



# TECO®REACH Cyprinid Vitellogenin ELISA

For Serum, Plasma, WBH & Mucus

# **REACH** Cyprinid Vitellogenin ELISA

Carp (Caprinus carpio)
Goldfish (Carassius auratus)
Zebrafish (Danio rerio)
Fathead Minnow (Pimephales promelas)

Instructions for Use English

Catalogue No. TE1040 For Research Use Only

## **Symbol Description**



Kit Instructions



Lot Number



Expiry Date



**96** Tests



Storage Temperature



Manufacturer



Intended use



TE1040



Attention

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# **Contents of the TECO®** *REACH* **Cyprinid Vitellogenin ELISA-Kit:**

YMBOL	DESCRIPTION	FORMAT
1	96-well plate coated with cypVTG Antibody	1 plate
	12 break apart strips of 8 wells (12 x 8 in total), in a frame.	
	Ready to use.	
NSB	Wells for Non-Specific Binding (NSB)	1x
	1 break apart strip of 8 wells coated without specific binding antibodies	
S	Standard Stock lyophilized	2 x
	35 ng	
<b>C</b> 1	Control C1 lyophilized	2 x
	Concentration see Certificate of Analysis.	
C2	Control C2 lyophilized	2 x
	Concentration see Certificate of Analysis.	
C3	Control C3 lyophilized	2 x
	Concentration see Certificate of Analysis.	2.0
RS	Inter-Assay Reference Standard Stock lyophilized	1x
	Concentration see Certificate of Analysis	
2	Wash Buffer 50x	1 x 30 ml
	Dilute 1:50 with deionized Water.	
3	Dilution Buffer	1 x 55 ml
	Ready to use.	
4	Matrix Solution	1 x 7 ml
	Ready to use.	
5	Biotinylated Antibody (Biotin-AB)	1 x 12 ml
	Ready to use.	
6	Streptavidin Peroxidase Conjugate (SA- HRP Conj.)	1 x 12 ml
	Ready to use.	
7	TMB Substrate	1 x 12 ml
	Ready to use.	
8	Stop Solution – 1 M HCI	1 x 12 ml
	1 M hydrochloric acid, ready to use.	

## **Storage**

The kit has to be stored at 2-8 °C until expiry date. Do not freeze. Store unused reagents at 2-8 °C.

## Instruction for Use

The TECO®REACH Cyprinid Vitellogenin ELISA Kit is a sensitive enzyme linked immunosorbent assay for the quantitative determination of vitellogenin (VTG) in fish serum, plasma, whole body homogenate (WBH) and mucus. This ELISA kit is specifically designed to fulfill the requirements for the vitellogenin testing in serum or plasma within REACH regualtion as mentioned in EC regulation Nr. 440/2008 (REACH) from July 10th 2015/Document D039048/03 (1).

## **Background**

In oviparous animals, vitellogenin (VTG) is an estrogen induced yolk precursor protein mainly synthesized in the liver to be deposited in the maturing oocytes, where it is split in the yolk proteins lipovitellin 1, lipovitellin 2 and phosvitin. These yolk proteins serve as nourishment storage for the developing embryos. Non-physiological induction of vitellogenin in males or in juvenile fish is thought to indicate an estrogen mediated endocrine disruption. Therefore VTG determination is one of the core endpoints in screening and testing for endocrine disrupting chemicals standardized in the OECD Guidelines for the testing of chemicals for estrogenic activity (2,3,4).

Normally vitellogenin is measured in blood samples or whole body homogenate (WBH) - both sample types require invasive and destructive treatment of the fish. Blood is difficult to collect, in particular where very small fish are concerned, or in approaches where the animals must survive sampling. This is particularly important in field monitoring in order to avoid impact on the population under investigation (6).

