

Human C-Peptide

96-Well Plate

Cat. # EZHCP-20K, EZHCP-20BK

HUMAN C-PEPTIDE ELISA KIT 96-Well Plate (Cat. #EZHCP-20K, EZHCP-20BK)

I.	Intended Use	2
II.	Principles of Procedure	2
III.	Reagents Supplied	2
IV.	Storage and Stability	4
٧.	Reagent Precautions	4
VI.	Materials Required But Not Provided	6
VII.	Sample Collection And Storage	6
VIII.	Assay Procedure	7
IX.	Microtiter Plate Arrangement	9
Χ.	Calculations	11
XI.	Assay Characteristics	11
XII.	Correlation Graph of HCP ELISA vs RIA	14
XIII.	Human C-Peptide Standard Curve	14
	Quality Controls	15
XV.	Troubleshooting Guide	15
XVI.	Replacement Reagents	15
XVII.	Ordering Information	16
XVIII.	References	16

HUMAN C-PEPTIDE ELISA KIT 96-Well Plate (Cat. #EZHCP-20K, EZHCP-20BK)

I. INTENDED USE

This kit is for non-radioactive quantification of Human C-Peptide (HCP) in serum, plasma and other biological media. One kit is sufficient to measure 38 unknown samples in duplicate. *This kit is for Research Use Only. Not for Use in Diagnostic Procedures.*

II. PRINCIPLES OF PROCEDURE

This assay is based, sequentially, on: 1) capture of Human C-Peptide from samples by a monoclonal antibody immobilized to the wells of a microtiter plate, 2) binding of the biotinylated monoclonal HCP antibody to capture Human C-Peptide molecules, 3) wash away of unbound materials including free materials from samples and free detection antibody, 4) conjugation of SA-HRP (Poly-HRP-labeled streptavidin) enzyme to the biotinylated antibodies, and 5) quantification of bound detection conjugate by monitoring SA-HRP enzyme activity in the presence of TMB (tetramethylbenzidine) substrates. The enzyme activity is measured spectrophotometrically by the absorbency at 450 nm due to production of the photometric product. Since the amount of photometric product is directly proportional to the concentration of Human C-Peptide in the unknown sample, the latter can be derived by interpolation from a reference curve generated in the same assay with reference standards of known concentrations of HCP.

III. REAGENTS SUPPLIED

Each kit is sufficient to run one 96-well microtiter plate and contains the following reagents:

A. Human C-Peptide ELISA Plate

Coated with anti-Human C-Peptide Monoclonal Antibody

Quantity: 1 plate

Preparation: Ready to use

Note: Unused strips should be resealed in the foil pouch with the desiccant

provided and stored at 2-8°C.

B. Adhesive Plate Sealer

Quantity: 1 Sheet

Preparation: Ready to use

C. 10X HRP Wash Buffer Concentrate

10X concentrate of 50 mM TBS Buffer containing 0.05% Tween 20

Quantity: 2 bottles containing 50 mL each Preparation: Dilute 1:10 with deionized water

III. REAGENTS SUPPLIED (continued)

D. Human C-Peptide Standards

Human C-Peptide in Assay Buffer: 0.2, 0.5, 1, 2, 5, 10, and 20 ng/mL

Quantity: 0.5 mL/vial Preparation: Ready to use

Note: The standard(s) in this kit have been calibrated to an International Reference standard, NIBSC code # 84/510, Version 03.

E. ELISA Human C-Peptide Quality Controls 1 and 2

Human C-Peptide in QC Buffer

Quantity: 0.5 mL/vial Preparation: Ready to use

F. Matrix Solution

Quantity: 1 mL

Preparation: Ready to use

G. Assay Buffer

0.05M PBS containing 0.025 M EDTA, 1% BSA, with 0.08%

Sodium Azide, with proprietary reagent, pH 7.4

Quantity: 8 mL

Preparation: Ready to use

H. Human C-Peptide Detection Antibody

Biotinylated anti-Human C-Peptide Monoclonal Antibody

Quantity: 3.0 mL

Preparation: Ready to use

I. Enzyme Solution

Pre-titered Streptavidin-Horseradish Peroxidase Conjugate (SA-HRP)

Quantity: 12 mL

Preparation: Ready to use

J. Substrate (TMB)

3, 3', 5, 5'-tetramethylbenzidine (TMB)

Quantity: 12 mL

Preparation: Ready to use

K. ELISA Stop Solution

0.3M HCI

Quantity: 12 mL

Preparation: Ready to use

IV. STORAGE AND STABILITY

Recommended storage for kit components is 2-8°C.

