassette would not have allowed for targeted homologous recombination. The insertion was also not located at the locus but instead, preceded the gene from which the alternate promoter was obtained. However, the expression cassette that was inserted contains a functional promoter and terminator, flanking a gene containing no mutations and the insertion does not interrupt any other genes.

### Overexpression of the endogenous haem co-factor

Haem is an important co-factor required for the oxygen-transporting role of leghemoglobin and provides the dietary source of iron. It is also a required co-factor for the function of several critical haem-proteins in yeast thus haem could become a limiting factor in the production of recombinant leghemoglobin (Franken et al. 2011; Krainer et al. 2015). To solve this potential issue, the applicant chose to increase the level of endogenous haem by introducing the genes encoding the eight enzymes used to synthesis haem. The enzymes have been well characterised and the biosynthesis pathway is the same in yeast, plants and animals (reviewed in Layer et al. 2010; Franken et al. 2011).

Expression cassettes were generated to introduce haem enzyme genes, targeted to a single locus. The haeme enzyme genes and associated regulatory sequences were all sourced from the host organism. The expression cassettes also contained antibiotic resistance genes sourced from non-host organisms, flanked by Cre-Lox sequences (Dale and Ow 1991). The expression cassettes were transformed into the host as linearised fragments of DNA, free of vector backbone sequences, using electroporation. After selection of positive transformants, the antibiotic resistance genes were excised from the transformed cells using the Cre-Lox system.

Characterisation of the inserted haem synthesis pathway genes were performed by PCR and whole genome sequencing. The characterisation identified that at least one functional copy of every enzyme was inserted at the locus. For all enzymes, there is at least one copy where the DNA sequences are as expected.

During the initial PCR screen, it was observed that one of the enzyme genes existed in both full and truncated forms. A subsequent expression cassette was designed to replace the truncated versions but the reinsertion of the expression cassette did not repair the truncated gene variants. Furthermore, some mutations were created in the recombination event. FSANZ noted that the replacement expression cassette only contained a promoter sequence and gene coding region with no terminator sequence thus it would be highly unlikely any transcripts would be stable or translated. In addition, peptide mapping focusing on the specific enzyme product identified only the full length protein with the expected peptide sequence, confirming that the product is as expected. A full insert of this enzyme was achieved in the original transformation event targeting the locus.

### Overexpression of the endogenous transcription factor Mxr1

A common regulatory element used for the expression of recombinant proteins in *Pichia* is the promoter for the alcohol oxidase gene (pAOX1). This promoter was used for several of the recombinant haem enzyme genes and for one of the expression cassettes for leghemoglobin. To ensure sufficient activation of all the genes driven by pAOX1, the applicant chose to increase expression of the main transcription factor for pAOX1, known as Mxr1 (Lin-Cereghino et al. 2006).

A single expression cassette was generated, containing the Mxr1 gene from the host, flanked by the *Pichia* AOX1 promoter and terminator sequences. FSANZ noted that in GRN737 (US FDA 2018) the terminator sequence was declared as tFDH, however from the DNA sequence provided to FSANZ, we can confirm that the terminator sequence is tAOX1. As there were no selective marker genes included in the expression cassette, a plasmid containing an antibiotic resistance gene was co-transformed. After selection of positive transformants, plasmid curing was used to generate transformants no longer containing the antibiotic resistance plasmid.

The applicant noted that due to the cloning strategy employed for Mxr1, six extra N-terminal amino acids are encoded in the final protein. Analysis of the final leghemoglobin product using mass spectrometry showed that the protein engineered Mxr1 protein was not present.

Characterisation of the inserted Mxr1 insert confirmed a single copy is present in the locus and there are no mutations or rearrangements.

***Absence of the selective marker genes:*** To confirm absence of the antibiotic resistance genes used to select positive transformants, which were subsequently removed by either cre-lox excision or plasmid curing, results from three methods were provided: growth on selective agar; PCR targeting the antibiotic resistance gene; and whole genome sequencing. Results obtained from the final production strain MXY0541 showed absence of all the antibiotic resistance genes used. These analyses confirmed that both the curing and cre-lox excision processes were successful.

***Inheritance and genetic stability of the inserted DNA:*** Genetic stability and inheritance of the introduced genes was confirmed by analysis of cultures at the end of a standard fermentation run (150-200 generations). PCR analysis targeting all the inserted genes showed that all the inserts were present and at the expected sizes, confirming inheritance and stability.

***Presence of novel DNA in the final product:*** A genomic DNA preparation of the final leghemoglobin preparation was performed and quantified using NanoDrop spectrophotometry. The data shows that the amount of DNA present in the the Preparation is approximately 300 mg/l. As the amount of the Preparation added to the burgers will range from 5-10 ml, the amount of DNA would be approximately 1.5-3 mg (i.e. 0.3mg/ml). This DNA will contain the novel DNA that encodes the recombinant soy leghemoglobin.

### 2.7.4 Conclusion

Several expression cassettes were generated to introduce ten genes into *P. pastoris* in order to express soy leghemoglobin. The data provided by the applicant demonstrated that for every gene introduced, there is at least one full copy of the insert that has been integrated into the host. In some situations, more than one copy of a gene has been inserted. In one situation, a truncated version of a single gene has been inserted multiple times. All the gene inserts were shown to be stably integrated over several generations. Data was also provided confirming successful removal of the antibiotic selection markers used during the transformant process. The conclusion from this assessment is the production strain MXY0541 poses no safety concerns.

# 3 Risk assessment summary and conclusion

FSANZ has conducted a comprehensive assessment following the internationally recognised risk analysis framework. The assessment was based on a weight of evidence approach, combining information and scientific evidence provided by the applicant with independent sources.

In conducting the risk assessment of the soy leghemoglobin and the Preparation, a number of criteria have been addressed, including the safety of the *P. pastoris* host strain, novel proteins, toxicity of the the Preparation and a nutritional and dietary intake assessment. The safety assessment of the source organism and novel proteins concluded there were no public health and safety concerns. The source organism is a well characterised yeast with a recognised safe history of use for the production of food enzymes. It is neither pathogenic nor toxigenic.

The novel soy leghemoglobin was shown to be equivalent to that expressed in soybean and was shown to be expressed as a holoprotein. Analyses of the potential allergenicity or toxicity of all the novel proteins, including soy leghemoglobin and the *Pichia* proteins, did not identify any significant similarities to known allergens or toxins. The proteins were shown to be susceptible to pepsin digestion and were denatured at standard cooking temperatures and in acidic conditions that mimic the stomach environment. The shelf life and specifications of the the Preparation are also appropriate for addition to meat analogue products.

The applicant submitted *in vitro* genotoxicity studies in bacterial and mammalian cells and an oral toxicity study in rats. These studies are intended to confirm the outcome of the compositional and bioinformatic analysis conducted as a part of the safety assessment. No hazard was identified in the submitted studies. The Preparation was not genotoxic *in vitro* and did not cause adverse effects in short-term toxicity studies in rats. The NOAEL of freeze-dried Preparation in a 28-day dietary toxicity study in rats was 1536 mg/kg bw/day, the highest dose tested. This dose corresponds to 1421 mg/kg bw/day TOS.

Mean and P90 estimated dietary intakes of the Preparation at the maximum proposed use level were 20 – 60 mg/kg bw/day TOS and 45 – 124 mg/kg bw/day TOS, respectively. Mean and P90 estimated dietary intakes of the Preparation at the likely use level were 11 – 32 mg/kg bw/day TOS and 24 – 68 mg/kg bw/day TOS, respectively. The estimated intakes of the Preparation for both scenarios are considered to be conservative and over-estimate consumption as it is unlikely that consumers will eat meat analogue products containing soy leghemoglobin in the same amounts or with the same frequency they currently consume minced meat and poultry products, and vegetarian meat alternatives (particularly over a long period of time).

