invitrogen USER GUIDE

Rat IL-17A ELISA Kit

Enzyme-linked Immunosorbent Assay for quantitative detection of rat IL-17A

Catalog Numbers BMS635 or BMS635TEN

Pub. No. MAN0016907 Rev. A.0 (30)



WARNING! Read the Safety Data Sheets (SDSs) and follow the handling instructions. Wear appropriate protective eyewear, clothing, and gloves. Safety Data Sheets (SDSs) are available from **thermofisher.com/support**.

Product description

The Rat IL-17A ELISA Kit is an enzyme-linked immunosorbent assay for the quantitative detection of rat IL-17A.

Summary

A new family of cytokines, Interleukin-17, has recently been defined that reveals a distinct ligand-receptor signaling system. There is high evidence for its importance in the regulation of immune responses. IL-17A was first characterized and six IL-17 family members (IL-17A-F) have subsequently been described. IL-17A, a homodimeric cytokine of about 32 kDa, is largely produced by activated memory T-lymphocytes, but stimulates innate immunity and host defense. IL-17A and IL-17F both mobilize neutrophils partly through granulopoeisis and CXC chemokine induction, as well as increased survival locally. IL-17A and IL-17F production by T-lymphocytes is regulated by IL-23 independent of T-cell receptor activation.

The T help 1 (Th1) and Th2 cell classification has until recently provided the framework for understanding CD4(+) T-cell biology and the interplay between innate and adaptive immunity. Recent studies have defined a previously unknown arm of the CD4(+) T-cell effector response, the Th17 lineage. This subset of T-cells produces interleukin 17, which is highly proinflammatory and induces severe autoimmunity. Whereas IL-23 serves to expand previously differentiated T(H)-17 cell populations, IL-6 and transforming growth factor- β (TGF- β) induce the differentiation of T(H)-17 cells from naive precursors.

Increasing evidence shows that IL-17 family members play an active role in inflammatory diseases, autoimmune diseases, and cancer. The IL-17 signaling system is operative in disparate tissues such as articular cartilage, bone, meniscus, brain, hematopoietic tissue, kidney, lung, skin and intestine. Thus, the evolving IL-17 family of ligands and receptors may play an important role in the homeostasis of tissues in health and disease beyond the immune system. Increased levels of IL-17 have been associated with several conditions, including airway inflammation, rheumatoid arthritis, intraperitoneal abscesses and adhesions, inflammatory bowel disease, allograft rejection, psoriasis, cancer and multiple sclerosis.

For literature update refer to our website.

Principles of the test

An anti-rat IL-17A coating antibody is adsorbed onto microwells.

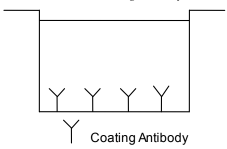


Fig. 1 Coated microwell

Rat IL-17A present in the sample or standard binds to antibodies adsorbed to the microwells. A biotin-conjugated anti-rat IL-17A antibody is added and binds to rat IL-17A captured by the first antibody.

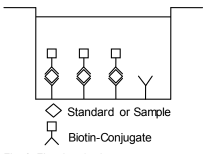


Fig. 2 First incubation

Following incubation unbound biotin-conjugated anti-rat IL-17A antibody is removed during a wash step. Streptavidin-HRP is added and binds to the biotin-conjugated anti-rat IL-17A antibody.

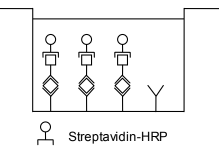


Fig. 3 Second incubation

Following incubation unbound Streptavidin-HRP is removed during a wash step, and substrate solution reactive with HRP is added to the wells.

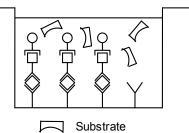


Fig. 4 Third incubation

A colored product is formed in proportion to the amount of rat IL-17A present in the sample or standard. The reaction is terminated by addition of acid and absorbance is measured at 450 nm. A standard curve is prepared 7 rat IL-17A standard dilutions and rat IL-17A sample concentration determined.