Recently, several cell types have been shown to produce VTG after estrogen stimulation, including those of the epidermal mucosa (5). Further studies showed that both VTG and estrogen receptor genes are expressed in epidermal cells. Immunoaffinity and mass fin gerprint analysis showed induction of identical VTG peptides in liver and epidermis (7). VTG contents in the serum demonstrated a similar dose-response pattern in the epidermis and the blood using the TECO®Cyprinid Vitellogenin ELISA (7). Even though the VTG concentration in the skin mucus is an order of magnitude lower than in blood serum or in body homogenates (containing liver tissue), the skin mucosa is very well suited as a matrix to determine exogenous VTG induction caused by environmental chemicals with affinity to estrogen receptors. By using a highly sensitive ELISA in combination with an unique sampling and extraction system the determination of mucosa born VTG determination has the following advantages:

- Simple and highly standardized sampling technique and sample preparation.
- Strictly defined matrix without protease contamination caused by non-target tissues or lymphatic fluid.
- Non-destructive and thereby allowing several subsequent samplings in order to record a kinetic of VTG induction with a maximum known to appear within 1-2 weeks after exposure. Therefore mucosa tests are compatible with acute as well as chronical OECD test methods.
- Epithelial organized epidermis is directly exposed to exogenous estrogens and thereby allowing a direct comparison with in vitro test using estrogen sensitive vitellogenin producing fish cell lines.
- Lower degree of interference with endogenous VTG production (in females) and bio concentration or enterohepatic circulation of the effective estrogen (xenoestrogen) and thereby showing a clear dose response relationship.
- Stability of standards and samples if prescribed storage conditions are observed.

## References

[1] Document D39048/03
COMMISSION REGULATION (EU) No .../.. of XXX amending, for the purpose of its adaptation to technical progress, Regulation (EC) No 440/2008 laying down test methods pursuant to Regulation (EC) No 1907/2006 of the European Parliament and of the Council on the Registration, Evaluation, Authorisation and Restriction of Chemicals (REACH) (Text with EEA relevance) European Commission, July 2015: http://data.consilium.europa.eu/doc/document/ST-10886-2015-ADD-2/en/pdf (2015)

#### OECD (2009), Test No. 229

Fish Short Term Reproduction Assay. OECD Guidelines for the Testing of Chemicals, Section 2, OECD Publishing. OECD (2009), Test No. 230

21-day Fish Assay: A Short-Term Screening for Oestrogenic and Androgenic Activity, and Aromatase Inhibition. OECD Guidelines for the Testing of Chemicals, Section 2, OECD Publishina.

#### OECD (2011), Test No. 234

Tish Sexual Development Test. OECD Guidelines for the Testing of Chemicals, Section 2, OECD Publishing.

[5] Moncaut, N., Lo Nostro, F., Maggese M. C. (2003)

Vitellogenin detection in surface mucus of the South American cichlid fish Cichlasoma dimerus (Heckel, 1840) induced by estradiol-17b. Effects on liver and gonads. Aquatic Toxicology 63, 127-137.

Allner B., Gönna von der S., Griebeler E.M., Nikutowski N., Schaat A., Reproductive functions of wild fish as bioindicators of reproductive toxicants in the

aquatic environment. ESPR Environ. Sci. Pollut. Res., 17, 505-518. Lerche C.F., Schmidt T., Schneider K, Willner M, Stahlschmidt-Allner P. (2016)

Kinetic determination of vitellogenin induction in the epidermis of cyprinid and perciform fishes: Evaluation of sensitive Enzyme-Linked Immunosorbent Assays (ELISAs). Environ Toxicol Chem. 2016 May 6. DOI: 10.1002/etc.3475. [Epub ahead of print]

## **Assay Principle**

The TECO®REACH Cyprinid Vitellogenin ELISA kit is a 96 well homologue immuno-capture ELISA product using homologue antibodies and homologue VTG standard material. Samples are incubated with the vitellogenin specific antibody coated microtiter plate. After unbound material is washed out, a polyclonal biotinylated antibody binds to the vitellogenin. In the following incubation step, a streptavidin-peroxidase conjugate binds to the biotinylated antibody. In the final substrate reaction, the color development is directly proportional to the amount of vitellogenin in the sample. The standard range of the TECO® REACH Cyprinid Vitellogenin EIA Kit is between 0 and 35 ng/ml. In order to avoid additional sample dilution this kit provides an optional standard range extension up to 70 ng/ml.

