

Enzyme Research Laboratories

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Matched-Pair Antibody Set for ELISA of Canine Factor IX antigen (CFIX)

Sufficient reagent for 4 x 96 well plates

Product #: CFIX-EIA
Lot #: Sample
Expiry Date: Sample

Store at -10 to -20°C

For Research Use Only Not for use in diagnostic procedures.

Description of Factor IX (FIX)

Factor IX (FIX, Christmas Factor) is a vitamin K-dependent glycoprotein produced in the liver and circulates in human plasma at a concentration of 5 µg/ml (87 nM). The biological importance of FIX is demonstrated in Haemophilia B (Christmas disease), an X-linked congenital bleeding disease resulting from a quantitative (low activity and low antigen) or qualitative (low activity and normal antigen) defect in FIX function.

In its zymogen form FIX is a single chain molecule of 55,000 daltons. It contains two EGF-like domains and an amino-terminal domain containing 12 γ -carboxy-glutamic acid (Gla) residues. These Gla residues allow FIX to bind divalent metal ions and participate in calcium-dependent binding interactions. The activation of FIX occurs by limited proteolysis in the presence of calcium by activated factor XI (FXIa) and/or by a complex of VIIa/tissue factor/phospholipid and activated Factor X between residues Arg^{146} -Ala 147 and between Arg^{180} -Val 181 . The terminal activated product in either case is F.IXa_{\beta}, a two-chain enzyme consisting of a heavy chain (28,000 daltons), a light chain (18,000 daltons) and an activation peptide product of 11,000 daltons. FIX can also be cleaved into inactive products by thrombin and by elastase. The activity of FIXa_{\beta} in plasma is inhibited by antithrombin and this inhibition is accelerated approximately 1000-fold in the presence of optimal concentrations of heparin $^{1\cdot3}$.

Principle of Sandwich-style ELISA

Purified antibody to FIX is coated onto the wells of a microtitre plate. The plates are washed and plasma or other fluids containing canine FIX are applied. The coated antibody will capture the canine FIX in the sample. After washing the plate to remove unbound material, a peroxidase conjugated second antibody to FIX is added to the plate to bind to the captured canine FIX. After washing the plate to remove unbound conjugated antibody, the peroxidase activity is expressed by incubation with o-phenylenediamine (OPD). After a fixed development time the reaction is quenched with the addition of $\rm H_2SO_4$ and the colour produced is quantified using a microplate reader. The colour generated is proportional to the concentration of canine FIX present in the sample.

Supplied Materials:

- **1. Capture Antibody (CFIX-EIA-C):** One yellow-capped vial containing 0.4 ml of polyclonal anti-FIX antibody for coating plates.
- **2. Detecting Antibody (CFIX-EIA-D):** One red-capped vial containing 0.4 ml of peroxidase-conjugated polyclonal anti-FIX antibody for detection of captured FIX.

Note: Antibodies are supplied in a 50% (v/v) glycerol solution for storage at -10 to -20°C. Keep vials tightly capped. Do not store in frost-free freezers.

Materials Required but not Provided:

- 1. Coating Buffer: 50 mM Carbonate
- 1.59g of Na_2CO_3 and 2.93g of $NaHCO_3$ up to 1 litre. Adjust pH to 9.6. Store at 2-8°C up to 1 month.
- 2. PBS: (base for wash buffer)

8.0g NaCl, 1.15g Na₂HPO₄, 0.2g KH₂PO₄ and 0.2g KCl, up to 1 litre. Adjust pH to 7.4, if necessary. Store up to 1 month at $2-8^{\circ}$ C, discard if there is evidence of microbial growth.

3. Wash Buffer: PBS-Tween (0.1%,v/v) To 1 litre of PBS add 1.0 ml of Tween-20. Check that the pH is 7.4. Store at 2-8°C up to 1 week.

4. Sample Diluent: HBS-BSA-EDTA-T20

5.95g HEPES (free acid), 1.46 g NaCl, **0.93 g Na₂EDTA**, 2.5 g Bovine Serum Albumin (Sigma, RIA grade) dissolved in 200 ml H_2O . Add 0.25 ml of Tween-20, check and adjust pH to 7.2 with NaOH, then make up to a final volume of 250 ml with H_2O . Aliquot and store frozen at -20°C.

- **5. Substrate Buffer:** Citrate-Phosphate buffer pH 5.0 2.6g Citric acid and 6.9g Na_2HPO_4 up to a final volume of 500 ml with purified H_2O . Store at 2-8°C up to 1 month.
- **6. OPD Substrate:** (o-Phenylenediamine.2HCl) $\underline{\text{Toxic!}}$ (5mg tablets: Sigma # P-6912). Make up immediately before use. Dissolve 5mg OPD in 12 ml substrate buffer then add 12 μ l 30% H₂O₂. Do not store.
- 7. Stopping Solution: 2.5 M H₂SO₄

<u>Caution: VERY CORROSIVE!</u> <u>GENERATES HEAT ON DILUTION!</u> Where stock sulphuric acid is 18 Molar, add 13.9 ml to 86 ml H_2O . Store at room temperature.

8. Other:

Microplates, 96-well Immulon 4-HBX Microplate washer (optional) Microplate reader.