The MOEs between the NOAEL of 1421 mg/kg bw/day TOS in the rat oral toxicity study and estimated dietary intakes at the maximum proposed use level ranged between 20 – 70 for mean intakes and between 10 – 30 at the 90th percentile. At likely use levels, MOEs for mean and P90 estimated dietary intakes ranged between 40 – 130 and 20 – 60, respectively. These MOEs are not considered to be of concern given that: a sufficient body of knowledge exists on the safety of the source organism, the soy leghemoglobin and *Pichia* proteins will be digested like other other dietary proteins and do not share any significant similarities to known allergens or toxins; and the conservative nature of the dietary intake assessment which is likely to overestimate intakes over a long period of time.

The nutrition assessment focused on the relative bioavailability of haem iron from soy leghemoglobin compared to haem iron from animal sources. Haem iron from soy leghemoglobin is expected to have similar bioavailability to haem iron from mammalian haem proteins (e.g. myoglobin present in muscle tissue). The absence of meat proteins in the proposed meat analogue product may decrease haem iron bioavailability. However, because iron absorption is regulated tightly by the body, and the proposed meat analogue products have higher total iron content relative to comparison foods, any decrease in haem iron bioavailability should not result in a nutritional disadvantage to consumers in Australia and New Zealand.

The estimated intakes of iron (with the additional iron contribution from soy leghemoglobin) for all population age/sex groups assessed for both the Australian and New Zealand populations are below the ULs for iron. The estimated iron intakes in FSANZs assessment, for both the *maximum proposed use level* and *likely use level* scenarios, are considered to be conservative and an overestimation of actual iron intakes. It is unlikely that consumption of meat analogue products containing soy leghemoglobin would pose a risk of iron exceedances to the Australian and New Zealand populations, including at levels up to 0.8% soy leghemoglobin.

As of March 2020, the applicant advised they had sold approximately 100,000,000 quarter-pound (113 g) servings of meat analogue products containing soy leghemoglobin. Its post-marketing surveillance has identified one complaint per 600,000 servings based on the current formulation (released on the market in the US in early 2019), but none of these complaints has been confirmed as an adverse event due to consumption of these products.

The assessment of soy leghemoglobin and the Preparation concluded that there are no public health and safety concerns associated with its use in meat analogue products at the proposed level of up to 0.8% soy leghemoglobin.

# 4 References

ABS (2014) National Nutrition and Physical Activity Survey, 2011–12, Basic Confidentialised Unit Record Files (CURF). Australian Bureau of Statistics, Commonwealth of Australia, Canberra

ABS (2015a) Australian Health Survey: Usual Nutrient Intakes, 2011-12. Table 4:Essential minerals (and caffeine). Australian Bureau of Statistics, Commonwealth of Australia, Canberra

ABS (2015b) Australian Health Survey: Nutrition – Supplements, 2011-12. Iron supplement intake. Australian Bureau of Statistics, Commonwealth of Australia, Canberra

Ahmad M, Hirz M, Pichler H, Schwab H (2014) Protein expression in *Pichia pastoris*: recent achievements and perspective for heterologous protein production. Appl Microbiol Biotechnol 98: 5301-5317

Appleby CA (1974) In: The Biology of Nitrogen Fixation (ed) Quispel, A. (North Holland, Amsterdam) pp. 521-554

Appleby CA, Bergersen FJ (1980) Preparation and experimental use of leghaemoglobin. In: Methods for Evaluating Biological Nitrogen Fixation, FJ Bergersen (ed) (Chichester: Wiley) pp. 315-335

Appleby CA (1984) Leghemoglobin and *Rhizobium* respiration. Ann Rev Plant Physiol 35:443-478

Aréchaga-Ocampo E, Saenz-Rivera J, Sarath G, Klucas RV, Arredondo-Peter R (2001) Cloning and

expression analysis of hemoglobin genes from maize (*Zea mays* ssp. mays) and teosinte (*Zea mays* ssp. parviglumis). Biochim Biophys Acta – Gene Structure ad Expression 1522(1): 1-8

Arrendondo-Peter R, Hargrove MS, Moran JF, Sarath G, Klucas RV (1998) Plant hemoglobins. Plant Physiol 118:1121-1125

Au AP and Reddy MB (2000) Caco-2 cells can be used to assess human iron bioavailability from a semipurified meal. J Nutr 130: 1329-1334

Banerjee H, Verma M (2000) Search for a novel killer toxin in yeast *Pichia pastoris*. Plasmid 43(2):181-183

Bothwell TH, Charlton RW, Cook JD, Finch CA (1979) Iron metabolism in man. Blackwell Scientific Publications, Oxford

Björn-Rasmussen E, Hallberg L, Isaksson B, Arvidsson B (1974) Food iron absorption in man applications of the two-pool extrinsic tag method to measure haem and nonhaem iron absorption from the whole diet. J Clin Invest 53(1):247-255

Brady JR, whittaker CA, Tan MC, Kristensen II DL, Ma D, Dalvie NC, Love KR, Love JC (2019) Comparative genome‐scale analysis of Pichia pastoris variants informs selection of an optimal base strain. Biotechnol Bioeng. 2019:1-13 <https://doi.org/10.1002/bit.27209>

Braun-Galleani S, Dias JA, Coughlan AY, Ryan AP, Byrne KP, wolfe KH (2019) Genomic diversity and meiotic recombination among isolates of the biotech yeast *Komagataella phaffii* (*Pichia pastoris*). Microb Cell Fact 18:e211

Carpenter CE and Mahoney AW (1992) Contributions of heme and nonheme iron to human nutrition. Crit Rev Food Sci Nutr 31(4): 333-367

Champagne ET (1988) Effects of pH on mineral-phytate, protein-mineral-phytate, and mineral-fiber interactions Possible consequences of atrophic gastritis on mineral bioavailability from high-fiber foods. J Am Coll Nutr 7(6): 499-508

Cho S, Kang SM, Seong P, Kang G, Choi S, Kwon E, Moon S, Kim D, Park B (2014) Physico-chemical meat qualities of loin and top round beef from Holstein calves with different slaughtering ages. Korean J Food Sci Anim Resour 34(5):674-682

Codex (2009) Foods derived from modern biotechnology 2nd edition. Codex Alimentarius Commission, Rome. [http://www.fao.org/docrep/pdf/011/a1554e/a1554e00.pdf. Accessed 16 October 2019](http://www.fao.org/docrep/pdf/011/a1554e/a1554e00.pdf.%20Accessed%2016%20October%202019)

Cregg JM, Barringer KJ, Hessler AY, Madden R (1985) *Pichia pastoris* as a host system for transformations. Mol Cell Biol 5(12):3376-3385

CSIRO (2006) [Shelf-life testing: methods for determining the claimable life of meat products](https://meatupdate.csiro.au/data/MEAT_TECHNOLOGY_UPDATE_06-2.pdf). MTU 2/06 – April 2006. Accessed 1 November 2019

Dale EC, Ow DW (1991) Gene transfer with subsequent removal of the selection gene from the host genome. Proc Natl Acad Sci USA 88(23): 10558-10562

De Schutter K, Lin YC, Tiels P, Van Hecke A, Glinka S, Weber-Lehmann J, Rouzé P, Van de Peer Y, Callewaert N (2009) Genome sequence of the recombinant protein production host *Pichia pastoris*. Nat Biotechnol 27(6):561-566

Delaney B, Astwood JD, Cunny H, Eichen Conn R, Herouet-Guicheney C, MacIntosh S, Meyer LS, Privalle L, Gao Y, Mattsson J, Levine M, ILSI International Food Biotechnology Committee Task Force on Protein Safety (2008) Evaluation of protein safety in the context of agricultural biotechnology. Food Chem Toxicol 46(S2):S71–S97

EC (2008) Commission Regulation (EC) No. 440/2008. B.10: Mutagenicity – in vitro mammalian chromosome aberration test

