## High Sensitive Peanut ELISA Kit II

The Quantitative Determination for<br>Peanut Protein in Food

96 Assays

For Research or Laboratory Use Only. Not for Use in Diagnostic Procedures. Please read full descriptions in this manual before use.

Manufactured by:<br>Morinaga Institute of Biological Science, Inc. (MloBS)<br>2-1-16 Sachiura, Kanazawa-ku, Yokohama 236-0003, Japan<br>E-mail: info_miobs_e@morinaga.co.jp<br>Website: https://www.miobs-e.com

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For optimal results, follow instructions exactly. Failure to do so may lead to inaccurate results.

Reproducible results depend on careful pipetting technique, maintaining incubations at the specified temperature for the specified time, complete washing, and thorough mixing of all solutions.

## INTENDED USE AND USER

High Sensitive Peanut ELISA Kit II is a sandwich enzyme immunoassay for the quantitative determination of peanut protein in processed or unprocessed food. This kit is designed to be used by quality control personnel or other trained professionals.

Table 1 : Detectable protein in this kit

| Kit | Antigen | Detectable Protein |
| :---: | :---: | :---: |
| High Sensitive <br> Peanut ELISA Kit II | Ara h2, Ara h6 | Whole peanut protein |

## PRINCIPLE OF ELISA

Peanut protein in the samples is extracted, centrifuged and filtered prior to the sandwich ELISA. Antigen is bound to the polyclonal antibody coated wells of the microplate module. This results in the formation of an antigen-antibody complex in the wells. Unbound materials are removed by washing. Subsequently, the enzyme-conjugated antibody is bound to the already bound antigen-antibody complex, forming an
antibody-antigen-antibody sandwich. A second washing step removes the excess conjugated antibody. Addition of enzyme substrate results in color development due to the enzyme bound to the complex. After addition of the stop solution, the color intensity of the solutions can be determined by the absorbance at 450 nm . The intensity of the color developed is directly proportional to the concentration of the peanut protein in the food. The concentration of peanut protein corresponding to the measured absorbance is determined by preparing a standard curve, and adjusting for a further dilution factor if necessary.

KIT COMPONENTS

| Label | Name of component | Content | Quantity |
| :---: | :--- | :---: | :---: |
| A | Reagent A (10X Concentrate) | 100 mL | 1 bottle |
| B | Reagent B (10X Concentrate) | 100 mL | 1 bottle |
| C | Reagent C (10X Concentrate) | 100 mL | 1 bottle |
| D | Antibody-coated Microplate Module | $6 \times 8 \mathrm{well}$ <br> modules | 2 packs |
| E | Peanut Standard (32 ng/mL) | 1 mL | 2 vials |
| F | Enzyme-conjugated Antibody | 13 mL | 1 bottle |
| G | Enzyme Substrate (TMB Solution) | 13 mL | 1 bottle |
| H | Stop Solution (1N Sulfuric Acid) | 13 mL | 1 bottle |
| I | Wash Buffer (20X Concentrate) | 50 mL | 1 bottle |
|  | Frame for mounting the microplate <br> module |  | 1 piece |
|  | Microplate cover |  | 1 piece |

Note: Peanut Standard ( $32 \mathrm{ng} / \mathrm{mL}$ ) (E) is made from the NIST Peanut Butter SRM2387.

## REQUIRED MATERIALS (NOT INCLUDED)

1. Distilled water (or deionized water)
2. Micropipettes and disposable tips ranging from 50 to $1000 \mu \mathrm{~L}$
3. Graduated cylinders
4. Polypropylene centrifuge tubes ( 50 mL size) for sample extraction
5. Polypropylene micro tubes ( $1-2 \mathrm{~mL}$ size) for preparation of working standard solutions and dilution of sample
6. pH test paper
7. Homogenizer/blender for sample preparation (if necessary)
8. Water bath for boiling or horizontal shaker, for sample extraction
9. Centrifuge
10. Vortex mixer
11. Aspirator for washing procedure, or optional microplate washer
12. Microplate reader with a 450 nm filter, and a filter for any wavelength from 600 to 650 nm , inclusive

## PERFORMANCE CHARACTERISTICS

High Sensitive Peanut ELISA Kit II is a tool to determine the presence of peanut protein in foods.

| Sample preparation time | Approx. 10 min |
| :--- | :--- |
| Sample extraction time | Approx. 30 min, or overnight |
| Time for ELISA analysis | Approx. 2 h |
| Assay sensitivity | $0.2 \mathrm{ppm}(0.2 \mathrm{~g}$ peanut protein $/ \mathrm{g}$ food $)$ |
| Detectable concentration range | 0.5 to 32 ppb peanut protein |

