**AccuDiag™ Fish ELISA**

**Test** | **Fish ELISA**  
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Recovery | 93 - 117%  
Incubation Time | 60 min  
Sensitivity | 1.4 ppm  

**INTENDED USE**

Fishes belong to the most frequent elicitors of food allergies. The allergies are predominantly induced by the low-molecular, calcium-binding muscle protein parvalbumin. The protein is characterized by its high heat resistance and stability against denaturing agents and proteolytic enzymes. Predominantly in regions with a high consumption of fish like Scandinavia, Japan or the Mediterranean countries, fish allergies represent a heavy health problem. The symptoms are ranging from inflammation of the skin over gastrointestinal and respiratory problems up to life-threatening anaphylactic shock. In spite of the high biodiversity most patients react with allergic symptoms to several fish species due to the high cross-reactivity between the fish allergens.

For fish-allergic persons hidden fish allergens in food are a critical problem. Already very low amounts of fish can cause allergic reactions, which may lead to anaphylactic shock in severe cases. Because of this, fish-allergic persons must strictly avoid the consumption of fish containing food. Cross-contamination, mostly in consequence of the production process, is often noticed. This explains why in many cases the existence of fish residues in food cannot be excluded. For this reason sensitive detection systems for fish residues in foodstuffs are required.

The DAI Fish ELISA represents a highly sensitive detection system for fish, based on the trans-species allergen parvalbumin. It is particularly capable of the quantification of fish residues in wine, soups, sauces, crackers, surimi and Asian products.

**TEST PRINCIPLE**

The DAI Fish ELISA is based on the principle of the enzyme linked immunosorbent assay. An antibody directed against fish proteins is bound on the surface of a microtiter plate. Fish containing samples or standards are given into the wells of the microtiter plate. After 20 minutes incubation at room temperature, the wells are washed with diluted washing solution to remove unbound material. A peroxidase conjugated second antibody directed against fish proteins is given into the wells and after 20 minutes of incubation the plate is washed again. A substrate solution is added and incubated for 20 minutes, resulting in the development of a blue color. The color development is terminated by the addition of a stop solution, and the color turns yellow. The yellow color is measured photometrically at 450 nm. The concentration of fish is directly proportional to the color intensity of the test sample.

**MATERIALS AND COMPONENTS**

**Materials provided with the test kits**

The kit contains reagents for 96 determinations. They have to be stored at 2-8°C. Expiry data are found on the labels of the bottles and the outer package.

1. Microtiter plate consisting of 12 strips with 8 breakable wells each, coated with anti-fish antibodies.
2. Cod Standards (0, 4, 10, 40, 100 ppm of cod): 5 vials with 4.0 mL each, dyed red, ready-to-use.
5. Stop Solution (0.5 M H$_2$SO$_4$): 15 mL, ready-to-use.
6. Extraction and sample dilution buffer (Tris): 2 x 120 mL as 10x concentrate, dyed red. Dilute 1+9 with distilled water. Stored at 4°C the diluted buffer is stable for at least one week. If during the cold storage crystals precipitate, the concentrate should be warmed up to 37°C for 15 minutes.
7. Washing Solution (PBS + Tween 20): 60 mL as 10x concentrate. Dilute 1+9 with distilled water. Stored at 4°C the diluted buffer is stable for at least 4 weeks. If during the cold storage crystals precipitate, the concentrate should be warmed up to 37°C for 15 minutes.
8. Plastic bag to store unused microtiter strips.

**Materials required but not provided**

**Instrumentation**
1. 100 - 1000 µL micropipettes
2. Volumetric flask
3. Analytical balance
4. Mortar, mixer
5. Water bath
6. Centrifuge
7. ELISA reader (450 nm)

**Reagents**

Double-distilled water

**SAMPLE PREPARATION**

Due to the high risk of cross-contamination all applied instruments like applicator, mortar, glass vials etc. have to be cleaned thoroughly before and after each sample. Fish proteins could adsorb to different surfaces. To identify possible cross-contamination caused by previous extractions it is strongly recommended to note the sequence of the extractions.