EC (2008) Commission Regulation (EC) No. 440/2008. B.13/14: Mutagenicity – reverse mutation test using bacteria

EFSA BIOHAZ Panel (2017) Update of the list of QPS-recommended biological agents intentionally added to food or feed as notified to EFSA 6: suitability of taxonomic units notified to EFSA until March 2017. EFSA Journal 15(7):4884

EFSA BIOHAZ Panel (2018) Update of the list of QPS-recommended biological agents intentionally added to food or feed as notified to EFSA 7: suitability of taxonomic units notified to EFSA until September 2017. EFSA Journal 16(1):5131

Eurofins (2017) *In vitro* mammalian chromosome aberration test in human lymphocytes with soy leghaemoglobin preparation. Eurofins BioPharma Product Testing Munich GmbH, Germany. Study number 160931

França RC, Conceição FR, Mendonça M, Haubert L, Sabadin G, de Oliveira PD, Amaral MG, Silva WP, Moreira ÂN (2015) *Pichia pastoris* X-33 has probiotic properties with remarkable antibacterial activity against *Salmonella Typhimurium*. Appl Microbiol Biotechnol 99(19):7953–7961

Franken ACW, Lokman BC, Ram AFJ, Punt PJ, van den Hondel CAMJJ, de weert S (2011) Heme biosynthesis and its regulation: towards understanding and improvement of heme biosynthesis in filamentous fungi. Appl Microbiol Biotechnol 91:447-460

Fraser RZ, Shitut M, Agrawal P, Mendes O, Klapholz S (2018) Safety Evaluation of Soy Leghemoglobin Protein Preparation Derived From *Pichia pastoris*, Intended for Use as a Flavor Catalyst in Plant-Based Meat. Int J Toxicol 37: 241-262

FSANZ (2009) Principles and practices of dietary exposure assessment for food regulatory purposes. Report prepared by Food Standards Australia New Zealand, Canberra

<http://www.foodstandards.gov.au/publications/Pages/Principles-and-Practices-of-Dietary.aspx>

FSANZ (2016a) AUSNUT 2011-13 – Food Nutrient Database. Australian Government, Canberra.

[www.foodstandards.gov.au/science/monitoringnutrients/ausnut/ausnutdatafiles/Pages/foodnutrient.aspx](http://www.foodstandards.gov.au/science/monitoringnutrients/ausnut/ausnutdatafiles/Pages/foodnutrient.aspx)

FSANZ (2016b) Safe Food Australia 3rd edition. A guide to the Food Safety Standards. Chapter 3 of the Australia New Zealand Food Standards Code (applies to Australia only)

Garrocho-Villegas V, Gopalasubramaniam SK, Arredondo-Peter R (2007) Plant hemoglobins: What we know six decades after their discovery. Gene 398:78-85

Gil de los Santos JR, Storch OB, Fernandes CG, Gil-Turnes C (2012) Evaluation in broilers of the probiotic properties of *Pichia pastoris* and a recombinant *P. pastoris* containing the *Clostridium perfringens* alpha toxin gene. Vet Microbiol 156(3-4):448-451

Gil de los Santos D, Gil de los Santos JR, Gil-Turnes C, Gaboardi G, Silva LF, França R, Fernandes CG, Conceição FR (2018) Probiotic effect of *Pichia pastoris* X-33 produced in parboiled rice effluent and YPD medium on broiler chickens. PLoS ONE 13(2):e0192904

Glahn RP, Lee OA, Yeung A, Goldman MI, Miller DD (1998) Caco-2 cell ferritin formation predicts nonradiolabeled food iron availability in an in vitro digestion/Caco-2 cell culture model. J Nutr 128:1555-1561

Hallberg L, Björn-Rasmussen E, Howard L, Rossander L (1979) Dietary haem iron absorption: a discussion of possible mechanisms for the absorption-promoting effect of meat and for the regulation of iron absorption. Scand J Gastroenterol 14(7):769-779

Hallberg L (1983) Iron requirements and bioavailability of dietary iron. Experientia Suppl 44:223–244

Hallberg L, Rossander L and Skanberg AB (1987) Phytates and the inhibitory effect of bran on iron absorption in man. Am J Clin Nutr 45:988-996

Hallberg L (1998) Combating iron deficiency: daily administration of iron is far superior to weekly administration. Am J Clin Nutr 68(2):213-217

Hargrove, MS, Wilkinson AJ, Olson JS (1996) Structural factors governing hemin dissociation from metmyoglobin. Biochemistry 35(35):11300-11309

Holzhauser T, Wackermann O, Ballmer-Weber BK, Bindslev-Jensen C, Scibilia J, Perono-Garoffo L, Utsami S, Poulsen LK, Vieths S (2009) Soybean (Glycine max) allergy in Europe: Gly m 5 (β-conglycinin) and Gly m 6 (glycinin) are potential diagnostic markers for severe allergic reactions to soy. J Allergy Clin Immunol 123(2):452-458

Hill RD (2012) Non-symbiotic haemoglobins - What's happening beyond nitric oxide scavenging? AoB Plants 2012:pls004

Hill R, Hargrove M, Arredondo-Peter R (2016) Phytoglobin: a novel nomenclature for plant globins accepted by the globin community at the 2014 XVIII conference on Oxygen-Binding and Sensing Proteins. F1000Res 5:212

Hooda J, Shah A, Zhang L (2014) Heme, an essential nutrient from dietary proteins, critically impacts diverse physiological and pathological processes. Nutrients 6(3):1080-1102

Hunt MC, Sorheim O and Slinde E (1999) Color and heat denaturation of myoglobin forms in ground beef. J Food Sci 64(5):847-851

Hurrell R and Egli I (2010) Iron bioavailability and dietary reference values. Am J Clin Nutr 91:1461s–1467s

IFBC (1990). Biotechnologies and food: assuring the safety of foods produced by genetic modification. Regul Toxicol Pharmacol 12(3, Part 2):vii-S196. DOI:10.1016/S0273-2300(05)80069-8

Ito K, Sjölander S, Sato S, Movérare R, Tanaka A, Söderström, Borres M, Poorafshar M, Ebisawa M Lm IgE to Gly m 5 and Gly m 6 is associated with severe allergic reactions to soybean in Japanese children. J Allergy Clin Immunol 128(3):673-675

Jin Y, He X, Andoh-Kumi K, Fraser RZ, Lu M, Goodman RE (2018) Evaluating potential risks of food allergy and toxicity of soy leghemoglobin expressed in *Pichia pastoris*. Mol Nutr Fod Res 62:e1700297

Krainer FW, Capone S, Jäger M, Vogl T, Gerstmann M, Glieder A, Herwig C, Spadiut O (2015) Optimizing cofactor availability for the production of recombinant heme peroxidase in *Pichia pastoris*. Microb Cell Fact 14:e4

Kubo H (1939) Über das hamoprotein aus den wurzelknollchen von leguminosen (About the haemoprotein from the root nodules of legume). Acta Phytochim [Tokyo] 11(1):195-200

Kurtzman CP (2009). Biotechnological strains of *Komagataella* (*Pichia*) *pastoris* are *Komagataella phaffii* as determined from multigene sequence analysis. J Ind Microbiol Biotechnol 36(11):1435-1438

Layer G, Reichelt J, Jahn D, Heinz DW (2010) Structure and function of enzymes in heme biosynthesis. Protein Sci 19:1137-1161

Li YT, Hsieh YL, Henion JD (1993) Studies on heme binding in myoglobin, haemoglobin and cytochrome *c* by ion spray mass spectrometry. J Am Soc Mass Spectrom 4:631-637

Lin-Cereghino GP, Godfrey L, de la Cruz BJ, et al (2006) Mxr1p, a key regulator of the methanol utilization pathway and peroxisomal genes in *Pichia pastoris*. Mol Cell Biol, 26:883–897