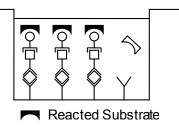


Fig. 5 Stop reaction

Reagents provided

Reagents for rat IL-17A ELISA BMS635 (96 tests)

1 aluminum pouch with a Microwell Plate (12 strips of 8 wells each) coated with monoclonal antibody to rat IL-17A

1 vial (70 $\mu L)$ Biotin-Conjugate anti-rat IL-17A monoclonal antibody 1 vial (150 $\mu L)$ Streptavidin-HRP

2 vials rat IL-17A Standard lyophilized, 200 pg/mL upon reconstitution

1 bottle (12 mL) Sample Diluent

1 vial (5 mL) Assay Buffer Concentrate 20x (PBS with 1% Tween $^{\text{\tiny M}}$ 20, 10% BSA)

1 bottle (50 mL) Wash Buffer Concentrate 20x (PBS with 1% Tween [™] 20)

1 vial (15 mL) Substrate Solution (tetramethyl-benzidine)

1 vial (15 mL) Stop Solution (1M Phosphoric acid)

4 adhesive Films

Reagents for rat IL-17A ELISA BMS635TEN (10x96 tests)

10 aluminum pouches with a Microwell Plate (12 strips of 8 wells each) coated with monoclonal antibody to rat IL-17A

10 vials (70 $\mu L)$ Biotin-Conjugate anti-rat IL-17A monoclonal antibody 10 vials (150 $\mu L)$ Streptavidin-HRP

 $10\ vials\ rat\ IL-17A\ Standard\ lyophilized, 200\ pg/mL\ upon\ reconstitution$

7 bottles (12 mL) Sample Diluent

2 vials (5 mL) Assay Buffer Concentrate 20x (PBS with 1% Tween $^{^{\text{\tiny TM}}}$ 20, 10% BSA)

6 bottles (50 mL) Wash Buffer Concentrate 20x (PBS with 1% Tween 20)

10 vials (15 mL) Substrate Solution (tetramethyl-benzidine) 1 vial (100 mL) Stop Solution (1M Phosphoric acid) 40 adhesive Films

Storage instructions – ELISA kit

Store kit reagents between 2°C and 8°C.

Immediately after use remaining reagents should be returned to cold storage (2°C to 8°C). Expiry of the kit and reagents is stated on labels.

Expiry of the kit components can only be guaranteed if the components are stored properly, and if, in case of repeated use of one component, this reagent is not contaminated by the first handling.

Sample collection and storage instructions

Cell culture supernatant, serum and plasma (EDTA, citrate, heparin) were tested with this assay. Other biological samples might be suitable for use in the assay.

Remove serum or plasma from the clot or cells as soon as possible after clotting and separation.

Pay attention to a possible *Hook Effect* due to high sample concentrations (see "Calculation of results" on page 4).

Samples containing a visible precipitate must be clarified prior to use in the assay. Do not use grossly hemolyzed or lipemic specimens.

Samples should be aliquoted and must be stored frozen at -20°C to avoid loss of bioactive rat IL-17A. If samples are to be run within 24 hours, they may be stored at 2°C to 8°C (refer to "Sample stability" on page 6).

Avoid repeated freeze-thaw cycles. Prior to assay, the frozen sample should be brought to room temperature slowly and mixed gently.

Materials required but not provided

- 5 mL and 10 mL graduated pipettes
- 5 µL to 1000 µL adjustable single channel micropipettes with disposable tips
- 50 μL to 300 μL adjustable multichannel micropipette with disposable tips
- Multichannel micropipette reservoir
- Beakers, flasks, cylinders necessary for preparation of reagents
- Device for delivery of wash solution (multichannel wash bottle or automatic wash system)
- Microwell strip reader capable of reading at 450 nm (620 nm as optional reference wave length)
- · Glass-distilled or deionized water
- Statistical calculator with program to perform regression analysis