## Materials required and not supplied

- Pipettes 10 μl 1000 μl
- Multichannel pipettes for 50 μl 100 μl
- Graduated cylinders for reconstituting or diluting reagents
- Manual Aspiration System or automatic washer for ELISA plates
- Aqua dest
- Vortex mixer
- ELISA plate reader suitable for 96 well formats and capable of measuring at 450 nm (Reference: 590-650 nm). For extended standard range: ELISA plate reader suitable for 96 well formats and capable of measuring at 405 nm and 450 nm (Reference: 590-650 nm)
- ELISA plate shaker (500 rpm) (orbital shaker)
- Software package for data generation and analysis

## Fish plasma from male control fish to prepare a fortification sample

For mucus samples: Extraction Buffer and validated Sampling Swabs are not part of this kit. Please order "TECO® Mucus Collection Set, TE1034" separately.

## **Warnings and Precautions**

This kit is for in vitro use by professional persons only.

#### Follow the instructions carefully.

Observe expiration dates stated on the labels and the specified stability for reconstituted reagents. Refer to "Materials Safety Data Sheet" for more detailed safety information. Material of animal origin used in the preparation of this kit has been obtained from animals certified as healthy but these materials should be handled as potentially infectious.

TECOmedical AG is not liable for loss or harm caused by non-observance of the Kit instructions.

- 1. For research use only.
- 2. Treat all specimen samples as potentially biohazardous material. Follow General Precautions when handling contents of this kit.
- 3. Disposal of containers and unused contents should be done in accordance with federal and local regulatory requirements.
- 4. Use the supplied reagents as an integral unit prior to the expiration date indicated on the package label.
- 5. Store assay reagents as indicated.
- 6. Do not use coated strips if pouch is punctured.
- 7. Test each sample in duplicate.
- 8. Use of multichannel pipettes or repeat pipettors is recommended to ensure the timely delivery of liquids.
- 9. a. 1 M hydrochloric acid is caustic and can be harmful for skin, eyes and mucosae.
  - b. Handle TMB with care. Do not ingest. Avoid contact with skin, eyes, or clothing. Should there be any contact, wash with water. If ingested, call a physician.
- 10. A mercury-free preservative is used. Incidental contact with or ingestion of buffer solutions may cause irritation of skin, eyes or mouth. Should there be any contact, wash with water. If ingested, call a physician.

## **Preparation of Reagents**

## 1 Microtiter plate

12 break apart strips of 8 wells (96 in total) in a frame and sealed in a foil bag. Fit strip wells firmly into the frame. After opening, return any unused wells to the original foil package and seal.

Store at 2-8 °C until expiration date.

## NSB Wells for Non-Specific Binding (NSB)

1 break apart strip of 8 wells sealed in a foil bag. Fit strip wells firmly into the frame of the microtiter plate. After opening, return any unused wells to the original foil package and seal.

Store at 2-8 °C until expiration date.

## S Standard Stock - 35 ng

2 vials of lyophilized standard containing stabilized cyprinid vitellogenin.

Store at 2-8 °C until expiration date.

## C Cyprinid Vitellogenin Controls 1, 2 and 3

2 vials each of Control 1, Control 2 and Control 3 containing containing Cyprinid vitellogenin (Concentration see Certificate of Analysis).

Store at 2-8 °C until expiration date.

## RS Inter-Assay Reference Standard Stock

1 vial of lyophilized reference standard containing stabilized cyprinid vitellogenin.

(Concentration see Certificate of Analysis)

Store at 2-8 °C until expiration date.

#### 2 Wash Buffer (50x) concentrated

1 vial of 30 ml Wash Buffer concentrate. Dilute the 50 times concentrate with deionized water up to 1500 ml. The diluted washing solution is stable for 4 weeks at 2-8 °C.

Store undiluted at 2-8 °C until expiration date.

#### 3 Dilution Buffer

1 vial of 55 ml, ready for use. Store at 2-8 °C until expiration date.

#### 4 Matrix Solution

1 vial of 7 ml, ready for use. Store at 2-8 °C until expiration date.

#### **Biotinylated Antibody (Biotin-AB)**

1 vial of 12 ml, ready for use. Store at 2–8 °C until expiration date.

#### 6 Streptavidin Peroxidase Conjugate (SA-HRP Conj.)

1 vial of 12 ml, ready for use. Store at 2-8 °C until expiration date.

## 7 TMB Substrate

1 vial of 12 ml of H<sub>2</sub>O<sub>2</sub>stabilized Tetramethylbenzidine.

Ready for use. Store at 2-8 °C until expiration date.

## 8 Stop Solution – 1 M HCl

1 vial of 12 ml of 1 M hydrochloric acid. Ready for use.