Lira-Ruan V, Sarath G, Klucas RV, Arredondo-Peter R (2001) Synthesis of hemoglobins in rice (*Oryza sativa* var. Jackson) plants growing in normal and stress conditions. Plant Sci 161(2): 279-287

Lombardi‐Boccia G, Martinez‐Dominguez B, Aguzzi A (2002) Total haem and non‐haem iron in raw and cooked meats. J Food Sci 67(5):1738-1741

López MA, Martos FC (2004) Iron availability: An updated review. Int J Food Sci Nutr 55(8):597-606

[Lynch SR](https://www.ncbi.nlm.nih.gov/pubmed/?term=Lynch%20SR%5BAuthor%5D&cauthor=true&cauthor_uid=4038429), [Dassenko SA](https://www.ncbi.nlm.nih.gov/pubmed/?term=Dassenko%20SA%5BAuthor%5D&cauthor=true&cauthor_uid=4038429), [Morck TA](https://www.ncbi.nlm.nih.gov/pubmed/?term=Morck%20TA%5BAuthor%5D&cauthor=true&cauthor_uid=4038429), [Beard JL](https://www.ncbi.nlm.nih.gov/pubmed/?term=Beard%20JL%5BAuthor%5D&cauthor=true&cauthor_uid=4038429), [Cook JD](https://www.ncbi.nlm.nih.gov/pubmed/?term=Cook%20JD%5BAuthor%5D&cauthor=true&cauthor_uid=4038429) (1985) Soy protein products and heme iron absorption in humans. Am J Clin Nutr 41(1):13-20

MoH (2003) NZ food NZ children: Key results of the 2002 national children's nutrition survey. Ministry of Health, Wellington

MoH (2005) 2002 National Children's Nutrition Survey: National Confidentialised Unit Record File (CURF). Ministry of Health, Wellington

MoH (2012) 2008-9 New Zealand Adult Nutrition Survey: National Confidentialised Unit Record File (CURF). Ministry of Health, Wellington

NHMRC, MoH (2006) Nutrient reference values for Australia and New Zealand. National Health and Medical Research Council and New Zealand Ministry of Health, Canberra, Australia

OECD (1997) Guidelines for the testing of chemicals, Section 4. Bacterial reverse mutation test. Test Guideline No. 471

OECD (2008) Guidelines for the testing of chemicals, Section 4. Repeated dose 28-day oral toxicity study in rodents. Test Guideline No. 407

OECD (2012) Revised consensus document on compositional considerations for new varieties of soybean [Glycine max (L.) Merr.]: Key food and feed nutrients, anti-nutrients, toxicants and allergens. Series on the Safety of Novel Foods and Feeds No. 25

OECD (2014) Guidelines for the testing of chemicals, Section 4. *In vitro* mammalian chromosomal aberration test. Test Guideline No. 473

Ofori-Anti AO, Ariyarathna H, Chen L, Lee HL, Pramod SN, Goodman RE (2008) Establishing objective detection limits for the pepsin digestion assay used in the assessment of genetically modified foods. Regul Toxicol Pharmacol 52:94-103

Ogawa T, Tsuji H, Bando N, Kitamura K, Zhu YL, Hirano H, Nishikawa K (1993) Identification of the soybean allergenic protein, Gly m Bd 30K, with the soybean seed 34-kDa oil-body-associated protein. Biosci Biotechnol Biochem 57(6):1030-1033

Ordway GA, Garry DJ (2004) Myoglobin: an essential hemoprotein in striated muscle. Journal of Experimental Biology, 207: 3441-3446. doi: 10.1242/jeb.01172

Ott T, van Dongen JT, Günther C, Krusell L, Desbrosses G, Vigeolas H, Bock V, Czechowski T, Geigenberger P, Udvardi MK (2005) Symbiotic leghemoglobins are crucial for nitrogen fixation in legume root nodules but not for general plant growth and development. Curr Biol 15(6):531-535

PFR, MoH (2019) New Zealand Food Composition Database. New Zealand Food Composition Database Online Search. New Zealand Institute for Plant & Food Research Limited and Ministry of Health

New Zealand Food Composition Database 2019. New Zealand Food Composition Database Online Search. The New Zealand Institute for Plant & Food Research Limited and Ministry of Health. https://www.foodcomposition.co.nz/search

Product Safety Labs (2016a) Soy leghemoglobin preparation, purified soy leghemoglobin preparation and bovine erythrocytes: A 14-day dietary toxicity/palatability study in rats. Product Safety Labs, USA. Study number 43167

Product Safety Labs (2016b) Soy leghemoglobin preparation: Bacterial reverse mutation test (Ames test). Product Safety Labs, USA. Study number 42759

Product Safety Labs (2017a) Soy leghemoglobin preparation: A 28-day dietary study in rats. Product Safety Labs, USA. Study number 43166

Product Safety Labs (2017b) Soy leghemoglobin preparation: An investigative 28-day dietary study in rats with a 14-day pre-dosing estrous cycle determination. Product Safety Labs, USA. Study number 44856

Proulx AK, Reddy MB (2006) Iron bioavailability of hemoglobin from soy root nodules using a Caco-2 cell culture model. J Agric Food Chem 54(4):1518-1522

Ross EJ, Shearman L, Mathiesen M, Zhou YJ, Arredondo-Peter R, Sarath G, Lucas RV (2001) Nonsymbiotic hemoglobins in rice are synthesized during germination and in differentiating cell types. Protoplasma 218(3-4):125–133

Safdar I, Khan S, Islam I, Ali MK, Bibi Z, Waqas M (2018) *Pichia pastoris* expression system: a potential candidate to express protein in industrial and biopharmaceutical domains. Biomedical Letters 4(1):1-14

Selb R, Wal JM, Moreno FJ, Lovik M, Mills C, Hoffmann-Sommergruber K, Fernandez A (2017) Assessment of endogenous allergenicity of genetically modified plants exemplified by soybean – Where do we stand? Food Chem Toxicol 101:139-148

Smagghe BJ, Hoy JA, Percifield R, Kundu S, Hargrove MA, Sarath G, Hilbert JL, Watts RA, Dennis ES, Peacock WJ, Dewilde S, Moens L, Blouin GC, Olsen JS, Appleby CA (2009) Correlations between oxygen affinity and sequence classifications of plant hemoglobins. Biopolymers 91:1083-1096

Sørensen HP (2010) Towards universal systems for recombinant gene expression. Microbial Cell Factories 9:e27

Spohner SC, Muller H, Quitmann H, Czermak P (2015) Expression of enzymes for the usage in food and feed industry with *Pichia pastoris*. J Biotechnol 202:118-134

Sturmberger L, Chappell T, Geier M, Krainer F, Day KJ, Vide U, Trstenjak S, Schiefer A, Richardson T, Soriaga L, Darnhofer B, Birner-Gruenberger R, Glick BS, Tolstorukov I, Cregg J, Madden K, Glieder A (2016) Refined *Pichia pastoris* reference genome sequence. J Biotechnol 235:121-131

Texas A&M Institute (2019) Meat Science: Meat Color. College Station (TX): Texas A&M Institute. Available at: <https://meat.tamu.edu/ansc-307-honors/meat-color/> Accessed: 20 October, 2019

Thomas K, Aalbers M, Bannon GA, Bartels M, Dearman RJ, Esdaile DJ, Fu TJ, Glatt CM, Hadfield N, Hatzos C, Hefle SL, Heylings JR, Goodman RE, Henry B, Herouet C, Holsapple M, Ladics GS, Landry TD, MacIntosh SC, Rice EA, Privalle LS, Steiner HY, Teshima R, van Ree R, Woolhiser M, Zawodny J (2004) A multi-laboratory evaluation of a common *in vitro* pepsin digestion assay protocol used in assessing the safety of novel proteins. Regul Toxicol Pharmacol 39:87-98

Trout GR (1989) Variation in myoglobin denaturation and color of cooked beef, pork, and turkey meat as influenced by pH, sodium chloride, sodium tripolyphosphate, and cooking temperature. J Food Sci 54(3):536-540

University of Otago, MoH (2011) A focus on nutrition: Key findings of the 2008/09 New Zealand adult nutrition survey. Chapter 4: Nutrient intakes and dietary sources: Micronutrients. Ministry of Health, Wellington, NZ

US FDA (2007a) Toxicological Principles for the Safety Assessment of Food Ingredients (Redbook 2000), IV.C.4.a.

