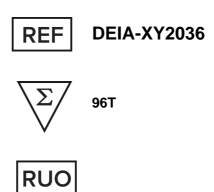




9-HODE ELISA kit



This product is for research use only and is not intended for diagnostic use.

For illustrative purposes only. To perform the assay the instructions for use provided with the kit have to be used.

Creative Diagnostics

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PRODUCT INFORMATION

Intended Use

For detection of 9-HODE in test samples.

General Description

Linoleic acid, the predominant polyunsaturated fatty acid (PUFA) in the human diet, can be metabolized by cyclooxygenase, lipoxygenase and P450 enzymes. The hydroxyoctadecadienoic acid (HODE) derivatives of linoleic acid, 9(R)-HODE, 9(S)-HODE and 13(S)-HODE, are the most widely distributed of the known linoleic acid metabolites. These compounds exhibit interesting biologic activities, including the regulation of platelet function, maintenance of vascular thromboresistance and transduction of the cellular responses to certain growth factors. HODE derivatives may also influence certain pathological states including psoriasis, the development of atherosclerosis and the development of cancer. This assay measures the level of total 9-HODE, which includes both 9(S)-HODE and 9(R)-HODE, in biological samples.

Principles of Testing

This kit is a competitive enzyme-linked immunoassay (ELISA). Briefly, the 9-HODE present in the samples or standards competes with 9(±)-HODE conjugated to horseradish peroxidase [9(±)-HODE HRP] for binding to an antibody specific for 9(±)-HODE that is precoated on a microplate. The peroxidase activity of 9(±)-HODE-HRP results in color development when a substrate is added. The intensity of the color is proportional to the amount of 9(±)-HODE-HRP bound and is inversely proportional to the amount of unconjugated 9-HODE present in the samples or standards.

Reagents And Materials Provided

- 1. Coated Plate: 96-well microplate coated with anti-9(±)-HODE antibody. 1 plate, 4°C.
- 2. **9(±)-HODE Standard:** 0.05 mg/mL 9(±)-HODE standard solution in ethanol. 20 μ L, 4°C.
- 3. **5x Wash Buffer:** Buffer used to wash the plate. 50 mL, 4°C.
- 4. 5x Dilution Buffer: Buffer used for diluting kit components and samples. 25 mL, 4°C.
- 5. **TMB Substrate:** TMB substrate used for color development. 25 mL, 4°C.
- 6. 9(±)-HODE-HRP Conjugate: 9(±)-HODE horseradish peroxidase concentrated conjugate. 320 μL, 4°C.

Materials Required But Not Supplied

- 1. Microplate reader with a 450 nm filter
- 2. Adjustable micropipettes (10 – 1000 μL) and tips
- 3. Deionized water
- 3 N Sulfuric Acid (H₂SO₄)

EXTRACTION MATERIALS

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- 1. Chloroform (CHCl₃)
- 2. Butylated Hydroxytoluene (BHT)
- 3. Triphenyl Phosphine (TPP)
- 4. Magnesium Chloride (MgCl₂)
- 5. Potassium Hydroxide (KOH)
- pH 3 Water 6.
- 7. Methanol (MeOH)
- 8. Sodium Chloride (NaCl)
- 9. Ethyl Acetate
- 10. HCI (concentrated and 1 N)
- 11. Nitrogen Gas (N₂)

Storage

- 1. Store the components of this kit at the temperatures specified on the labels.
- 2. Unopened reagents are stable until the indicated kit expiration date.
- 3. Desiccant bag must remain in foil pouch with unused strips. Keep pouch sealed when not in use to maintain a dry environment. Remove excess air before sealing

Specimen Collection And Preparation

- Urine can be assayed after diluting with Dilution Buffer.
- Plasma and most other mediums will need to be extracted.

EXTRACTION REAGENTS NEEDED

- Folch Solution + 0.005% BHT (w/v) + 0.05% TPP (w/v): 2:1 CHCl₃ :MeOH 1.
- 2. Folch Solution + 0.005% BHT (w/v): 2:1 CHCl3: MeOH
- 3. 0.43% MgCl₂: 0.43% MgCl₂ (w/v) in deionized water
- 4. MeOH + 0.005% BHT (w/v)
- 15% KOH: 15% KOH (w/v) in deionized water 5.
- 6. pH 3 Water: deionized water brought to a pH of 3 with HCl
- 0.9% NaCl: 0.9% NaCl (w/v) in deionized water 7.