Store at 2-8°C until expiration date.

## **Preparation Standard curve**

Standards have to be prepared freshly before use. Add 500  $\mu$ l **Dilution Buffer** 3 to the vial. Let it stand for 15-30 minutes and then vortex.

Now, the Standard stock contains 70 ng/ml of stabilized cyprinid vitellogenin (Standard A, Std A).

Preparation of the standard curve with **Dilution Buffer 3**:

ID	Concentration ng/ml	Dilution Buffer μΙ	Standard solution
Std A	70	Add 500 μl	
Std B	35	200	200 μl Std A
Std C	11.7	200	100 μl Std B
Std D	3.9	200	100 μl Std C
Std E	1.3	200	100 μl Std D
Std F	0.4	200	100 μl Std E
Std G	0	200	

## **Preparation Kit Controls**

Cyprinid Vitellogenin Controls 1, 2 and 3 have to be prepared freshly before use. Add 500 µl **Dilution Buffer** to the vial. Let it stand for 15-30 minutes and then vortex.

Reconstituted and diluted standards and controls are for single use only and are stable for 4 hours at room temperature (20-28°C). Do not store reconstituted standards and controls.

## **Preparation Fortification Sample**

According to the EC regulation Nr. 440/2008 (REACH) from July 10th 2015/ Document D039048/03 [1] the fortification sample should be prepared by spiking plasma from male control fish with a defined concentration of vitellogenin. Dependent on fish species, the volume of plasma obtained from one or more male control fish might be limiting and from ethical point of view critical. For this reason we recommend to add suitable amount of diluted plasma samples (e.g. 1:10 000) of male control fish to the Inter-Assay Reference Stock vial resulting in a concentration which should exceed the vitellogenin levels of the male control fish by factor 10 to 100. Let the vial stand for 15-30 minutes and then vortex. Final plasma dilution and concentration of the fortification sample are dependent on the fish species used in exposition experiment. Examples given in following table:

Plasma Matrix (Diluted plasma sample from male fish)	Fortification sample Final concentration in ng/ml
Add 200 μl	25
Add 500 μl	10
Add 1000 μl	5

## **Preparation and Stability of Samples**

#### **Preparation of Samples**

#### **Serum** or Plasma

Store fresh samples immediately after collection at -20°C or lower until assayed.

Recommended sample thawing: A simple and fast method is to place the frozen serum or plasma samples in normal tap cold water (15-20°C). They should be thawed within 10 to 15 minutes. For assay, samples should be pre-diluted dependent on fish species with **Dilution Buffer 3**, e.g 1: 10 000 (two times 1:100). Optimal sample dilution for cyprinid species may differ. Mature female fish may have elevated vitellogenin.

#### Whole Body Homogenate (WBH)

Store fresh WBH samples immediately after preparation below -20°C until assayed. For assay, WBH samples should be pre-diluted dependent on fish species with **Dilution Buffer 3**, e.g 1: 10 000 to 1: 1 000 000.

#### Mucus

Collect mucus as described in the TECO® Mucus Collection Set TE1034.

For assay, add 500 µl Extraction Buffer (**TECO® Mucus Collection Set, TE1034**) to the swab 15-30 min before vortex. For more determinations (e.g. total protein, Cortisol etc.) the swabs should be removed from each vial and discared before vitellogenin measurement. Before pipetting sample into wells repeat vortexing the sample. In most studies, this sample dilution should be used for sample measurements as a matter of routine. For dose response curves or if concentrations of estrogenic induced fish are required, a further pre-dilution of mucus samples of 1:10 to 1:100 with **Dilution Buffer** 3 may be necessary.

#### Sample Stability (Mucus samples from Cyprinids)

Mucus-containing swabs can be stored several months at <-20°C. After addition of Extraction Buffer the samples are stable up to 4 hours at room temperature (20-28°C).

#### Stability of sample vitellogenin

Avoid repeated freeze/thaw cycles.

## Correction of vitellogenin results by the protein concentration

Independently from the assay procedure, various factors may influence the final amount of biological samples added into the Vitellogenin ELISA (e.g. total amount of blood collected into the prefilled sample tubes; effectiveness of homogenization; amount of mucus on the swab etc.). In order to obtain the correct analytical result, all samples may be corrected by the protein concentration by using in parallel a colorimetric protein determination. The Dilution Buffer and the Extraction Buffers in the VTG kits are protein free and may be used as Standard buffers and for sample dilution in the protein assays. This sample dilution may differ from the optimal sample dilution in the vitellogenin assay.