US FDA (2007b) Toxicological Principles for the Safety Assessment of Food Ingredients (Redbook 2000), IV.C.1.a.

US FDA (2018) FDA has no questions: Soy leghemoglobin preparation from a strain of *Pichia pastoris*. GRN No. 737 <https://www.accessdata.fda.gov/scripts/fdcc/?set=GRASNotices&id=737&sort=GRN_No&order=DESC&startrow=1&type=basic&search=737>

Vainshtein BK, Harutyunyan EH, Kuranova IP, Borisov VV, Sosfenov NI, Pavlovsky AG, Grebenko AI, Konareva NV (1975) Structure of leghaemoglobin from lupin root nodules at 5 angstrom resolution. Nature 254:163-164

Vaghefi N, Nedjaoum F, Guillochon D, Bureau F, Arhan P, Bouglé D (2002) Influence of the extent of hemoglobin hydrolysis on the digestive absorption of heme iron. An in vitro study. J Agric Food Chem 50(17):4969-4973

Vogl T, Hartner FS, Glieder A (2013) New opportunities by synthetic biology for biopharmaceutical production in *Pichia pastoris*. Curr Opin Biotechnol 24(6):1094-1101

Weinborn V, Pizarro F, Olivares M, Brito A, Arredondo M, Flores S, Valenzuela C (2015) The effect of plant proteins derived from cereals and legumes on heme iron absorption. Nutrients 7(11):8977-86

WHO (2009) Safety evaluation of certain food additives. Prepared by the sixty-ninth meeting of the Joint FAO/WHO Expert Committee on Food Additives (JECFA). WHO Food Additives Series, 60. World Health Organization, Geneva.

Yamada Y, Matsuda M, Maeda K, Mikata K(1995) The phylogenetic relationships of methanol-assimilating yeasts based on the partial sequences of 18S and 26S ribosomal RNAs: the proposal of *Komagataella* gen. nov. (*Saccharomycetaceae*). Biosci Biotechnol Biochem 59(3):439-444

Yun S, Habicht JP, Miller DD, Glahn RP (2004) An in vitro digestion/Caco-2 cell culture system accurately predicts the effects of ascorbic acid and polyphenolic compounds on iron bioavailability in humans. J Nutr. 134:2717-2721

# Appendix 1: FSANZ Approach to the dietary intake assessment

A dietary intake or exposure assessment is the process of estimating how much of a nutrient or food chemical a population, or population subgroup consumes. Dietary intake of nutrients is estimated by combining food consumption data with food composition data. The process of doing this is called ‘dietary modelling’.

***Dietary intake = food chemical concentration x food consumption***

FSANZ’s approach to dietary modelling is based on internationally accepted procedures for estimating intake of nutrients (FSANZ, 2009). Different dietary modelling approaches may be used depending on the assessment, the type of food chemical, the data available and the risk assessment questions to be answered. In the majority of assessments FSANZ uses the food consumption data from each person in the national nutrition surveys to estimate their individual dietary intake. Population summary statistics such as the mean intake or a high percentile intake are derived from the ranked individual person’s intakes from the nutrition survey. In some cases, as used in this assessment for estimating total iron intakes, FSANZ will use the usual mean and high percentile usual intakes of nutrients that are published based on nutrition survey data.

An overview of how dietary intake assessments are conducted and their place in the FSANZ Risk Analysis Process is provided on the FSANZ website[[13]](#footnote-14).

FSANZ has a custom-built computer program called ‘Harvest’ to calculate nutrient intakes and dietary exposures to food chemicals. Harvest was used for this assessment to:

* estimate the dietary intake of the Preparation
* compare the baseline iron intake of meat eaters to non-meat eaters
* extract the combined consumption amounts for minced meat and poultry products and vegetarian meat alternatives for Australia and New Zealand to estimate population iron intakes with the additional contribution from soy leghemoglobin.

Further detailed information on conducting dietary intake assessments at FSANZ is provided in *Principles and Practices of Dietary Exposure Assessment for Food Regulatory Purposes* (FSANZ 2009)[[14]](#footnote-15).

## A1.1 Food consumption data used

The most recent food consumption data from national nutrition surveys were used to estimate the combined consumption amounts for minced meat and poultry products or vegetarian meat alternatives for Australia and New Zealand. Dietary intake assessments based on food consumption data from national nutrition surveys provide the best estimation of actual consumption of a food and the resulting estimated dietary intake for the Australian and New Zealand populations. However, it should be noted that national nutrition survey data do have limitations. The national nutrition survey data used for these assessments, the design of these surveys and the key attributes, including survey limitations, are set out below.

Further information on the national nutrition surveys used to conduct dietary intake assessments is available on the FSANZ website at: <http://www.foodstandards.gov.au/science/exposure/Pages/dietaryexposureandin4438.aspx>.

### 2011–12 Australian National Nutrition and Physical Activity Survey (2011–12 NNPAS)

The 2011–12 Australian National Nutrition and Physical Activity Survey (NNPAS) undertaken by the Australian Bureau of Statistics is the most recent food consumption data for Australia. This survey includes dietary patterns of a sample of 12,153 Australians aged 2 years and above. The survey used a 24-hour recall method for all respondents, with 64% of respondents also completing a second 24-hour recall on a second, non-consecutive day. The data were collected from May 2011 to June 2012 (with no enumeration between August and September 2011 due to the Census). Only those respondents who had two days of food consumption data (n=7,735) were used to derive the consumer consumption amounts for meat, poultry and vegetarian meat alternatives. The survey data was weighted for use in all calculations. Consumption and respondent data from the survey were incorporated into the Harvest program from the Confidentialised Unit Record Files (CURF) data set (ABS, 2014).

The baseline iron intakes from food and beverages, and dietary supplements, used for the Australian population in this assessment are those derived from the 2011-12 survey and published by the ABS (ABS, 2015a; ABS, 2015b). The NCI method that is used to estimate usual nutrient intakes was used to estimate the usual mean and high percentile iron intakes from food and beverages. The baseline data on intakes from dietary supplements is based on consumers who had taken iron from supplements the day prior to interview, and excludes pregnant and breastfeeding females. These intakes were used deterministically to estimate iron intakes from the diet in addition to the iron from the addition of soy leghemoglobin to meat analogue products.

### 2002 New Zealand National Children’s Nutrition Survey (2002 NZ NCNS)

The 2002 NZ NCNS was a cross-sectional and nationally representative survey of 3,275 New Zealand children aged 5-14 years. The data were collected during the school year from February to December 2002. The survey used a 24-hour food recall and provided information on food and nutrient intakes, eating patterns, frequently eaten foods, physical activity patterns, dental health, anthropometric measures and nutrition-related clinical measures. It was also the first children’s nutrition survey in New Zealand to include a second day diet recall data for about 15% of the respondents, and dietary intake from both foods (including beverages) and dietary supplements. For the consumption amounts for minced meat and poultry products and vegetarian meat alternatives in this assessment, only the first (day 1) 24-hour recall food consumption data for all consumers was used as not enough respondents completed a second 24 hour recall to enable a reliable or representative two day average to be calculated. The survey data were weighted for use in all calculations. Consumption and respondent data from the survey were incorporated into the Harvest program from the Confidentialised Unit Record Files (CURF) data set (MoH, 2005).