EXTRACTION PROTOCOL

Note: This is a suggested protocol. Varying compositions of biological fluids may alter extraction efficiency. It is therefore important to measure the 9(±)-HODE concentration of a parallel "spike" sample (i.e. a biological sample to which a known amount of 9(±)-HODE is added prior to extraction) in order to determine extraction efficiency.

Freeing Esterified 9-HODE

From Plasma or Other Fluids:

Add 20 mL of Folch Solution + 0.005% BHT + 0.05% TPP to a 50 mL conical tube and place on ice.

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- 2. Add 1 mL of plasma or other fluid.
- 3. Shake or vortex well for 1 minute.
- Add 10 mL of ice cold 0.43% MgCl₂ and shake or vortex well for 1 minute. 4.
- Centrifuge for 3 minutes at 2500 xg and 4°C. 5.
- 6. Discard the top layer and transfer the bottom organic layer to a new 50 mL tube, being careful not to transfer any protein layer that may be present.
- 7. Evaporate the organic layer under N₂.
- Add 0.5-2 mL of MeOH + 0.005% BHT (depending on the amount of lipid present) and an equal volume of 8. 15% KOH, swirling after each addition.
- Incubate the sample at 37°C for 30 minutes. 9.
- 10. Adjust to pH 3 with 1 N HCl using approximately 2.5 times the volume of 15% KOH that was added.
- 11. Dilute with pH 3 Water so that the volume of MeOH added is ≤5% of the total volume. The sample is now ready for liquid phase extraction as described below.

From Tissue Samples:

- Add 20 mL of Folch Solution + 0.005% BHT to a 40 mL flat bottom tube and place on ice.
- 2. Weigh 0.5 to 1 gram of tissue and add to tube on ice.
- Shake or vortex well for 1 minute. 3.
- 4. Homogenize with a blade homogenizer or sonicator for 30 seconds.
- 5. Allow to stand under N2 in a sealed tube for one hour at room temperature, vortexing occasionally.
- 6. Add 4 mL of 0.9% NaCl.
- Vortex vigorously and centrifuge for 3 minutes at 2500 x g and 4°C.
- Discard the upper layer and transfer the lower phase to a new 50 mL conical tube, carefully avoiding the protein layer.
- Evaporate under N2.
- 10. Add 2-4 mL of MeOH + 0.005% BHT and an equal volume of 15% KOH, swirling after each addition.
- 11. Incubate at 37°C for 30 minutes.
- 12. Adjust to pH 3 with 1 N HCl using approximately 2.5 times the volume of 15% KOH that was added.
- 13. Dilute to 40–80 mL with pH 3 Water so the MeOH is ≤5% of the total volume. The sample is now ready for liquid phase extraction as described below.

Extraction from Plasma, Serum or Tissue Culture Medium

- Acidify to pH 3 with concentrated HCl.
- Extract with 3x the sample volume of water saturated Ethyl Acetate. Centrifuge at a low speed or allow to 2. stand until phases separate.
- Remove the organic (upper) phase and transfer it to a new container, being careful not to contaminate it with the aqueous phase.
- Repeat steps 2 and 3, combining the organic phases with that from the first extraction. 4.
- 5. Evaporate completely under N2 or in a centrifugal evaporator.

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Bring samples up in 25 µL Methanol, then add 975 µL Dilution Buffer. (If solubility is a problem, it may be necessary to increase the pH.)

NOTE: This ELISA assay is sensitive to differences in pH among samples and/or standards. Hence, it is critical to ensure that all samples and standards are adjusted to the same pH prior to running the assay

Reagent Preparation

- 1. 5x Dilution Buffer: Add 25 mL of 5x Dilution Buffer to 100 mL of deionized water.
- 2. 5x Wash Buffer: Add 50 mL of 5x Wash Buffer to 200 mL of deionized water.
- 9(±)-HODE-HRP Conjugate: Add 300 μL of Conjugate to 11.70 mL Dilution Buffer.