## **Assay Procedure**

All determinations (NSB, standards, fortification sample, controls, diluted male control samples and samples) should be assayed in duplicate. When performing the assay, the standards, controls and samples should be pipetted as fast as possible (<15 minutes). To avoid distortions due to differences in incubation times, HRP Conjugate, Substrate Solution and Stop Solution should be added to the plate in the same order and with the same time interval as the samples. A multichannel pipette is essential. Allow all reagents to stand at room temperature (20–28°C) for at least 30 minutes. During all incubation steps, plates should be sealed with the adhesive foil or a plastic cover. For light protection, incubate in a dark chamber or cover plate with aluminum foil.

- 1. Allocate the wells of the Microtiter plate 1 for NSB, standards, controls, fortification sample, diluted male control sample and samples.
- 2. Pipette 50 µl Matrix solution 4 (multichannel pipette) into each well.
- 3a. Standard range (0-35 ng/ml): Add 50 μl of each prepared standard ( B G ), prepared controls ( C1 and C2 ), prepared fortification sample, the diluted male plasma sample (used to prepare the fortification sample) and (pre-diluted) samples into the corresponding wells.
- 3b. Extended standard range (0-70 ng/ml): Add 50 μl of each prepared standard ( A G ), prepared controls ( C1 , C2 and C3 ), prepared fortification sample, the diluted male plasma sample (used to prepare the fortification sample) and (pre-diluted) samples into the corresponding wells.
- 4. Cover the wells and incubate the plate for  $120 \pm 10$  min at room temperature (20–28°C) on a shaker (500 rpm).
- 5. After incubation, aspirate the contents of the wells and wash 5 times with 350 μl diluted Wash Buffer 2. The use of an automatic plate washer is recommended.
- 6. Following the last washing step, pipette 100  $\mu$ l of the Biotinylated AB 5 in each well (multichannel pipette).
- 7. Cover the wells and incubate the plate for  $60 \pm 5$  min at room temperature ( $20-28^{\circ}$ C) on a shaker (500 rpm).
- 8. After incubation, wash the wells 5 times with Wash Buffer as described in step 5.
- 9. Following the last washing step, pipette 100 μl of the SA-HRP Conjugate 6 in each well (multichannel pipette).
- 10. Cover the wells and incubate the plate for  $30 \pm 5$  min at room temperature ( $20-28^{\circ}$ C) on a shaker (500 rpm).
- 11. After incubation, wash the wells 5 times with Wash Buffer as described in step 5.
- 12. Pipette 100 μl of the TMB Substrate Solution 7 in each well (multichannel pipette).
- 13. Incubate the plate for 15-30 min, in the dark, at room temperature (20–28°C) on a shaker (500 rpm).
- 14. Stop the reaction by adding 100 μl of Stop Solution 8 (multichannel pipette).
- 15a. Standard range (0-35 ng/ml): Measure color reaction within 10 minutes at 450 nm (reference filter between 590-650 nm).
- 15b. Extended standard range (0-70 ng/ml): Measure color reaction within 10 minutes at 450 nm (reference filter between 590-650 nm) and at 405 nm (reference filter 590-650 nm). First, calculate the sample values between 0 and 35 ng/ml (using Std G Std B) by using 450 nm. Thereafter, values between 35 and 70 ng/ml should be calculated using 405 nm reading (using Std G and Std E- Std A).

## **Result Analysis**

#### **Establishing the Standard Curve**

A calibration curve can be established by plotting standard concentration on the x-axis (linear scale) against the absorbance of the standards on the y-axis (linear scale). The vitellogenin concentrations in mucus can then be read off the calibration curve.

A 4-parameter curve fit should be used for automatic data reduction. If samples were pre-diluted, the concentration will be obtained by multiplying the value read off the calibration curve by the dilution factor. There is no dilution correction for mucus necessary, if the 0.5 ml Extraction buffer is added to the swab. Samples with higher absorbance values than standard A should be tested again pre-diluted with Dilution Buffer. This additional dilution has to be taken in account for the concentration calculation.