Precautions for use

- All reagents should be considered as potentially hazardous. We
 therefore recommend that this product is handled only by those
 persons who have been trained in laboratory techniques and that it
 is used in accordance with the principles of good laboratory
 practice. Wear suitable protective clothing such as laboratory
 overalls, safety glasses, and gloves. Care should be taken to avoid
 contact with skin or eyes. In the case of contact with skin or eyes
 wash immediately with water. See material safety data sheet(s)
 and/or safety statement(s) for specific advice.
- Reagents are intended for research use only and are not for use in diagnostic or therapeutic procedures.
- Do not mix or substitute reagents with those from other lots or other sources.
- Do not use kit reagents beyond expiration date on label.
- Do not expose kit reagents to strong light during storage or incubation.
- Do not pipet by mouth.
- Do not eat or smoke in areas where kit reagents or samples are handled.
- Avoid contact of skin or mucous membranes with kit reagents or samples.
- Rubber or disposable latex gloves should be worn while handling kit reagents or samples.
- Avoid contact of substrate solution with oxidizing agents and metal.

- · Avoid splashing or generation of aerosols.
- To avoid microbial contamination or cross-contamination of reagents or samples that may invalidate the test, use disposable pipette tips and/or pipettes.
- Use clean, dedicated reagent trays for dispensing the conjugate and substrate reagent.
- Exposure to acid inactivates the conjugate.
- Glass-distilled water or deionized water must be used for reagent preparation.
- Substrate solution must be at room temperature prior to use.
- Decontaminate and dispose samples and all potentially contaminated materials as if they could contain infectious agents.
 The preferred method of decontamination is autoclaving for a minimum of 1 hour at 121.5°C.
- Liquid wastes not containing acid and neutralized waste may be mixed with sodium hypochlorite in volumes such that the final mixture contains 1.0% sodium hypochlorite. Allow 30 minutes for effective decontamination. Liquid waste containing acid must be neutralized prior to the addition of sodium hypochlorite.

Preparation of reagents

- 1. Buffer Concentrates should be brought to room temperature and should be diluted before starting the test procedure.
- 2. If crystals have formed in the Buffer Concentrates, warm them gently until they have completely dissolved.

Wash buffer (1x)

- Pour entire contents (50 mL) of the Wash Buffer Concentrate (20x) into a clean 1000 mL graduated cylinder. Bring to final volume of 1000 mL with glass-distilled or deionized water. Mix gently to avoid foaming.
- 2. Transfer to a clean wash bottle and store at 2°C to 25°C. Please note that Wash Buffer (1x) is stable for 30 days.
- 3. Wash Buffer (1x) may also be prepared as needed according to the following table:

Number of Strips	Wash Buffer Concentrate (20x) (mL)	Distilled Water (mL)
1 - 6	25	475
1 - 12	50	950

Assay buffer (1x)

- 1. Pour the entire contents (5 mL) of the Assay Buffer Concentrate (20x) into a clean 100 mL graduated cylinder. Bring to final volume of 100 mL with distilled water. Mix gently to avoid foaming.
- 2. Store at 2°C to 8°C. Please note that the Assay Buffer (1x) is stable for 30 days.
- 3. Assay Buffer (1x) may also be prepared as needed according to the following table:

Number of Strips	Assay Buffer Concentrate (20x) (mL)	Distilled Water (mL)
1 - 6	2.5	47.5
1 - 12	5.0	95.0

Biotin-Conjugate

Note: The Biotin-Conjugate should be used within 30 minutes after dilution

Make a 1:100 dilution of the concentrated Biotin-Conjugate solution with Assay Buffer (1x) in a clean plastic tube as needed according to the following table:

Number of Strips	Biotin-Conjugate (mL)	Assay Buffer (1x) (mL)
1 - 6	0.03	2.97
1 - 12	0.06	5.94

Streptavidin-HRP

Note: The Streptavidin-HRP should be used within 30 minutes after dilution

Make a 1:200 dilution of the concentrated Streptavidin-HRP solution with Assay Buffer (1x) in a clean plastic tube as needed according to the following table:

Number of Strips	Streptavidin-HRP (mL)	Assay Buffer (1x) (mL)
1 - 6	0.03	5.97
1 - 12	0.06	11.94

Rat IL-17A standard

- 1. Reconstitute rat IL-17A standard by addition of distilled water.
- 2. Reconstitution volume is stated on the label of the standard vial. Swirl or mix gently to insure complete and homogeneous solubilization (concentration of reconstituted standard = 200 pg/mL).
- 3. Allow the standard to reconstitute for 10-30 minutes. Mix well prior to making dilutions.