STANDARD CURVE PREPARATION

The 9(±)-HODE Standard is provided as a 0.05 mg/mL stock solution in ethanol. Make a 1000 ng/mL working standard stock solution by adding 980 µL of Dilution Buffer directly to the vial containing 20 µL of 9(±)-HODE Standard. Use the table on the following page to construct an eight-point standard curve.

Standard	9(±)-HODE Conc. (ng/mL)	Vol. of Dilution Buffer (µL)	Transfer Vol. (µL)	Transfer Source	Final Vol. (µL)
S ₇	500	400	400	Stock	600
S ₆	100	800	200	S7	900
S ₅	10	900	100	S ₆	600
S ₄	5	400	400	S ₅	600
S ₃	1	800	200	S4	600
s_2	0.5	400	400	S ₃	600
s_1	0.1	800	200	S ₂	1000
В0	0	1000	백	딸	600

Assay Procedure

PROCEDURAL NOTES: 9(±)-HODE has been reported to bind to glassware. The use of plasticware (polypropylene) or silanated glassware is therefore recommended for all procedures involving the standards, enzyme conjugate, or samples containing 9-HODE.

To minimize errors in absorbance measurements due to handling, wipe the exterior bottom of the microplate wells with a lint-free paper towel prior to inserting into the plate reader.

- Add 100 µL of Standards or Samples to the corresponding wells on the microplate in duplicate. See Scheme I for a sample plate layout.
- 2. Add 100 μL of diluted 9(±)-HODE-HRP Conjugate to each well. Incubate at room temperature for two hours.
- 3. Wash the plate three times with 300 µL of diluted Wash Buffer per well.
- 4. Add 200 µL of TMB Substrate to each well. Incubate at room temperature for 45-60 minutes.
- Add 50 μ L of 3 N H₂SO₄ to each well to stop the reaction.
- 6. Read the plate at 450 nm.

NOTE: If accounting for substrate background, use 2 wells as blanks (BLK) with only 150 µL TMB Substrate in the wells. Subtract the average of these absorbance values from the absorbance values of the wells being

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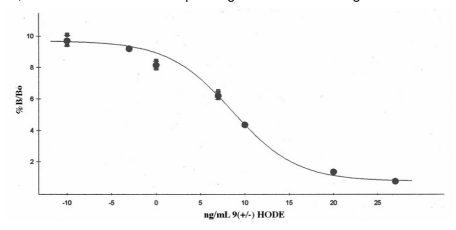
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assayed.