#### **Typical Results**

(Example only, not for use in calculation of actual results.)

Sample	Conc. ng/ml	OD 450 nm	OD 405 nm
Std A	70.0	-	1.058
Std B	35.0	2.090	0.656
Std C	11.7	0.739	0.241
Std D	3.9	0.295	0.103
Std E	1.3	0.120	0.049
Std F	0.4	0.061	-
Std G	0	0.019	0.017

Table 1 Reader values of a typical standard curve.

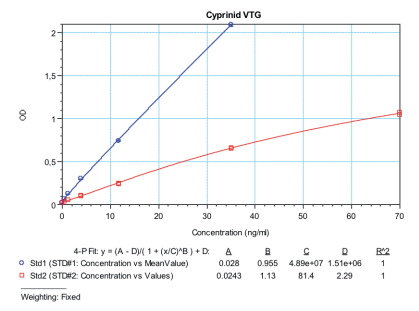


Figure 1 Standard curves by using a 4-parameter curve fit (4-PL) using measurement at 450 nm and 405 nm.

## Recovery of fortification sample

The fortification sample should be determined together with the diluted male plasma used for spiking each day vitellogenin assays are performed in the laboratory. The corresponding recovery has to be reported together with the results obtained at that specific day.

	Measured concentration of male control plasma ng/ml	Added Inter-Assay Reference Standard* ng/ml	Expected concentration	Measured concentration	Recovery in % Measured/Expected Concentration x 100
			ng/ml	ng/ml	
Example	0,6	5,0	5,6	5,8	5,8/5,6 x 100 = 103,6

<sup>\*</sup>see table in chapter "Preparation Fortification Sample"

## **Test Performance**

Mucus vitellogenin concentration are expressed in ng/ml swab extract (0.5 ml buffer/swab). For serum, plasma or WBH samples, concentration has been corrected by the pre-dilution.

Standard range: 0.4 ng/ml – 35 ng/ml (undiluted samples) Extended standard range: 0.4 ng/ml - 70 ng/ml (undiluted samples)

LLOQ < 0.4 ng/ml LLD 0.036 ng/ml

The LLD (lower limit of detection) is defined as the corresponding concentration of the mean OD zero standard plus 3 SD.

The mean coefficient of determination (R2) of 10 standard curves was 1.0.

Sample	Mean	SD	cv
#1	2,1	0,08	3,6
#2	17,1	0,44	2,6

Table 2 Intra-assay coefficient of variation (CV).

Sample	Mean	SD	cv
Control 1	2,1	0,13	6,1
Control 2	16,9	0,50	3,0

*Table 3 Inter-assay coefficient of variation (CV).* 

		Before addition	Added	Expected	Measured	Recovery	Mean	SD
Species	Sample	ng/ml	ng/ml	ng/ml	ng/ml	%	%	%
Fathead	1	0,1	6,1	6,2	7,2	116	112	3,3
minnow	2	0,1	6,1	6,2	7,0	113		
	3	1,8	6,1	7,9	8,8	111		
	4	0,0	6,1	6,1	6,6	108		
Zebrafish	6	0,0	6,1	6,1	6,8	111	110	4,0
	7	0,0	6,1	6,1	7,0	115		
	8	0,0	6,1	6,1	6,5	107		
	9	0,0	6,1	6,1	6,5	107		

Table 4
Recovery of vitellogenin
spiked to mucus samples of
untreated cyprinids.