All components are shipped and stored at 2-8°C. Once opened, liquid standards and controls can be stored up to 30 days at 2-8°C. Refer to expiration dates on all reagents prior to use. Do not mix reagents from different kits unless they have the same lot numbers

V. REAGENT PRECAUTIONS

A. Sodium Azide or Proclin

Sodium Azide or Proclin has been added to some reagents as a preservative. Although the concentrations are low, Sodium Azide and Proclin may react with lead and copper plumbing to form highly explosive metal azides. Dispose of unused contents and waste in accordance with international, federal, state, and local regulations

B. Hydrochloric Acid

Hydrochloric Acid is corrosive and can cause eye and skin burns. It is harmful if swallowed and can cause respiratory and digestive tract burns. Avoid contact with skin and eyes.

Note: See Full Labels of Hazardous components on next page.

Full labels of hazardous components in this kit:

Ingredient, Cat #		Full Label	
Human C-Peptide Detection Antibody	E1020	<u>(i)</u>	Warning. Causes serious eye irritation. IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.
ELISA Quality Control 1 & 2	E6020-K	<u>(i)</u>	Warning. Causes serious eye irritation. IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.
Human C-Peptide Standards	E8020-K		Warning. Causes serious eye irritation. IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.
Stop Solution	ET-TMB		Warning. May be corrosive to metals.
10X HRP 10X Wash Buffer Concentrate	EWB-HRP	<u>(i)</u>	Warning. May cause an allergic skin reaction. Wear protective gloves. IF ON SKIN: Wash with plenty of soap and water.

VI. MATERIALS REQUIRED BUT NOT PROVIDED

- 1. Pipets with tips, 10 μL-200 μL
- 2. Multi-channel pipette, 50 μL-300 μL
- 3. Buffer and Reagent Reservoirs
- 4. Vortex Mixer
- 5. Refrigerator
- 6. Deionized Water
- 7. Microtiter plate reader capable of reading absorbency at 450 nm
- 8. Microtiter Plate Shaker
- 9. Absorbent Paper or Cloth

VII. SAMPLE COLLECTION AND STORAGE

- 1. Human C-Peptide must be protected from proteolysis during assay procedures and sample storage. Trasylol (Aprotinin) at a concentration of 500 KIU per mL of serum or plasma should be added to samples to protect from proteolysis.
- 2. To prepare serum samples, whole blood is directly drawn into a Vacutainer® serum tube that contains no anticoagulant. Let blood clot at room temperature for 30 minutes. Promptly centrifuge the clotted blood at 2,000 to 3,000 x g for 15 minutes at $4 \pm 2^{\circ}$ C. Transfer and store serum samples in separate tubes.
- 3. Samples can be stored at 4°C if they will be tested within 3 hours of collection. For longer storage, specimens should be stored at ≤ -20°C. Avoid multiple (>3) freeze/thaw cycles. Aliquot samples before freezing if necessary.
- 4. To prepare plasma samples, whole blood should be collected into Vacutainer[®] EDTA-plasma tubes and centrifuged immediately after collection. Observe the same precautions in the preparation of serum samples.
- 5. If heparin is to be used as an anticoagulant, the effect on the assay outcome at the dose of heparin used should be pre-determined.
- 6. Avoid using samples with gross hemolysis or lipemia.

VIII. ASSAY PROCEDURE

Pre-warm all reagents to room temperature (20-25°C) immediately before setting up the assay.

- 1. Dilute the concentrated 10X Wash Buffers 10 fold. Mix the entire contents of both bottles of 10X Wash Buffer with 900 mL distilled or deionized water.
- 2. Remove the required number of strips from the Microtiter Assay Plate. Assemble the strips in an empty plate holder and fill each well with 300 μL of 1X Wash Buffer. Incubate at room temperature for 5 minutes. Decant Wash Buffer and remove the residual amount from all wells by inverting the plate and tapping it smartly onto absorbent towels several times. Do not let wells dry before proceeding to the next step.
- 3. Wash the wells two additional times with 1X Wash Buffer, 300 µL per well per wash. Decant and tap after each wash to remove residual buffer.
- 4. Add 40 μL of Assay Buffer into each Blank, Standard, and QC well. Refer to Section IX for suggested well orientations.
- 5. Add 50 µL of the Assay Buffer into each Sample well.
- 6. Add 10 µL Assay Buffer to assay background well, i.e. the blank wells #A1 and #B1.
- 7. Add 10 µL of Matrix Solution into each Blank, Standard, and QC well.
- 8. Add 10 µL of Standards in duplicate into appropriate wells.
- 9. Add 10 µL of QC1 and QC2 in duplicate into appropriate wells.
- 10. Add 10 µL of serum or plasma samples in duplicate into appropriate wells.
- 11. Add 20 µL of the Detection Antibody into each well. For best results all additions should be completed within one hour.
- 12. Cover the plate with plate sealer.