Baseline iron intakes from food and beverages used for New Zealand Children aged 5 to 14 years are those that were published with the 2002 NZ CNS (MoH, 2003). They are adjusted intakes derived from the distribution of usual intakes of iron using the PC-SIDE computer software. A search of the published literature found no iron intake data from dietary supplements for the New Zealand population that is equivalent to that published for the Australian 2011-12 NNPAS. It was assumed in this assessment that the average intake of iron from dietary supplements for the New Zealand population (for those who consume supplements) is the same as the Australian population.

### 2008-9 New Zealand Adult Nutrition Survey (2008-9 NZ ANS)

The 2008-9 NZ ANS provides comprehensive information on the dietary patterns of a sample of 4,721 respondents aged 15 years and above. The survey was conducted on a stratified sample over a 12 month period from October 2008 – October 2009. The survey used a 24-hour recall methodology with 25% of respondents also completing a second 24-hour recall. The information collected in the 2008 NZANS included food and nutrient intakes, dietary supplement use, socio-demographics, nutrition related health, and anthropometric measures. For the consumption amounts for minced meat and poultry products and vegetarian meat alternatives in this assessment, only the first day (day 1) 24-hour recall food consumption data for all consumers was used as not enough respondents completed a second 24 hour recall to enable a reliable or representative two day average to be calculated. The survey data were weighted for use in all calculations. Consumption and respondent data from the survey were incorporated into the Harvest program from the Confidentialised Unit Record Files (CURF) data set (MoH, 2012).

Baseline iron intakes from food and beverages used for New Zealanders 15 years and above are those published with the 2008-9 NZ ANS (University of Otago and MoH, 2011). They are adjusted intakes derived from the distribution of usual intakes of iron using the PC-SIDE computer software.

### Food groups and consumption amounts used in the dietary intake assessment

FSANZ dietary intake assessment assumes that consumers may choose to eat meat analogue products containing soy leghemoglobin in the same amounts they currently consume of traditional minced meat and poultry products, and vegetarian meat alternatives. The food groups used to derive the consumption amounts are listed in [Table A1.1](#TableA1_1). The consumption amounts for Australia and New Zealand for consumers of these foods is shown in [Table A1.2](#TableA1_2) and shown for each age/sex group in [Table A1.3](#TableA1_3). The consumption amounts include foods reported as eaten and where used in processed foods or mixed dishes, based on FSANZ recipe database in Harvest.

*Table A1.1:* Food groups used to derive consumption amounts for meat analogue products containing soy leghemoglobin

|  |  |  |
| --- | --- | --- |
| *Food groups* | *Foods included in this group* | *Foods not included in this group* |
| Minced meat products | All minced meat including sausages and luncheon meat. | Meat in whole or cut pieces. |
| Minced poultry products | All minced poultry including sausages and luncheon meat. | Poultry in whole or cut pieces. |
| Vegetarian meat alternatives | Vegetable patties, tofu, tempeh, vegetarian sausages and other meat analogues, falafel and nut meat. | Vegetable fritters (e.g. corn fritters). |

*Table A1.2:* Mean and 90th percentile consumption amounts for minced meat and poultry products and vegetarian meat alternatives for the Australian and New Zealand populations

|  |  |  |  |
| --- | --- | --- | --- |
| ***Country*** | ***Age group (years)*** | ***Mean consumption (g/day)*** | ***Consumption at 90th percentile (g/day)*** |
| **Australia\*** | 2 years and above  | 63 | 139 |
| **New Zealand**^ | 5 – 14 years  | 111 | 222 |
| 15 years and above  | 116 | 257 |

\* 2011-12 Australian National Nutrition and Physical Activity Survey. Based on consumption data from consumers with two days of data only.

**^** 2002 New Zealand National Children’s Nutrition Survey and the 2008–09 New Zealand Adult Nutrition Survey. Based on day 1 consumption data only from consumers only.

*Table A1.3:* Mean and 90th percentile consumption amounts of minced meat and poultry products and vegetarian meat alternatives by age group and sex

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| ***Country*** | ***Sex*** | ***Age group (years)*** | ***Mean consumption (g/day)*** | ***Consumption at 90th percentile (g/day)*** |
| **Australia**\* | Male | 2-3 | 34 | 71 |
| 4-8 | 50 | 97 |
| 9-13 | 63 | 144 |
| 14-18 | 93 | 190 |
| 19-30 | 80 | 182 |
| 31-50 | 75 | 164 |
| 51-70 | 72 | 171 |
| >= 71 | 71 | 156 |
| Female | 2-3 | 34 | 92 |
| 4-8 | 37 | 82 |
| 9-13 | 49 | 102 |
| 14-18 | 44 | 98 |
| 19-30 | 59 | 124 |
| 31-50 | 54 | 123 |
| 51-70 | 58 | 140 |
| >= 71 | 53 | 117 |
| **New Zealand**^ | Male | 5-6 | 95 | 174 |
| 7-10 | 122 | 224 |
| 11-14 | 144 | 307 |
| 15-18 | 139 | 274 |
| 19-30 | 155 | 396 |
| 31-50 | 138 | 290 |
| 51-70 | 106 | 269 |
| >= 71 | 94 | 188 |
| Female | 5-6 | 84 | 145 |
| 7-10 | 98 | 196 |
| 11-14 | 97 | 200 |
| 15-18 | 116 | 227 |
| 19-30 | 107 | 255 |
| 31-50 | 99 | 234 |
| 51-70 | 91 | 179 |
| >= 71 | 84 | 197 |

\* 2011-12 Australian National Nutrition and Physical Activity Survey. Based on consumption data from consumers with two days of data only.

**^** 2002 New Zealand National Children’s Nutrition Survey and the 2008–09 New Zealand Adult Nutrition Survey. Based on day 1 consumption data from consumers only.

## A1.2 Concentration data used

### The soy leghemoglobin preparation

The applicant has proposed a maximum level of 0.8% soy leghemoglobin for use in meat analogue products. The applicant has stated that the likely use levels are 0.45% soy leghemoglobin in beef analogue products and 0.25% soy leghemoglobin in pork analogue products. The dietary intake assessment assumes that these levels apply to the raw product, and that the Preparation would be further concentrated where moisture loss occurs during cooking. There are no other sources or existing permissions for soy leghemoglobin in foods currently on the market in Australia and New Zealand.

Population statistics (mean and 90th percentile) for soy leghemoglobin intakes were derived from each individual’s intake of soy leghemoglobin which were then converted to the corresponding amount of the Preparation, based on the commercial Preparation containing up to 9% soy leghemoglobin, and having a TOS content of 22%.

### Iron

Based on the atomic mass of iron (55.845) and the soy leghemoglobin protein containing the iron (16,131 Da), 0.35% of the mass of the protein is iron. For every gram of soy leghemoglobin added to a meat analogue product it is expected that 3.5 mg of iron has been contributed. If soy leghemoglobin is added to raw meat analogue products at the maximum proposed use level 0.8% (8 g/kg) the expected concentration of iron in these products (raw form) is 28 mg/kg. If soy leghemoglobin is added to raw meat analogue products at the use level of 0.45% (4.5 g/kg) the expected concentration of iron in these products (raw form) is 15.75 mg/kg. If soy leghemoglobin is added to raw meat analogue products at the use level of 0.25% (2.5 g/kg) the expected concentration of iron in these products (raw form) is 8.75mg/kg.