The standard has to be used immediately after reconstitution and cannot be stored.

External standard dilution

- Label 7 tubes, one for each standard point: S1, S2, S3, S4, S5, S6, S7.
- 2. Then prepare 1:2 serial dilutions for the standard curve as follows: Pipette 225 μL of Sample Diluent into each tube.
- 3. Pipette 225 μL of reconstituted (concentration of standard = 200.0 pg/mL) into the first tube, labeled S1, and mix (concentration of standard 1 = 100.0 pg/mL).
- Pipette 225 µL of this dilution into the second tube, labeled S2, and mix thoroughly before the next transfer.
- 5. Repeat serial dilutions 5 more times thus creating the points of the standard curve (see Figure 6).

Sample Diluent serves as blank.

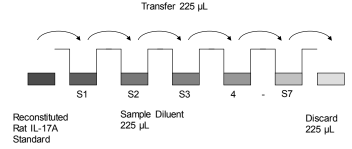


Fig. 6 Dilute standards - tubes

Test protocol

- Determine the number of microwell strips required to test the desired number of samples plus appropriate number of wells needed for running blanks and standards. Each sample, standard, blank and optional control sample should be assayed in duplicate. Remove extra microwell strips from holder and store in foil bag with the desiccant provided at 2°C to 8°C sealed tightly.
- 2. Wash the microwell strips twice with approximately $400~\mu$ L Wash Buffer per well with thorough aspiration of microwell contents between washes. Allow the Wash Buffer to sit in the wells for about 10-15 seconds before aspiration. Take care not to scratch the surface of the microwells.

After the last wash step, empty wells and tap microwell strips on absorbent pad or paper towel to remove excess Wash Buffer. Use the microwell strips immediately after washing. Alternatively microwell strips can be placed upside down on a wet absorbent paper for not longer than 15 minutes. Do not allow wells to dry.

3. Standard dilution on the microwell plate (Alternatively the standard dilution can be prepared in tubes - see "External standard dilution" on page 3.): Add 100 μL of Sample Diluent in duplicate to all standard wells. Pipette 100 μL of prepared standard (see "Rat IL-17A standard" on page 3 concentration = 200 pg/mL) in duplicate into well A1 and A2 (see Table 1). Mix the contents of wells A1 and A2 by repeated aspiration and ejection (concentration of standard 1, S1 = 100.0 pg/mL), and transfer 100 μL to wells B1 and B2, respectively (see Figure 7). Take care not to scratch the inner surface of the microwells. Continue this procedure 5 times, creating two rows of rat IL-17A standard dilutions ranging from 100.0 to 1.6 pg/mL. Discard 100 μL of the contents from the last microwells (G1, G2) used.

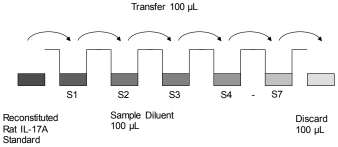


Fig. 7 Dilute standards - microwell plate

Table 1 Example of the arrangement of blanks, standards and samples in the microwell strips.

	•				
	1	2	3	4	
А	Standard 1 100.0 pg/mL	Standard 1 100.0 pg/mL	Sample 1	Sample 1	
В	Standard 2 50.0 pg/mL	Standard 2 50.0 pg/mL	Sample 2	Sample 2	
С	Standard 3 25.0 pg/mL	Standard 3 25.0 pg/mL	Sample 3	Sample 3	
D	Standard 4 12.5 pg/mL	Standard 4 12.5 pg/mL	Sample 4	Sample 4	
Е	Standard 5 6.3 pg/mL	Standard 5 6.3 pg/mL	Sample 5	Sample 5	
F	Standard 6 3.1 pg/mL	Standard 6 3.1 pg/mL	Sample 6	Sample 6	
G	Standard 7 1.6 pg/mL	Standard 7 1.6 pg/mL	Sample 7	Sample 7	
Н	Blank	Blank	Sample 8	Sample 8	

In case of an external standard dilution (see "External standard dilution" on page 3), pipette 100 μL of these standard dilutions (S1 - S7) in the standard wells according to Table 1.