## REAGENT PREPARATION

## Sample Extraction Solution

Prepare the Sample Extraction Solution by mixing Reagent $A(A)$,
Reagent $\boldsymbol{B}(\boldsymbol{B})$, Reagent $\boldsymbol{C}(\boldsymbol{C})$ and distilled water at a ratio of 1:1:1:7.
The following example would prepare enough Sample Extraction Solution to assay 16 samples:

| Reagent A (10X Concentrate) (A) | 40 mL |
| ---: | ---: |
| Reagent B (10X Concentrate) (B) | 40 mL |
| Reagent C (10X Concentrate) (C) | 40 mL |
|  | Distilled water 280 mL <br>  Total <br>  400 mL |

## Note:

a. Reagent A (A) may produce crystals after refrigerated storage. These crystals must be re-dissolved completely in a water bath at $30-37^{\circ} \mathrm{C}$ $\left(86-99^{\circ} \mathrm{F}\right)$ prior to use. The fully re-dissolved Reagent A (A) can be stored at $20-30^{\circ} \mathrm{C}\left(68-86^{\circ} \mathrm{F}\right)$.
b. Sample Extraction Solution can be stored at $4^{\circ} \mathrm{C}$ for preparing Diluent
II. If Sample Extraction Solution forms a precipitate after refrigerated storage, then the solution must be warmed in a water bath at $20-30^{\circ} \mathrm{C}$ (68-86 ${ }^{\circ}$ F) to re-dissolve the precipitate prior to use.

## Diluent I

Diluent I is used for preparing Diluent II and for diluting the Sample Extract.
Dilute Reagent C (C) 10 -fold with distilled water.
The following is an example of preparing Diluent $I$ :

| Reagent C (10X Concentrate) (C) | 5 mL |
| ---: | ---: |
| Distilled water | 45 mL |
| Total | 50 mL |

## Diluent II

Diluent II is used to prepare Working Peanut Standard, and to further dilute the Working Sample Solution.
Dilute Sample Extraction Solution, 20 -fold with Diluent I.
The following is an example of preparing Diluent II:

| Sample Extraction Solution | 1 mL |
| ---: | ---: |
| Diluent I | 19 mL |
| Total | 20 mL |

## Washing Solution

Dilute Wash Buffer (I) 20 -fold with distilled water, and gently mix until the solution is homogeneous.

Peanut Standard ( $32 \mathrm{ng} / \mathrm{mL}$ ) (E)
Peanut Standard ( $\mathbf{3 2} \mathrm{ng} / \mathrm{mL}$ ) ( $E$ ) is used as a 32 ppb standard as well as for preparing the diluted working standards described in ASSAY
PROTOCOL b.

## ASSAY PROTOCOL

It is strongly recommended to use disposable polypropylene tubes, and care should be taken to clean up all equipments and materials so as to be free from cross-contamination, since the assay is highly sensitive.
a. Sample Preparation

Extraction of peanut protein from the test food sample can be performed by two different methods depending on customer's convenience.

## (Short Time Extraction Method)

1. Grind and mix up the test food sample to homogeneity with a contamination-free homogenizer/blender.
2. A disposable polypropylene centrifuge tube, containing 1.0 g of the homogenized sample and 19 mL of Sample Extraction Solution, is capped tightly and vortexed for 30 seconds.
3. Incubate capped tube in a boiling water bath at $100^{\circ} \mathrm{C}\left(212^{\circ} \mathrm{F}\right)$ for 10 minutes.
4. Place the tube in running water to cool it down to ambient temperature (approximately 10 minutes).
5. Vortex the tube for 30 seconds.
6. Check the fluid pH with pH test paper, and neutralize ( $\mathrm{pH} 6-8$ ) with HCl or NaOH , if required.
7. Centrifuge the tube at $3,000 \times g$ for 20 minutes at $20-30^{\circ} \mathrm{C}\left(68-86^{\circ} \mathrm{F}\right)$, and the supernatant is retained as Sample Extract. (Filter the supernatant with filter paper, if necessary.)
8. Dilute the Sample Extract by 20 -fold with Diluent I (see REAGENT PREPARATION), and the diluted solution is used as the Working Sample Solution for ELISA.
Note: If further sample dilution is required, dilute the Working Sample Solution with Diluent II.

## (Overnight Extraction Method)

1. Grind and mix up the test food sample to homogeneity with a contamination-free homogenizer/blender.
2. A disposable polypropylene centrifuge tube, containing 1.0 g of the homogenized sample and 19 mL of Sample Extraction Solution, is capped tightly and vortexed for 30 seconds.
3. Fix the capped centrifuge tube to a shaker horizontally, and oscillate at room temperature overnight (for at least 12 hours at 90-110 rpm) with a reciprocating motion of about 3 cm .
4. Follow the procedures 6-8 for Short Time Extraction Method.