Assay Procedure:

1. Coating of plates:

Dilute the capture antibody 1/100 in coating buffer (preferably in a polypropylene tube) and immediately add 100 μ l to every well in the plate. Incubate 2 hrs @ 22°C or overnight at 2-8°C.

2. Blocking:

Blocking is not required under the conditions described. Washing the plate with PBS-Tween is sufficient to block non-specific interactions.

Wash plate X 3 with wash buffer.

3. Samples:

Canine plasma is diluted 1/40 (100%) then serial 1/2's down to 1/1280 (3.13%). Sample plasmas are diluted 1/40 & 1/80. All dilutions are made in HBS-BSA-EDTA-T20 sample diluent. Apply 100 μ l to each well and incubate plate @ 22°C for 60 minutes. Wash plate X 3 with wash buffer.

4. Detecting Antibody:

Dilute the detecting antibody 1/100 in HBS-BSA-EDTA-T20 sample diluent and apply $100 \mu l$ to each well. Incubate plate @ $22^{\circ}C$ for 60 minutes. Wash plate X 3 with wash buffer.

5. OPD Substrate:

Apply 100 μ l of freshly prepared OPD substrate to every well. Allow colour to develop for 10-15 minutes then stop colour reaction with the addition of 50 μ l/well of 2.5 M H₂SO₄. The plate can be read at wavelength of 490 nm.

Calculation of Results:

The construction of a proper reference curve is of no less importance than any other aspect of the assay. A reference curve should be constructed by plotting the known concentration of standards versus absorbance. This can be done manually using graph paper, or by using curve-fitting computer software. In our experience, the dose response curves of most immunoassays tend to be sigmoid in shape. Although linear regions can be identified within the curve, the best overall fit is often obtained using an algorithm that provides a weighted theoretical model of fit throughout the entire curve, such as a 4-parameter or 5-parameter logistic curve fit ^{4,5}. In general, the simplest model that defines the concentration-response relationship should be used ⁶.

The "back-fit" test is a simple and reliable method to determine if a curve-fitting method is appropriate. In this test, the apparent concentrations for the absorbance values of each standard point are read from the reference curve. The derived values are compared to the assigned values. An appropriate curve fitting method will produce derived values that closely match assigned values throughout the range of the curve, within user-defined limits⁶. The coefficient of determination (R²) is a valuable indicator of the overall fit, but should not be used by itself in the selection of a curve fitting method, as a poor fit in a particular region of the

curve may not be evident from this value alone 5,6

In the quality control of this product we have determined that under the conditions described above, a reference curve that is constructed using serial dilutions of canine plasma, will produce a correlation coefficient (R²) of at least 0.980 using a semi-log fit, and an R² of at least 0.990 using a 4-parameter logistic curve fit algorithm. However, the performance characteristics of in-house assays developed using this product in other laboratories may vary slightly from ours. Different curve fitting methods may be employed but we recommend that the back-fit test be applied as evidence that the fitting method is appropriate.

Technical Notes:

- This paired antibody product is intended to facilitate the end user in establishing an in-house immunoassay for research purposes only. It must not be used for diagnostic applications. Assay validation is the responsibility of the end user and should be done according to user-defined protocols⁶.
- Reference calibrators should be of the same matrix and anticoagulant as the samples to be tested (example serum or plasma, citrate or EDTA)
- Do not use samples diluted less than 1/4, as falsely high readings may result.
- The optimal colour development time should be determined empirically as the time required to obtain an absorbance of at least 1.000 at 490 nm for the 100% reference point, not to exceed 20 minutes.
- Rheumatoid factor in samples may interfere in ELISA by binding to the capture and/or detecting antibodies.
- The wells should not be allowed to become dry. Keep plate covered or in a humid chamber during incubations.
- Antibodies are supplied in a 50% glycerol solution and can be centrifuged briefly in a micro-centrifuge to gather residual reagent from the cap and walls of the tube.

References:

- 1. Lawson, JH, Mann KG; Cooperative Activation of Human F.IX by the Human Extrinsic Pathway of Coagulation; JBC 266 pp11317-11327, 1991
 2. Enfield DL, Thompson AR; Cleavage and Activation of Factor IX by Serine
- Enfield DL, Thompson AR; Cleavage and Activation of Factor IX by Serine Proteases; Blood 64, pp 821-831, 1984
- **3.** Limentani SA, Furie BC, Furie B, in Hemostasis and Thrombosis, 3rd Edition, eds. RW Colman, J Hirsh, VJ Marder and EW Salzman, pp. 94-108, J.B. Lippincott Co., Philadelphia PA, USA, 1994.
- **4.** Nix,B, Wild D, in Immunoassays, A Practical Approach, editor J.P. Gosling, pp. 239-261, Oxford University Press, 2000.
- **5.** NCCLS. Evaluation of the Linearity of Quantitative Analytical Methods; Proposed Guidline Second Edition. NCCLS Document EP6-P2 (ISBN 1-56238-446-5, NCCLS, Wayne, Pennsylvania USA, 2001
- **6.** FDA Guidance for Industry. Bioanalytical Method Validation; May 2001, available on the internet: www.fda.gov/cder/guidance/index.htm