			Day 0		Day 4				
		Measured		Concentration		Measured		Concentration	Dilution
Species	Sample	ng/ml	Dilution	ng/ml	Sample	ng/ml	Dilution	ng/ml	recovery (%)
Fathead	1	0,08	1	0,08	16	0,15	10	1,5	
minnow	2	0,09	1	0,09		0,03	100	2,6	n/a
	3	2,20	1	2,20	17	45,07	10	450,7	
	4	0,08	1	0,08		4,44	100	443,5	98
	5	0,20	1	0,20	18	15,65	10	156,5	
	20	0,28	1	0,30		1,49	100	149,2	95
					19	44,12	10	441,1	
						4,48	100	447,5	101
					20.1	2,35	10	23,5	
						0,26	100	25,8	110
			Day 0				Day 4		
		Measured		Concentration		Measured		Concentration	Dilution
Species	Sample								
		ng/ml	Dilution	ng/ml	Sample	ng/ml	Dilution	ng/ml	recovery (%)
Zebrafish	6	0,01	Dilution 1	ng/ml 0,01	Sample 21	ng/ml 16,44	Dilution 10	ng/ml 164,4	recovery (%)
Zebrafish	6 7								recovery (%)
Zebrafish	-	0,01	1	0,01		16,44	10	164,4	• • • • • • • • • • • • • • • • • • • •
Zebrafish	7	0,01 0,01	1	0,01 0,01	21	16,44 1,77	10 100	164,4 177,4	
Zebrafish	7 8	0,01 0,01 0,01	1 1 1	0,01 0,01 0,01	21	16,44 1,77 11,58	10 100 10	164,4 177,4 115,8	108
Zebrafish	7 8 9	0,01 0,01 0,01 0,01	1 1 1 1	0,01 0,01 0,01 0,01	21	16,44 1,77 11,58 1,23	10 100 10 100	164,4 177,4 115,8 122,6	108
Zebrafish	7 8 9	0,01 0,01 0,01 0,01	1 1 1 1	0,01 0,01 0,01 0,01	21	16,44 1,77 11,58 1,23 9,56	10 100 10 100 100	164,4 177,4 115,8 122,6 95,6	108
Zebrafish	7 8 9	0,01 0,01 0,01 0,01	1 1 1 1	0,01 0,01 0,01 0,01	22 23	16,44 1,77 11,58 1,23 9,56 0,98	10 100 10 100 100 10	164,4 177,4 115,8 122,6 95,6 98,1	108

25

3.56

0,35

10

100

35.6

34,6

97

Table 5
Treatment effect on mucus vitellogenin and dilution linearity in two groups (day 0) and (day 4) of estradiol treatment (1 µg/l) in cyprinid species.

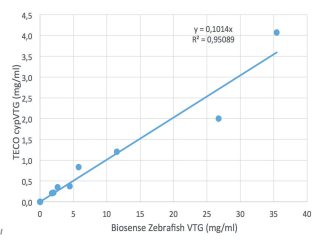


Figure 2
Relation between a commercial ELISA kit and TECO
Cyprinid Vitellogenin ELISA. 13 serum samples
obtained from Zebrafish were measured according
to the corresponding kit instructions.

## **Bisphenol A (BPA) treatment experiments**

## **BPA treatment in Fat head minnow**

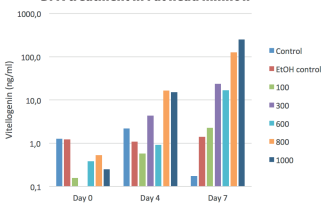


Figure 3
Mucus of five to six individuals per treatment
group were maintained per aquarium and
were exposed to up to 1000 µg BPA/I.

#### **BPA** treatment in male zebrafish

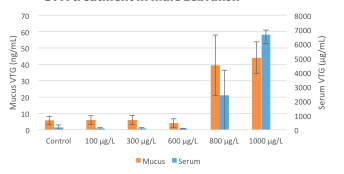


Figure 4
Vitellogenin in Mucus and serum of male zebrafish (2-5 fish per group) after 7 days of different BPA treatments.

## Normal fish vitellogenin concentrations

## **Zebrafish Specimens**



Figure 5
Vitellogenin in different tissues, serum and mucus of a male and a female zebrafish.
Tissue extracts were obtained according to OECD guidelines.

Please note, that mucus concentrations are expressed in ng/ml.

More data on file.

Notes		

# **TECO® REACH Cyprinid Vitellogenin**

**Mucus sample preparation: Quick Guide** 

Mucus samples have to be collected using the validated TECO Mucus Collection Set TE1034. This collection set also contains the required Extraction Buffer for sample extraction before assay run.

## **Procedure**

Take frozen sample swab tips (vials) selected for testing in the Vitellogenin ELISA out of the freezer.

Open all vials.

Add **500** µl Extraction Buffer\* (TECO Mucus Collection Set, TE1034) into each vial and wait **15-30 min.** 

Vortex the closed vials extensively.

For more determinations (e.g. total protein, Cortisol etc.) remove the swabs before vitellogenin assay and discard.