## b. Working Peanut Standard Preparation

1. Dispense 0.5 mL Diluent II into six polypropylene micro tubes labeled $0.5,1,2,4,8$ and 16 ppb .
2. Dispense 0.5 mL of Peanut Standard ( $\mathbf{3 2} \mathbf{n g} / \mathrm{mL}$ ) (E) into the 16 ppb tube, and mix thoroughly.
3. Dispense 0.5 mL of the freshly-prepared 16 ppb standard into the 8 ppb tube, and mix thoroughly.
4. Repeat this 2 -fold dilution series to prepare $4,2,1$ and 0.5 ppb standards.
5. Dispense 0.5 mL of Diluent II into the polypropylene micro tube labeled 0 ppb .

## c. ELISA

## (First reaction)

1. Unseal the Antibody-coated Microplate Module (D) after equilibrated to room temperature $\left(20-30^{\circ} \mathrm{C}, 68-86^{\circ} \mathrm{F}\right)$. Fix the module to the mounting frame (supplied).
2. Into each well, dispense $100 \mu \mathrm{~L}$ of standards ( $0,0.5,1,2,4,8,16$ and 32 ppb Working Peanut Standard) and Working Sample Solution, in (at least) duplicate.
Note: The test food sample presumed to contain the peanut protein greater than 32 ppb should be diluted with Diluent II (ASSAY PROTOCOL a.8, Note).
3. Cover the microplate with a microplate cover (supplied).
4. Incubate for 1 hour at $20-30^{\circ} \mathrm{C}\left(68-86^{\circ} \mathrm{F}\right)$.

## (Second reaction)

1. Remove the solution completely from each well by aspiration. (OR: Carefully pour the liquid out of the wells and tap the microplate frame upside down vigorously five times on paper towels to ensure complete removal of liquid from wells.)
2. Wash six times, by adding $300 \mu \mathrm{~L}$ Washing Solution per well followed by aspiration. After washing, remove any remaining solution by inverting and tapping the plate on a clean paper towel. (OR: Fill all the wells ( $300 \mu \mathrm{~L}$ of the Washing Solution in each), pour out the liquid again and tap. Repeat five more times.)
3. Add $100 \mu \mathrm{~L}$ Enzyme-conjugated Antibody (F) to each well.
4. Cover the microplate with a microplate cover and incubate for precisely 30 minutes at $20-30^{\circ} \mathrm{C}\left(68-86^{\circ} \mathrm{F}\right)$.

## (Enzyme reaction)

1. Completely aspirate the well contents and wash six times by filling with $300 \mu \mathrm{~L}$ Washing Solution per well, and aspirate. After the sixth washing, remove any remaining solution by inverting and tapping the plate on a clean paper towel. (OR: Completely pour the liquid out of the wells, then fill all the wells with $300 \mu \mathrm{~L}$ of the Washing Solution, pour out the liquid again and tap. Repeat five more times.)
2. Immediately, dispense $100 \mu \mathrm{~L}$ of Enzyme Substrate (G) into each well.
3. Cover the microplate with the microplate cover and incubate for precisely 20 minutes at $20-30^{\circ} \mathrm{C}\left(68-86^{\circ} \mathrm{F}\right)$. During the enzyme reaction, avoid exposing the microplate to light.
4. Stop the enzyme reaction by adding $100 \mu \mathrm{~L}$ of Stop Solution $(H)$ to each well.
5. Immediately measure absorbance using a microplate reader at 450 nm , subtract a reference wavelength between 600 and 650 nm , inclusive.
Note: The absorbance must be measured within 30 minutes after stopping the enzyme reaction.

## Summary of Procedure

(Extraction procedure)


Incubate capped tube in boiling water for 10 minutes.

Cool down the tube and vortex for 30 sec .


Adjust pH to 6.0-8.0.
Centrifuge and filter the supernatant if necessary.


Dilute Sample Extract 20 -fold with Diluent $I$.

## Overnight Extraction Method Grind/mince the sample

Weigh 1.0 g sample in a tube, add 19 mL Sample Extraction Solution.

Fix the tube to a shaker horizontally, and oscillate at room temperature overnight.

Adjust pH to 6.0-8.0.
Centrifuge and filter the supernatant if necessary.
$\downarrow$
Dilute Sample Extract 20 -fold with Diluent I.

## (ELISA procedure)

Pipette $100 \mu \mathrm{~L}$ Working Peanut Standard and Working Sample Solution.

Wash the wells 6 times with Washing Solution.