VIII. ASSAY PROCEDURE (continued)

- 13. Incubate at room temperature (20-25°C) for 2 hours while shaking on a microtiter plate shaker.
- 14. Remove plate sealer and decant solutions from the plate. Tap as before to remove residual solutions in the well.
- 15. Wash the wells 5 times with 1X Wash Buffer, 300 µL per well per wash. Decant and tap after each wash to remove residual buffer.
- 16. Add 80 µL of the Enzyme Solution into each well.
- 17. Cover the plate with plate sealer. Incubate 30 minutes at room temperature *while* shaking on a microtiter plate shaker.
- 18. Wash the wells 5 times with 1X Wash Buffer, 300 µL per well per wash. Decant and tap after each wash to remove residual buffer.
- 19. Add 80 µL of the Substrate Solution to each well, cover plate with sealer and shake on the plate shaker for **approximately** 12 to 19 minutes. Blue color should be formed in wells of Human C-Peptide standards with intensity proportional to increasing concentrations of Human C-Peptide.

NOTE: Please be aware that the color may develop more quickly or more slowly than the recommended incubation time depending on the localized room temperature. Please visually monitor the color development to optimize the incubation time.

- 20. Remove the plate sealer, and stop the reaction by adding 80 µL of Stop Solution into each well of the plate. Shake the plate by hand to ensure complete mixing of solution in all wells. The blue color should turn to yellow after acidification.
- 21. Wipe the bottom of the microtiter plate to remove any residue prior to reading on plate reader. Read absorbance at 450 nm in a plate reader within 5 minutes and ensure that there is no air bubbles in any well. The absorbance of highest Human C-Peptide standard should be approximately 2.4 ~ 2.8

Assay Procedure for Human C-Peptide ELISA Kit (Cat. # EZHCP-20K)

	Step 1	Step 2	Step 3	Step 4	Step 5-7	Step 8	Step 8-10	Step 11	Step 11-13	Step 14	Step 14	Step 14	Step 14
Well #		minutes.	Assay Buffer	Matrix Solution	Standards/ Controls/ Samples	Detection Antibody		Enzyme Solution		Substrate		Stop Solutio n	
A1, B1	ater.	5 minu Is.	50 μL	10 μL		20 μL		80 µL		80 µL		80 µL	
C1, D1	Dilute both bottles of 10X Wash Buffer with 900mL Deionized Water.	µL Wash Buffer to plate and incubate at room temperature for 5 i Remove residual buffer by tapping smartly on absorbent towels. Wash 2X with 300 µL Wash Buffer	40 µL	10 μL	10 μL of 0.2 ng/mL Standard	20 μL	ıture.	80 μL	rature .	80 µL	12-19 minutes at Room Temperature.	80 µL	
E1, F1	JL Deiol	empera absorbe fer	40 μL	10 μL	10 μL of 0.5 ng/mL Standard	20 µL	at Room Temperature. Wash Buffer	80 µL	Seal, Agitate, Incubate 30 minutes at Room Temperature Wash 5X with 300 µL Wash Buffer	80 µL	m Temp	80 µL	
G1, H1	th 900m	t room t irtly on ash Buff	40 μL	10 μL	10 μL of 1 ng/mL Standard	20 µL	at Room Ten Wash Buffer	80 µL	t Room ash Buf	80 µL	at Roo	80 µL	Ë
A2, B2	uffer wi	ubate a ing sma 0 µL Wa	40 μL	10 μL	10 μL of 2 ng/mL Standard	20 µL	ours at 0 µL Wa	80 µL	nutes a 0 µL W≀	80 µL	minutes	80 µL	at 450 n
C2, D2	Wash B	and inc by tappi with 30	40 μL	10 μL	10 μL of 5 ng/mL Standard	20 µL	Incubate 2 hours sh 5X with 300 µ∟	80 µL	te 30 mi with 30	80 µL	, 12-19 r	80 µL	rbance
E2, F2	of 10X	o plate buffer I /ash 2X	40 μL	10 μL	10 μL of 10 ng/mL Standard	20 µL	ate, Incubate 2 hours Wash 5X with 300 µL	80 µL	Incubat Iash 5X	80 µL	Seal, Agitate, Incubate	80 µL	Read Absorbance at 450 nm.
G2, H2	bottles	Buffer t esidual M	40 μL	10 μL	10 μL of 20 ng/mL Standard	20 µL	Seal, Agitate, Wa	80 µL	Agitate, M	80 µL	gitate, l	80 µL	Rea
A3, B3	te both	- Wash emove r	40 μL`	10 μL	10 μL of QC I	20 µL	Seal	80 µL	Seal, /	80 µL	Seal, A	80 µL	
C3, D3	Dilu	Add 300 µl	40 μL	10 μL	10 μL of QC II	20 µL		80 µL		80 µL		80 µL	
E3, F3 ↓		Adı	50 μL		10 μL of Sample	20 μL		80 µL		80 µL		80 µL	