Scheme I: Sample Plate Layout											
1	2	3	4	5	6	7	8	9	10	11	12
В0	B ₀	U_1	U_1	U9	U9	U17	U17	U25	U25	U33	U33
s_1	s_1	U_2	U_2	U_{10}	U_{10}	U_{18}	U_{18}	U26	U26	U34	U34
s_2	s_2	U_3	U_3	U_{11}	U_{11}	U19	U19	U_{27}	U_{27}	U35	U35
S3	S3	U4	U4	U_{12}	U_{12}	U20	U20	U28	U28	U36	U36
S ₄	S ₄	U_5	U_5	U_{13}	U_{13}	U_{21}	U_{21}	U29	U29	U37	U37
S ₅	S ₅	U_6	U_6	U_{14}	U_{14}	U_{22}	U_{22}	U30	U ₃₀	U38	U38
S ₆	S ₆	U7	U7	U_{15}	U ₁₅	U_{23}	U23	U31	U31	U39	U39
S 7	S 7	U8	U ₈	U16	U16	U24	U24	U32	U32	BLK	BLK
	B ₀ S ₁ S ₂ S ₃ S ₄ S ₅ S ₆	1 2 B0 B0 S1 S1 S2 S2 S3 S3 S4 S4 S5 S5 S6 S6	1 2 3 B0 B0 U1 S1 S1 U2 S2 S2 U3 S3 S3 U4 S4 S4 U5 S5 S5 U6 S6 S6 U7	1 2 3 4 B0 B0 U1 U1 S1 S1 U2 U2 S2 S2 U3 U3 S3 S3 U4 U4 S4 S4 U5 U5 S5 S5 U6 U6 S6 S6 U7 U7	1 2 3 4 5 B0 B0 U1 U1 U9 S1 S1 U2 U2 U10 S2 S2 U3 U3 U11 S3 S3 U4 U4 U12 S4 S4 U5 U5 U13 S5 S5 U6 U6 U14 S6 S6 U7 U7 U15	1 2 3 4 5 6 B0 B0 U1 U1 U9 U9 S1 S1 U2 U2 U10 U10 S2 S2 U3 U3 U11 U11 S3 S3 U4 U4 U12 U12 S4 S4 U5 U5 U13 U13 S5 S5 U6 U6 U14 U14 S6 S6 U7 U7 U15 U15	1 2 3 4 5 6 7 B0 B0 U1 U1 U9 U9 U17 S1 S1 U2 U2 U10 U10 U18 S2 S2 U3 U3 U11 U11 U19 S3 S3 U4 U4 U12 U12 U20 S4 S4 U5 U5 U13 U13 U21 S5 S5 U6 U6 U14 U14 U22 S6 S6 U7 U7 U15 U15 U23	1 2 3 4 5 6 7 8 B0 B0 U1 U1 U9 U9 U17 U17 S1 S1 U2 U2 U10 U10 U18 U18 S2 S2 U3 U3 U11 U11 U19 U19 S3 S3 U4 U4 U12 U12 U20 U20 S4 S4 U5 U5 U13 U13 U21 U21 S5 S5 U6 U6 U14 U14 U22 U22 S6 S6 U7 U7 U15 U15 U23 U23	1 2 3 4 5 6 7 8 9 B0 B0 U1 U1 U9 U9 U17 U17 U25 S1 S1 U2 U2 U10 U10 U18 U18 U26 S2 S2 U3 U3 U11 U11 U19 U19 U27 S3 S3 U4 U4 U12 U12 U20 U20 U28 S4 S4 U5 U5 U13 U13 U21 U21 U29 S5 S5 U6 U6 U14 U14 U22 U22 U30 S6 S6 U7 U7 U15 U15 U23 U23 U31	1 2 3 4 5 6 7 8 9 10 B0 B0 U1 U1 U9 U9 U17 U17 U25 U25 S1 S1 U2 U2 U10 U10 U18 U18 U26 U26 S2 S2 U3 U3 U11 U11 U19 U19 U27 U27 S3 S3 U4 U4 U12 U12 U20 U20 U28 U28 S4 S4 U5 U5 U13 U13 U21 U21 U29 U29 S5 S5 U6 U6 U14 U14 U22 U22 U30 U30 S6 S6 U7 U7 U15 U15 U23 U23 U31 U31	1 2 3 4 5 6 7 8 9 10 11 B0 B0 U1 U1 U9 U9 U17 U17 U25 U25 U33 S1 S1 U2 U2 U10 U10 U18 U18 U26 U26 U34 S2 S2 U3 U3 U11 U11 U19 U19 U27 U27 U35 S3 S3 U4 U4 U12 U12 U20 U20 U28 U28 U36 S4 S4 U5 U5 U13 U13 U21 U21 U29 U29 U39 S5 S5 U6 U6 U14 U14 U22 U22 U30 U30 U38 S6 S6 U7 U7 U15 U15 U23 U23 U31 U31 U39

Calculation

- 1. Average the blank absorbance values and subtract the average from each well.
- Average standard replicates (S1 through S7) and divide by the average of the B0 values and multiply by 100 to obtain the %B0 value.
- Graph the %B0 on the y-axis (linear) vs. the standard concentration on the x-axis (logarithmic) to obtain a 3. standard curve (see Figure 1 for a typical standard curve).
- Average the replicates of each unknown and divide by the average B0 values and multiply by 100 to obtain 4. the %B/B0, then determine the corresponding concentration using the standard curve



Specificity

- ±)-HODE 100.0%
- S)-HODE 100.0% 9(
- R)-HODE 100.0%
- 13(S)-Hydroxyoctadecadienoic Acid 1.2%
- 13(R)-Hydroxyoctadecadienoic Acid 1.2%
- oxo-Octadecadienoic Acid 1.2%
- 13- oxo-Octadecadienoic Acid 2.4%
- 11(S)-HETE 0.0%

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15(S)-HETE 0.0% Linoleic Acid 0.0%

Precautions

- 1. Use aseptic technique when opening and dispensing reagents.
- This kit is designed to work properly as provided and instructed. Additions, deletions or substitutions to the 2. procedure or reagents are not recommended, as they may be detrimental to the assay.

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