Before pipetting repeat vortexing the sample.

| Please read Kit instruction before using the Quick Guide

\*If necessary, the sensitivity of the vitellogenin determination may be increased by using 250  $\mu$ l instead of 500  $\mu$ l Extraction Buffer (TECO Mucus Collection Set, TE1034) into each vial and wait for 15-30 min. In order to correct the dilution factor, divide the final result obtained from the standard curve by factor 2.

# **TECO®** REACH Cyprinid Vitellogenin

**Quick Guide: Standard Range (0-35 ng/ml)** 

# Prepare Standards, Controls, Diluted Male Control Sample, Fortification Sample and Samples

- Add NSB wells into the microtiter plate frame
- Dilute Wash Buffer concentrate 1:50 with destilled water
- Allow all reagents to stand at room temperature (20-28°C) for at least 30 minutes

## **Assay Procedure**

Add **50 µL** Matrix solution **4** into each well (multichannel pipette)

Add **50 µL** of each standard **B** - **G** , prepared controls (**C1**, **C2**), fortification sample, diluted male control sample and (pre-diluted) samples into wells

Add **50 µL** Dilution Buffer into **NSB** wells

Incubate plate for 120±10 minutes on a shaker (500rpm) at RT (20-28°C)

Wash plate **5 times** using Wash Buffer **2** 

Add **100 µL** Biotinylated AB **5** in each well (multichannel pipette)

Incubate plate for 60±5 minutes on a shaker (500rpm) at RT (20-28°C)

Wash plate **5 times** using Wash Buffer **2** 

Add **100 µL** SA-HRP-Conjugate 6 in each well (multichannel pipette)

Incubate plate for 30±5 minuten on a shaker (500rpm) at RT (20-28°C)

Wash plate **5 times** using Wash Buffer **2** 

Add **100 µL** TMB Substrate Solution 7 in each well (multichannel pipette)

Incubate the plate for 15-30 minutes, in the dark, on a shaker (500rpm) at RT (20-28°C)

Add **100 µL** Stop Solution **8** in each well (multichannel pipette)

Measure the color reaction within 10 minutes at 450nm (reference filter between 560 - 650 nm)

Calculate the sample values between 0 and 35 ng/ml by using Std G- Std B A 4-parameter curve fit should be used for automatic data reduction

# **TECO®** REACH Cyprinid Vitellogenin

**QuickGuide: Extended Standard Range (0-70 ng/ml)** 

# Prepare Standards, Controls, Diluted Male Control Sample, Fortification Sample and Samples

- Add NSB wells into microtiter plate frame
- Dilute Wash Buffer concentrate 1:50 with destilled water
- Allow all reagents to stand at room temperature (20-28°C) for at least 30 minutes

## **Assay Procedure**

Add **50 µL** Matrix solution **4** into each well (multichannel pipette)

Add **50 µL** of each standard **A** - **G** , prepared controls (**C1**, **C2**, **C3**), fortification sample, diluted male control sample and (pre-diluted) samples into wells Add **50 µL** Dilution Buffer into **NSB** wells

Incubate plate for 120±10 minutes on a shaker (500rpm) at RT (20-28°C)

Wash plate **5 times** using Wash Buffer **2** 

Add **100 µL** Biotinylated AB **5** in each well (multichannel pipette)

Incubate plate for 60±5 minutes on a shaker (500rpm) at RT (20-28°C)

Wash plate **5 times** using Wash Buffer **2** 

Add **100 µL** SA-HRP-Conjugate **6** in each well (multichannel pipette)

Incubate plate for 30±5 minuten on a shaker (500rpm) at RT (20-28°C)

Wash plate **5 times** using Wash Buffer **2** 

Add **100 µL** TMB Substrate Solution 7 in each well (multichannel pipette)

Incubate the plate for 15-30 minutes, in the dark, on a shaker (500rpm) at RT (20-28°C)

Add **100 µL** Stop Solution **8** in each well (multichannel pipette)

Measure the color reaction within 10 minutes at 450nm and 405 nm (reference filter between 560 - 650 nm).

Using 450 nm: Calculate the sample values between 0 and 35 ng/ml by using Std G- Std B Using 405 nm: Calculate the sample values between 35 and 70 ng/ml by using Std G and Std E- Std A.

A 4-parameter curve fit should be used for automatic data reduction