## A1.3 Assumptions and limitations of the dietary intake assessment

Where significant uncertainties in the data exist, conservative assumptions were made in the intake assessment to ensure the estimated intakes of the Preparation and iron were not underestimated. Assumptions made in the dietary intake assessment include:

* Moisture loss from cooking meat analogue products containing soy leghemoglobin is equivalent to the loss from cooking the meat, poultry or vegetarian meat alternatives they are replacing in the assessment.
* In estimating the intakes for the *likely use level* scenario, minced pork and poultry products can be used as a proxy for meat analogue products that may contain soy leghemoglobin at 0.25%. All other minced meat products (including minced beef and lamb products) and vegetarian meat alternatives consumed in the national nutrition survey can be used as a proxy for meat analogue products that may contain soy leghemoglobin at 0.45%.
* Iron intakes that were published for each national nutrition survey reflect the current intakes.
* The iron intake of respondents who did not consume meat or any meat products on day 1 or day 2 of the survey are representative of the iron intake of a ‘non-meat eater’.
* The average intake of iron from dietary supplements for the New Zealand population (for those who consume supplements) is the same as the Australian population.
* The iron contributed by ingredients other than the soy leghemoglobin in meat analogue products is within the range of iron in meat, poultry and vegetarian meat alternative products currently on the market in Australia and New Zealand.
* Intakes of iron from added iron for foods with existing permissions in Schedule 17 of the Food Standards Code (for example, breakfast cereals) are captured by the baseline intake data published for each national nutrition survey.

In addition to the specific assumptions made in relation to this dietary intake assessment, there are a number of limitations associated with the nutrition surveys on which this assessment is based. A discussion of these limitations is included in Section 6 of the *Principles and Practices of Dietary Exposure Assessment for Food Regulatory Purposes* (FSANZ, 2009).

## References

ABS (2014) National Nutrition and Physical Activity Survey, 2011–12, Basic Confidentialised Unit Record Files. Australian Bureau of Statistics, Commonwealth of Australia, Canberra

FSANZ (2009) Principles and practices of dietary exposure assessment for food regulatory purposes. Report prepared by Food Standards Australia New Zealand, Canberra

<http://www.foodstandards.gov.au/publications/Pages/Principles-and-Practices-of-Dietary.aspx>

MoH (2005) 2002 National Children's Nutrition Survey: National Confidentialised Unit Record File (CURF). Ministry of Health, Wellington

MoH (2012) 2008-9 New Zealand Adult Nutrition Survey: National Confidentialised Unit Record File (CURF). Ministry of Health, Wellington

# Appendix 2: Estimated dietary intake of iron with additional contribution from soy leghemoglobin

For the dietary intake assessment of iron, FSANZ first compared the baseline intakes of iron from food for meat-eaters with that of non-meat eaters for the Australian and New Zealand population using Harvest. This was undertaken in order to determine whether non-meat eaters should be considered as a separate population group in estimating dietary intakes of iron with the additional contribution from soy leghemoglobin from meat analogue products. The results of this comparison are included in Table A2.1.

Respondents who consumed meat (including poultry, seafood and game) or meat containing mixed dishes or products (e.g. fish sauce) on either day of the nutrition surveys were defined for the purpose of the assessment as ‘meat eaters’. Those who did not consume meats or meat products were defined as ‘non-meat eaters’. Only the first day of data was used for comparison of the iron intake of meat eaters to non-meat eaters as this allowed for a larger sample size of non-meat eaters to calculate the mean intake. All data were weighted in the calculations.

*Table A2.1:* Current mean iron intake of meat eaters and non-meat eaters

|  |  |  |
| --- | --- | --- |
| ***Country*** | ***Age group (years)*** | ***Mean iron intake (mg/day)*** |
|  |  | ***Meat eaters*** | ***Non-meat eaters*** |
| **Australia\*** | 2 years and above  | 11 | 8 |
| **New Zealand**^ | 5 – 14 years  | 11 | 8 |
| 15 years and above  | 12 | 9 |

\*2011-12 Australian National Nutrition and Physical Activity Survey. Based on day 1 consumption data for all respondents.

**^** 2002 New Zealand National Children’s Nutrition Survey and the 2008–09 New Zealand Adult Nutrition Survey. Based on day 1 consumption data for all respondents.

For all surveys the mean intake of iron for non-meat eaters was below that of meat eaters. As this assessment considered only exceedances of the UL, it was determined that it was not necessary to consider the estimated iron intake of non-meat eaters as a special population group.

The estimated mean and high intakes of iron, including the contribution from soy leghemoglobin, for the Australian and New Zealand population are listed in Table A2.2 and Table A2.3.

*Table A2.2:* Estimated dietary intake of iron (mg/day) for the Australian population with additional contribution from soy leghemoglobin

|  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- |
| *Age group (years)* | *UL\**  | *Mean intake from food €* | *High intake from food (P95) €*  | *Mean intake from supplements ∑*  | *Intake from soy leghemoglobin (mean)£*  | *Intake from soy leghemoglobin (P90) £*  | *Mean total intake¥*  | *High total intakeΩ*  |
|  | ***Maximum proposed use level*** | ***Likely use level*** | ***Maximum proposed use level*** | ***Likely use level*** | ***Maximum proposed use level*** | ***Likely use level*** | ***Maximum proposed use level*** | ***Likely use level*** |
| *Males* |  |  |  |  |
| 2-3 | **20** | 8 | 12 | 3 | 1 | 1 | 2 | 1 | 12 | 12 | 17 | 16 |
| 4-8 | **40** | 9 | 13 | 3 | 1 | 1 | 3 | 2 | 13 | 13 | 19 | 18 |
| 9-13 | **40** | 12 | 17 | 4 | 2 | 1 | 4 | 2 | 18 | 17 | 25 | 23 |
| 14-18 | **45** | 13 | 19 | 4 | 3 | 1 | 5 | 3 | 20 | 18 | 28 | 26 |
| 19-30 | **45** | 13 | 19 | 8 | 2 | 1 | 5 | 3 | 23 | 22 | 32 | 30 |
| 31-50 | **45** | 13 | 19 | 7 | 2 | 1 | 5 | 3 | 22 | 21 | 31 | 29 |
| 51-70 | **45** | 12 | 18 | 4 | 2 | 1 | 5 | 3 | 18 | 17 | 27 | 25 |
| 71 + | **45** | 12 | 18 | 6 | 2 | 1 | 4 | 2 | 20 | 19 | 28 | 26 |
| *Females* |  |  |  |  |
| 2-3 | **20** | 7 | 10 | 3 | 1 | 1 | 3 | 1 | 11 | 11 | 16 | 14 |
| 4-8 | **40** | 8 | 12 | 3 | 1 | 1 | 2 | 1 | 12 | 12 | 17 | 16 |
| 9-13 | **40** | 9 | 14 | 5 | 1 | 1 | 3 | 2 | 15 | 15 | 22 | 21 |
| 14-18 | **45** | 9 | 14 | 10 | 1 | 1 | 3 | 1 | 20 | 20 | 27 | 25 |
| 19-30 | **45** | 9 | 14 | 12 | 2 | 1 | 3 | 2 | 23 | 22 | 29 | 28 |
| 31-50 | **45** | 9 | 14 | 8 | 2 | 1 | 3 | 2 | 19 | 18 | 25 | 24 |
| 51-70 | **45** | 10 | 15 | 5 | 2 | 1 | 4 | 2 | 17 | 16 | 24 | 22 |
| 71 + | **45** | 9 | 14 | 6 | 1 | 1 | 3 | 2 | 16 | 16 | 23 | 22 |

\*Upper Level for iron based on the Nutrient Reference Values for Australia and New Zealand (NHMRC and MoH 2006).

€Intake derived from 2011-12 NNPAS (ABS 2015a)**.**Published usual intakes for Australia report the 95th percentile intake; this has been used for the estimated high intakes from food for Australia.

**∑** Intake derived from 2011-12 NNPAS (ABS 2015b).

**£** Population statistics (mean and 90th percentile) were derived from each individual’s intake of soy leghemoglobin then converted to the corresponding amount of iron.

¥Sum of mean intake from food and beverages, dietary supplements and soy leghemoglobin.

ΩSum of high intake from food and beverages, mean intake from dietary supplements and high intake from soy leghemoglobin.