- 4. Add 100 μL of Sample Diluent in duplicate to the blank wells.
- 5. Add $50 \mu L$ of Sample Diluent to the sample wells.
- 6. Add 50 µL of each sample in duplicate to the sample wells.
- 7. Prepare Biotin-Conjugate (see "Biotin-Conjugate" on page 3).
- 8. Add 50 μL of Biotin-Conjugate to all wells.
- 9. Cover with an adhesive film and incubate at room temperature (18°C to 25°C) for 2 hours on a microplate shaker.
- Prepare Streptavidin-HRP (refer to "Streptavidin-HRP" on page 3).
- 11. Remove adhesive film and empty wells. Wash microwell strips 4 times according to point 2 of the test protocol. Proceed immediately to the next step.
- 12. Add 100 μL of diluted Streptavidin-HRP to all wells, including the blank wells.
- Cover with an adhesive film and incubate at room temperature (18°C to 25°C) for 1 hours on a microplate shaker.
- 14. Remove adhesive film and empty wells. Wash microwell strips 4 times according to point 2 of the test protocol. Proceed immediately to the next step.
- 15. Pipette 100 μL of TMB Substrate Solution to all wells.

- 16. Incubate the microwell strips at room temperature (18°C to 25°C) for about 30 minutes. Avoid direct exposure to intense light.
 - The color development on the plate should be monitored and the substrate reaction stopped (see next point of this protocol) before positive wells are no longer properly recordable.
 - Determination of the ideal time period for color development has to be done individually for each assay.
 - It is recommended to add the stop solution when the highest standard has developed a dark blue color. Alternatively the color development can be monitored by the ELISA reader at 620 nm. The substrate reaction should be stopped as soon as Standard 1 has reached an OD of 0.9-0.95.
- 17. Stop the enzyme reaction by quickly pipetting $100 \mu L$ of Stop Solution into each well. It is important that the Stop Solution is spread quickly and uniformly throughout the microwells to completely inactivate the enzyme. Results must be read immediately after the Stop Solution is added or within one hour if the microwell strips are stored at $2^{\circ}C$ to $8^{\circ}C$ in the dark.
- 18. Read absorbance of each microwell on a spectro-photometer using 450 nm as the primary wave length (optionally 620 nm as the reference wave length; 610 nm to 650 nm is acceptable). Blank the plate reader according to the manufacturer's instructions by using the blank wells. Determine the absorbance of both the samples and the standards.

Note: In case of incubation without shaking the obtained O.D. values may be lower than indicated below. Nevertheless the results are still valid.

Calculation of results

- Calculate the average absorbance values for each set of duplicate standards and samples. Duplicates should be within 20% of the mean value
- Create a standard curve by plotting the mean absorbance for each standard concentration on the ordinate against the rat IL-17A concentration on the abscissa. Draw a best fit curve through the points of the graph (a 5-parameter curve fit is recommended).
- To determine the concentration of circulating rat IL-17A for each sample, first find the mean absorbance value on the ordinate and extend a horizontal line to the standard curve. At the point of intersection, extend a vertical line to the abscissa and read the corresponding rat IL-17A concentration.
- If instructions in this protocol have been followed, samples have been diluted 1:2 (50 μ L sample + 50 μ L Sample Diluent) and the concentration read from the standard curve must be multiplied by the dilution factor (x 2).
- Calculation of samples with a concentration exceeding standard 1
 may result in incorrect, low rat IL-17A levels. Such samples
 require further external predilution according to expected rat
 IL-17A values with Sample Diluent in order to precisely
 quantitate the actual rat IL-17A level.
- It is suggested that each testing facility establishes a control sample of known rat IL-17A concentration and runs this additional control with each assay. If the values obtained are not within the expected range of the control, the assay results may be invalid.