The following sample preparation should be applied for solid samples:

1. To maximize homogeneity and representative-ness of the sample drawing, a minimum of 5 g sample should be pulverized finely in a mortar, impact mill etc.
2. 1 g of the homogenized mixture is suspended in 20 mL of pre-diluted extraction and sample dilution buffer. Afterwards the suspension is incubated for 15 min...
in a preheated water bath at 60°C. To ensure good homogeneity, the samples should be shaken every two minutes. The samples are centrifuged for 10 minutes at 2000 g. If it is not possible to separate the supernatant from the precipitate completely, the suspension should be filtered if necessary.

4. 100 μL of particle-free solution are applied per well. If the results of a sample are out of the measuring range, further dilution with the pre-diluted extraction and sample dilution buffer is necessary. The additional dilution has to be considered when calculating the concentration.

The following sample preparation should be applied for liquid samples:

1 mL of liquid sample is diluted in 19 mL of pre-diluted extraction and sample dilution buffer. Afterwards the suspension is incubated for 15 min in a preheated water bath at 60°C. To ensure good homogeneity, the samples should be shaken every two minutes. The process is continued at point 3 of solid sample extraction process.

**PRECAUTION**

Full compliance of the following good laboratory practices (GLP) will determine the reliability of the results:

1. Prior to beginning the assay procedure, bring all reagents to room temperature (20-25°C).
2. All reagents should be mixed by gentle inversion or swirling prior to use. Do not induce foaming.
3. Once the assay has been started, all subsequent steps should be completed without interruption and within the recommended time limits.
4. Replace caps in all the reagents immediately after use. Do not interchange vial stoppers.
5. Use a separate disposable tip for each specimen to prevent cross-contamination.
6. All specimens and standards should be run at the same time, so that all conditions of testing are the same.
7. Do not mix components from different batches.
8. Do not use reagents after expiration date.
9. Allow the reaction to develop in the dark (e.g. cupboard or drawer; the chromogen is light-sensitive) for 20 minutes at room temperature.

**HEALTH & SAFETY INSTRUCTIONS**

1. Do not smoke or eat or drink or pipet by mouth in the laboratory.
2. Wear disposable gloves whenever handling patient specimens.
3. Avoid contact of substrate and stop solution with skin and mucosa (possible irritation, burn or toxicity hazard). In case of contact, rinse the affected zone with plenty of water.
4. Handling and disposal of chemical products must be done according to good laboratory practices (GLP).

**ASSAY PROCEDURE**

The washing solution is supplied as 10x concentrate and has to be diluted 1+9 with double distilled water before use.

In any case the ready-to-use standards should be determined twofold. When samples in great quantities are determined, the standards should be pipetted once before the samples and once after the samples. For final interpretation the arithmetic mean is used for calculation.

In consideration of GLP and quality control requirements a duplicate measurement of samples is recommended.

The procedure is according to the following scheme:

1. Prepare samples as described above.
2. Pipet 100 μL ready-to-use standards or prepared samples in duplicate into the appropriate wells of the microtiter plate.
3. Incubate for 20 minutes at room temperature.
4. Wash the plate three times as follows: Discard the contents of the wells (dump or aspirate). Pipet 300 μL of diluted washing solution into each well. After the third repetition empty the wells again and remove residual liquid by striking the plate against a paper towel. The wash procedure is critical. Insufficient washing will result in poor precision and falsely elevated absorbencies.
5. Pipet 100 μL of conjugate (anti-fish-peroxidase) into each well.
6. Incubate for 20 minutes at room temperature.
7. Wash the plate as outlined in 4.
8. Pipet 100 μL of substrate solution into each well.
9. Allow the reaction to develop in the dark (e.g. cupboard or drawer; the chromogen is light-sensitive) for 20 minutes at room temperature.
10. Stop enzyme reaction by adding 100 μL of stop solution (0.5 M H₂SO₄) into each well. The blue color will turn yellow upon addition.
11. After thorough mixing, measure absorbance at 450 nm (reference wavelength 620 nm), using an ELISA reader. The color is stable for 30 minutes.

**RESULTS**

The ready-to-use standards are prepared for a direct determination of sample concentrations. The dilution of samples in the extraction process as described in the above stated sample preparation procedure is already considered. Additional dilution due to high sample concentration has to be accounted for.