*Table A2.3:* Estimated intake of iron (mg/day) for the New Zealand population with additional contribution from soy leghemoglobin

|  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- |
| *Age group (years)* | *UL\** | *Mean intake from food €* | *High intake from food (P90) €* | *Mean intake from supplements ∑* | *Intake from soy leghemoglobin (mean) £*  | *Intake from soy leghemoglobin (P90) £* | *Mean total intake ¥*  | *High total intake Ω* |
|  |  |  |  |  | ***Maximum proposed use level*** | ***Likely use level*** | ***Maximum proposed use level*** | ***Likely use level*** | ***Maximum proposed use level*** | ***Likely use level*** | ***Maximum proposed use level*** | ***Likely use level*** |
| *Males* |
| 5-6 | **40** | 10 | 13 | 3 | 3 | 1 | 5 | 3 | 16 | 15 | 21 | 19 |
| 7-10 | **40** | 12 | 16 | 4 | 3 | 2 | 6 | 3 | 20 | 18 | 26 | 24 |
| 11-14 | **40-45\*\*** | 14 | 19 | 4 | 4 | 2 | 9 | 5 | 22 | 21 | 31 | 27 |
| 15-18 | **45** | 14 | 19 | 4 | 4 | 2 | 8 | 4 | 22 | 20 | 31 | 27 |
| 19-30 | **45** | 14 | 20 | 8 | 4 | 2 | 11 | 6 | 27 | 25 | 39 | 34 |
| 31-50 | **45** | 14 | 20 | 7 | 4 | 2 | 8 | 4 | 25 | 23 | 35 | 31 |
| 51-70 | **45** | 13 | 19 | 4 | 3 | 2 | 8 | 4 | 20 | 19 | 31 | 27 |
| 71 + | **45** | 12 | 16 | 6 | 3 | 1 | 5 | 3 | 20 | 19 | 27 | 25 |
| *Females* |  |  |  |  |
| 5-6 | **40** | 9 | 11 | 3 | 2 | 1 | 4 | 2 | 14 | 13 | 18 | 16 |
| 7-10 | **40** | 10 | 13 | 5 | 3 | 1 | 5 | 3 | 18 | 16 | 23 | 21 |
| 11-14 | **40-45\*\*** | 10 | 15 | 10 | 3 | 1 | 6 | 3 | 23 | 22 | 30 | 28 |
| 15-18 | **45** | 9 | 13 | 10 | 3 | 2 | 6 | 3 | 23 | 21 | 29 | 26 |
| 19-30 | **45** | 10 | 12 | 12 | 3 | 2 | 7 | 4 | 25 | 24 | 31 | 28 |
| 31-50 | **45** | 10 | 14 | 8 | 3 | 1 | 7 | 3 | 21 | 20 | 28 | 25 |
| 51-70 | **45** | 10 | 14 | 5 | 3 | 1 | 5 | 3 | 18 | 17 | 24 | 21 |
| 71 + | **45** | 9 | 13 | 6 | 2 | 1 | 6 | 3 | 18 | 17 | 24 | 22 |

\* Upper Level for iron based on the Nutrient Reference Values for Australia and New Zealand (NHMRC and MoH 2006).

\*\*The UL for 9-13 years olds is 40 mg/day. The UL for 14 year olds is 45 mg/day.

€Intake derived from the 2002 NZ CNS and 2008-9 NZ ANS (Ministry of Health 2003; University of Otago and MoH 2011). The highest reported intake in these publications is the 90th percentile; this has been used for the estimated high intakes from food for New Zealand.

∑ Australian mean intake for supplements used (ABS 2015b). For 5-6 year olds the average used is for Australians aged 4-8 years. For 7-10 years olds the mean used is for Australians aged 9-13 years. For 11-14 year olds the average used is for Australians aged 14-18 years.

**£** Population statistics (mean and 90th percentile) were derived from each individual’s intake of soy leghemoglobin then converted to the corresponding amount of iron.

¥Sum of mean intake from food and beverages, dietary supplements and soy leghemoglobin. As the intakes in the table have been rounded to the nearest whole number the mean total intake for each age group may not always exactly equal the sum of the intakes displayed.

ΩSum of high intake from food and beverages, mean intake from dietary supplements and high intake from soy leghemoglobin. As the intakes in the table have been rounded to the nearest whole number the high total intake for each age group may not always exactly equal the sum of the intakes displayed.

## References

ABS (2015a) Australian health survey: Usual Nutrient Intakes, 2011-12. Table 4:Essential minerals (and caffeine). Australian Bureau of Statistics, Commonwealth of Australia, Canberra

ABS (2015b) Australian health survey: Nutrition – Supplements, 2011-12. Australian Bureau of Statistics, Commonwealth of Australia, Canberra

MoH (2003) NZ food NZ children: Key results of the 2002 national children's nutrition survey. Ministry of Health, Wellington

NHMRC, MoH (2006) Nutrient reference values for Australia and New Zealand. National Health and Medical Research Council and New Zealand Ministry of Health, Canberra, Australia

University of Otago, MoH (2011) A focus on nutrition: Key findings of the 2008/09 New Zealand adult nutrition survey. Chapter 4: Nutrient intakes and dietary sources: Micronutrients. University of Otago and Ministry of Health, Wellington

1. FSANZ will herein use the term ‘soy leghemoglobin preparation’ and ‘the Preparation’ not LegH Prep. The first is the name used in the draft variation to the Code and the latter is the applicant’s brand name. [↑](#footnote-ref-2)
2. A holoprotein is a protein bound to a prosthetic group (e.g. haem) [↑](#footnote-ref-3)
3. FSANZ will herein use the term ‘soy leghemoglobin preparation’ and ‘the Preparation’ not LegH Prep. The first is the name used in the draft variation to the Code and the latter is the applicant’s brand name. [↑](#footnote-ref-4)
4. The applicant refers to ‘ground’ meat products, however in Australia and New Zealand the term ‘minced’ meat is used and from herein will be referred to as such. [↑](#footnote-ref-5)
5. [www.uniprot.org/uniprot/P02236](http://www.uniprot.org/uniprot/P02236) [↑](#footnote-ref-6)
6. [web.expasy.org/compute\_pi/](https://web.expasy.org/compute_pi/) [↑](#footnote-ref-7)
7. [www.foodstandards.gov.au/consumer/foodallergies/Pages/Allergen-labelling.aspx](http://www.foodstandards.gov.au/consumer/foodallergies/Pages/Allergen-labelling.aspx) [↑](#footnote-ref-8)
8. [www.allergenonline.org/databasebrowse.shtml](http://www.allergenonline.org/databasebrowse.shtml) [↑](#footnote-ref-9)
9. [www.allergenonline.org](http://www.allergenonline.org) [↑](#footnote-ref-10)
10. The study sponsor provided the test laboratory with information indicating that the test article contained 6.74% soy leghemoglobin and this value was used to calculate study concentrations. Product Safety Laboratories prepared a GLP certificate of analysis and found soy leghemoglobin at 6.68%. This value was considered to be within an acceptable margin of error. [↑](#footnote-ref-11)
11. Nutrient reference values, Australia and New Zealand: <https://www.nrv.gov.au/> [↑](#footnote-ref-12)
12. [www.uniprot.org/uniprot/P02236](http://www.uniprot.org/uniprot/P02236) [↑](#footnote-ref-13)
13. [www.foodstandards.gov.au/science/riskanalysis/Pages/default.aspx](http://www.foodstandards.gov.au/science/riskanalysis/Pages/default.aspx) [↑](#footnote-ref-14)
14. [www.foodstandards.gov.au/science/exposure/documents/Principles%20\_%20practices%20exposure%20assessment%202009.pdf](http://www.foodstandards.gov.au/science/exposure/documents/Principles%20_%20practices%20exposure%20assessment%202009.pdf). [↑](#footnote-ref-15)