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Dispense 100 \muL Enzyme-conjugated Antibody (F).
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    \(\downarrow\)
    Incubate the microplate for 30 min at \(20-30^{\circ} \mathrm{C}\left(68-86^{\circ} \mathrm{F}\right)\).
    Wash the wells 6 times.
$\downarrow$
Dispense $100 \mu \mathrm{~L}$ Enzyme Substrate (G).
Incubate reaction for 20 min at $20-30^{\circ} \mathrm{C}\left(68-86^{\circ} \mathrm{F}\right)$ in the dark.
Stop the enzyme reaction by adding $100 \mu \mathrm{~L}$ Stop Solution (H).
Measure absorbance at 450 nm .

Calculate the concentration of peanut protein using the standard curve.

## RESULTS AND INTERPRETATION

1. Determine the mean absorbance for each set of Working Sample Solution and Working Peanut Standard. Use graphing software, the built-in graphing feature of the microplate reader, or graph paper to construct a standard curve by plotting the mean absorbance for each standard on the Y axis versus the corresponding standard concentration on the X axis.
Note: A standard curve should be generated for each assay.
For computer processing of the data, 4-parameter curve fit (Cubic regression) is recommended and a linear curve fit is also available.
2. The peanut protein concentration in the Working Sample Solution is interpolated from the standard curve using the mean absorbance of each observation.
3. If the mean absorbance of Working Sample Solution is greater than the absorbance of the 32 ppb standard, increase the dilution of Working Sample Solution to 40 -fold or greater as appropriate instead of 20 -fold, and then assay again.

The peanut protein content in a sample, in ppm, can be estimated using the following formula:

Peanut protein content $(\mathrm{ppm})=0 \mathrm{~V}_{\times}$Dilution $\mathrm{A}_{\times}$Dilution $\mathrm{B} \times 1 / 1,000$
OV: Observed value (ppb)
Dilution A: Dilution for ASSAY PROTOCOL a.2, nominal 20-fold.
Dilution B: Dilution for ASSAY PROTOCOL a.8, nominal 20-fold.

## TYPICAL STANDARD CURVE

## Peanut



## STORAGE CONDITIONS AND EXPIRY OF KIT

1. Store the kit at $2-8^{\circ} \mathrm{C}\left(35-46^{\circ} \mathrm{F}\right)$, but DO NOT FREEZE!
2. Use the kit at $20-30^{\circ} \mathrm{C}\left(68-86^{\circ} \mathrm{F}\right)$, while minimizing exposure time at this temperature. The kit must be returned to storage at $2-8^{\circ} \mathrm{C}\left(35-46^{\circ} \mathrm{F}\right)$ as soon as possible. Do not expose this kit to temperatures in excess of $30^{\circ} \mathrm{C}\left(86^{\circ} \mathrm{F}\right)$ !
3. Do not use the kit after the expiration date indicated on the outside box.

## WARNINGS AND PRECAUTIONS

(General)

1. Peanut protein is used in this kit. Users who are allergic to this protein should handle kit reagents and perform the assay with great care. In case of an allergic reaction, such as sneezing or itching, seek medical attention if the reaction is serious or prolonged.
2. This product is recommended for use only by personnel trained in analytical sample handling techniques, and is meant to be used in accordance with good laboratory practices.
3. Direct contact with Stop Solution (H) should be strictly avoided. In case of contact, immediately flush affected area with copious amounts of water, and seek medical attention if necessary. In case of ingestion, call the poison control center and obtain first aid treatment.
4. Because all chemicals should be considered potentially hazardous, it is advisable to wear suitable protective clothing, goggles and gloves.
5. Because the assay is highly sensitive, all the procedures should be performed in a clean environment using uncontaminated equipment/devices and tubes/containers in order to minimize the risk of cross-contamination from previous analyses.
(ELISA)
6. Do not combine reagents from different lots or other kits.
7. All reagents should be equilibrated at $20-30^{\circ} \mathrm{C}\left(68-86^{\circ} \mathrm{F}\right)$ before use.
8. A standard curve should be generated for each assay concurrently with the samples.
9. Assays should be performed at least in duplicate to ensure confidence in the measured values.
10. Care should be taken to pipette standard solutions and samples accurately.
11. Follow all procedures carefully.
12. Washing must be thorough in order to minimize background readings. Complete removal of reagents from the microplate wells is essential.
13. The enzyme substrate reaction should be performed shielded from light.

Note: In the case of processed foods, the detection efficiency or sensitivity of the assay may decrease, due to denaturation and insolubilization of peanut protein. Therefore, food samples that give a negative result may still contain peanut material which is either unreactive or present
at concentrations below the limit of detection. It should not be assumed that such foods are peanut free.

## TECHNICAL ASSISTANCE

For further technical assistance or troubleshooting advice, contact Morinaga Institute of Biological Science, Inc. or your local distributor.

Morinaga Institute of Biological Science, Inc. E-mail: info_miobs_e@morinaga.co.jp

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