1. Calculate the average optical density (OD 450 nm) for each set of reference standards or samples.
2. Construct a standard curve by plotting the mean optical density obtained for each cod reference standard against its concentration in ppm on semi-log graph paper with the optical density on the vertical (y) axis and the cod concentration on the horizontal (x) axis. Alternatively the evaluation can be carried out by software. In this case the 4-parameter method should be preferred.
3. Using the mean optical density value for each sample, determine the corresponding equivalent concentration of cod in ppm from the standard curve. Depending on experience and/or the avail-ability of computer capability, other methods of data reduction may be employed.
4. If the fish species of the sample is known, the amount of the appropriate species can be calculated by multiplying the test result (cod) with a species specific conversion factor:

<table>
<thead>
<tr>
<th>Fish Species</th>
<th>Conversion factor (F)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eel</td>
<td>18</td>
</tr>
<tr>
<td>Flounder</td>
<td>7.4</td>
</tr>
<tr>
<td>Perch</td>
<td>1.1</td>
</tr>
<tr>
<td>Trout</td>
<td>5.4</td>
</tr>
<tr>
<td>Pike</td>
<td>1.2</td>
</tr>
<tr>
<td>Herring (smoked)</td>
<td>67</td>
</tr>
<tr>
<td>Carp</td>
<td>1.3</td>
</tr>
<tr>
<td>Salmon</td>
<td>39</td>
</tr>
</tbody>
</table>
It has to be considered that the standardization as well as the conversion factors relate to fresh fish. For the interpretation of the test results, the grade of process of the respective food sample has to be accounted for. Validation experiments showed that cooked cod meat (20 min) resulted in a reactivity of 25% compared to fresh cod.

### Typical Standard Values

The following table contains an example for a typical standard curve. The binding is calculated as percent of the absorption of the 100 ppm standard. These values are only an example and should not be used instead of the standard curve which has to be measured in each new test.

<table>
<thead>
<tr>
<th>Cod (ppm)</th>
<th>% binding of 100 ppm</th>
</tr>
</thead>
<tbody>
<tr>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>40</td>
<td>63</td>
</tr>
<tr>
<td>10</td>
<td>24</td>
</tr>
<tr>
<td>4</td>
<td>14</td>
</tr>
<tr>
<td>0</td>
<td>7</td>
</tr>
</tbody>
</table>

### Performance Characteristics

#### Sensitivity

The limit of detection (LOD) of the DAI Fish ELISA test is 1.4 ppm (cod) for the standard curve.

Validation experiments with common matrices resulted in the following LODs (ppm):

| Wine (red) | 1.5 |
| Soup      | 1.3 |
| Worcester Sauce | 0.3 |
| Asian Sauce       | 2.1 |
| Cracker         | 0.5 |
| Surimi          | 1.8 |
| Spring Roll     | 1.3 |

The limit of quantification (LOQ) of the DAI Fish ELISA test is 4 ppm.

Due to the variety of sample matrices and their influence on the blank, results less than the LOQ should be treated as negative.

#### Cross-reactivity

For the following foods no cross-reactivity could be detected:

- Bean
- Potato
- Rye
- Buckwheat
- Pumpkin seed
- Pork
- Cashew
- Lamb
- Celery
- Egg
- Macadamia
- Mustard
- Pea
- Corn
- Sesame
- Peanut
- Almond
- Shrimp
- Barley
- Milk
- Soy
- Oat
- Brazil nut
- Sunflower seed
- Hazelnut
- Pecan
- Walnut
- Millet
- Pistachio
- Wheat
- Chicken
- Rice
- Onion
- Carrot
- Beef

### Linearity

The serial dilution of spiked samples (Wine, soup, Worcester sauce, Asian sauce, cracker, surimi and spring roll) resulted in a dilution linearity of 89–105%.

### Recovery

Mean recovery was determined by spiking samples with different amounts of cod:

| Wine (red) | 103% |
| Soup      | 117% |
| Worcester Sauce | 112% |
| Asian Sauce       | 103% |
| Cracker         | 99%  |
| Surimi          | 114% |
| Spring Roll     | 93%  |

### References

ISO 13485
ISO 9001

Diagnostic Automation/ Cortez Diagnostics, Inc.
21250 Califa Street, Suite 102 and 116,
Woodland Hills, California 91367 USA

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