IX. MICROTITER PLATE ARRANGEMENT

Standard Human C-Peptide ELISA

	1	2	3	4	5	6	7	8	9	10	11	12
А	Blank	2 ng/mL	EQC 1	etc								
В	Blank	2 ng/mL	EQC 1	etc								
С	0.2 ng/mL	5 ng/mL	EQC 2									
D	0.2 ng/mL	5 ng/mL	EQC 2									
E	0.5 ng/mL	10 ng/mL	Sample 1									
F	0.5 ng/mL	10 ng/mL	Sample 1									
G	1 ng/mL	20 ng/mL	Sample 2									
Н	1 ng/mL	20 ng/mL	Sample 2									

X. CALCULATIONS

The dose-response curve of this assay fits best to a sigmoidal 5-parameter logistic equation. The results of unknown samples can be calculated with any computer program having a 5-parameter logistic function.

XI. ASSAY CHARACTERISTICS

Sensitivity

The lowest level of Human C-Peptide that can be detected by this assay is 0.05 ng/mL.

Crossreactivity

The specificity of the Human C-Peptide ELISA is the ability to selectively measure the analytes in the presence of other like components in the sample matrix.

Human C-Peptide	100%
Intact Human	63%
Proinsulin	
Human Insulin	n.d.*
Porcine C-Peptide	n.d.*
Rat C-Peptide	n.d.*
Canine C-Peptide	n.d.*

n.d.: Not detectable at concentrations up to 200 ng/mL.

Precision

Within and Between Assay Variation

Sample No.	Mean HCP Levels ng/mL	Within % CV	Between % CV
1	1	4.75	8.72
2	3	2.95	5.00
3	7	1.60	

XI. ASSAY CHARACTERISTICS (continued)

The between assay variation of EMD Millipore Human C-Peptide ELISA kits were studied using two serum samples with varying concentrations of Human C-Peptide. The mean variation of each sample was calculated using results from eight separate assays with duplicate samples in each assay.

Recovery

Spike & Recovery of Human C-Peptide in Human Serum

Sample #	HCP	Expected	Observed	% of
	added	ng/mL	ng/mL	Recovery
	ng/mL			
1	0	2.31	2.31	100
	0.5	2.81	2.83	104
	2.0	4.31	4.21	95
	5.0	7.31	7.70	108
2	0	1.68	1.68	100
	0.5	2.18	2.19	101
	2.0	3.68	3.57	93
	5.0	6.68	6.77	102
3	0	2.05	2.05	100
	0.5	2.55	2.50	98
	2.0	4.05	3.92	91
	5.0	7.05	6.93	97
4	0	3.84	3.84	100
	0.5	4.34	4.34	100
	2.0	5.84	5.87	102
	5.0	8.84	9.34	111

Varying concentrations of Human C-Peptide were added to four human serum samples and the Human C-Peptide content was determined by ELISA. Mean of the observed levels from four duplicate determinations are shown.

Percent recovery = observed \div expected x 100%.

XI. ASSAY CHARACTERISTICS (continued)

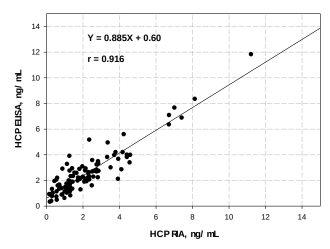
Linearity

Effect of Serum Dilution

Sample	Sample	Expected	Observed	% of
No.	Dilution	ng/mL	ng/mL	Expected
1	0	3.38	3.38	100
	2	1.69	1.83	108
	4	0.85	0.88	104
	8	0.42	0.43	102
2	0	7.24	7.24	100
	2	3.62	3.17	88
	4	1.81	1.65	92
	8	0.91	0.88	98
3	0	8.42	8.42	100
	2	4.21	3.60	86
	4	2.11	2.10	100
	8	1.05	1.01	96

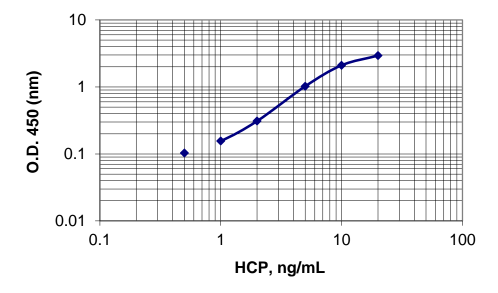
Dilutions of human sera containing varying concentrations of Human C-Peptide were analyzed. The mean Human C-Peptide level and percent of expected from four duplicates determinations are shown.

XII. CORRELATION GRAPH OF HCP ELISA vs RIA



Note: One hundard human serum samples were analized using this HCP \blacksquare JSA kit and Millipore's HCP \blacksquare A kit (Cat #: HCP20K).

XIII. HUMAN C-PEPTIDE STANDARD CURVE



XIV. QUALITY CONTROLS

The ranges for each Quality Control 1 and 2 are provided on the card insert or can be located at the EMD Millipore website emdmillipore.com using the catalog number as the keyword.

XV. TROUBLESHOOTING GUIDE

Low or No Signal with Standards

- * Standards were left at room temperature. Standards should be stored at -20°C.
- * Insufficient time for reaction with substrate. Allow substrate to react longer.
- * Kit reagents have expired.
- * Inadequate plate washing after sample incubation.
- * Too much washing after conjugate incubation can reduce signal.

High Background

- * Inadequate plate washing. After conjugate incubation, tap out plate on absorbent towels after decanting.
- * Cross contamination between neighboring wells.

Samples too High

* Dilute sample with assay buffer to bring HCP concentration within standard range.

Signal too High on Highest Standard

* Plate incubated too long with substrate. Discard substrate, wash plate once and add freshly prepared substrate. Check RFU in less time.

High Variance in RFU of Duplicates

- * Cross contamination in wells
- * Bubbles in substrate at time of reading
- * Loss of reagent or faulty pipetting in duplicates

XVI. REPLACEMENT REAGENTS

Reagents	Catalogue #
Human C-Peptide ELISA Plate with Plate Sealers	EP20
10X HRP Wash Buffer Concentrate	EWB-HRP
Human C-Peptide Standards (0.5 mL/vial)	E8020-K
ELISA HCP Quality Controls 1 & 2 (0.5 mL/vial)	E6020-K
Assay Buffer (8 mL/vial)	EABIR-2
Human C-Peptide Detection Antibody (3 mL/vial)	E1020
Enzyme Solution (12 mL/vial)	EHRP
Substrate (12 mL/vial)	ESS-TMB
Matrix Solution (1 mL/vial)	EMTX-CP
Stop Solution (12 mL/vial)	ET-TMB
10-pack of Human C-Peptide (HCP) ELISA kits	EZHCP-20BK

XVII.ORDERING INFORMATION

To place an order or to obtain additional information about our immunoassay products, please contact your Customer Service or Technical Support Specialist.

Contact information for each region can be found on our website:

emdmillipore.com/contact

Conditions of Sale

For Research Use Only. Not for Use in Diagnostic Procedures.

Safety Data Sheets (SDS)

Safety Data Sheets for EMD Millipore products may be ordered by fax or phone or through our website at emdmillipore.com/msds.

XVIII. REFERENCES

- Tijsen P. "Practice and Theory of Enzyme Immunoassays" in Burdon RH and Knippenberg PH (Ed.), <u>Laboratory Techniques in Biochemisrty and Molecular</u> <u>Biology</u>. Amsterdam/NY: Elsevier, 1985
- 2. O'Rahilly S & Moller DE 1992 Clin Endocrinol 36:121.
- 3. Reaven GM 1991 Diab Care 14:195.
- 4. Fontbonne AM & Eschwege EM 1991 Diab Care 14:461.
- 5. Horwitz DL et al., 1976 New Engl J Med 295:207.
- 6. Bommen M et al., 1984 Arch Dis Child 59:1096...
- 7. Field JB 1989 Endocrinol Metab Clin North Am 18:27.