 A representative standard curve is shown in Figure 8. This curve cannot be used to derive test results. Each laboratory must prepare a standard curve for each group of microwell strips assayed.

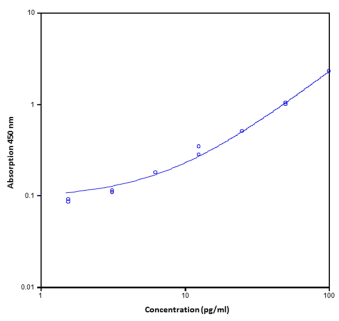


Fig. 8 Representative standard curve for rat IL-17A ELISA. Rat IL-17A was diluted in serial 2-fold steps in Sample Diluent. Do not use this standard curve to derive test results. A standard curve must be run for each group of microwell strips assayed.

Table 2 Typical data using the rat IL-17A ELISA

Measuring wavelength: 450 nm Reference wavelength: 620 nm

Standard	Rat IL-17A Concentration (pg/mL)	0.D. at 450 nm	Mean O.D. at 450 nm	C.V. (%)
1	100.0	2.297 2.300	2.299	0.1
2	50.0	1.037 0.999	1.018	1.8
3	25.0	0.509 0.506	0.508	0.3
4	12.5	0.344 0.282	0.313	9.9
5	6.3	0.178 0.180 0.179	0.4	
6	3.1	0.114 0.108	0.111	2.9
7	1.6	0.091 0.086	0.089	2.6
Blank	0	0.082 0.067	0.075	9.8

The OD values of the standard curve may vary according to the conditions of assay performance (e.g. operator, pipetting technique, washing technique or temperature effects). Furthermore shelf life of the kit may affect enzymatic activity and thus color intensity. Values measured are still valid.

Limitations

- Since exact conditions may vary from assay to assay, a standard curve must be established for every run.
- Bacterial or fungal contamination of either screen samples or reagents or cross-contamination between reagents may cause erroneous results.

- Disposable pipette tips, flasks or glassware are preferred, reusable glassware must be washed and thoroughly rinsed of all detergents before use.
- Improper or insufficient washing at any stage of the procedure will
 result in either false positive or false negative results. Empty wells
 completely before dispensing fresh wash solution, fill with Wash
 Buffer as indicated for each wash cycle and do not allow wells to
 sit uncovered or dry for extended periods.

Performance characteristics

Sensitivity

The limit of detection of rat IL-17A defined as the analyte concentration resulting in an absorbance significantly higher than that of the dilution medium (mean plus 2 standard deviations) was determined to be 1.0 pg/mL (mean of 6 independent assays).

Reproducibility

Intra-assay

Reproducibility within the assay was evaluated in 3 independent experiments. Each assay was carried out with 6 replicates of 7 serum samples containing different concentrations of rat IL-17A. Two standard curves were run on each plate. Data below show the mean rat IL-17A concentration and the coefficient of variation for each sample (see Table 3). The calculated overall intra-assay coefficient of variation was 8.5%.

 $\begin{tabular}{ll} \textbf{Table 3} & \textbf{The mean rat IL-17A concentration and the coefficient of variation for each sample} \\ \end{tabular}$

Sample	Experiment	Mean rat IL-17A Concentration (pg/mL)	Coefficient of Variation (%)
	1	77.62	4.8
1	2	78.87	4.9
	3	70.01	7.8
	1	48.59	7.2
2	2	40.97	7.4
	3	40.80	19.2
	1	21.35	5.3
3	2	21.05	5.1
	3	18.83	13.1
	1	13.56	3.8
4	2	10.88	7.7
	3	12.26	8.3
	1	70.74	4.2
5	2	72.92	5.7
	3	81.50	4.6
	1	24.60	4.8
6	2	25.17	6.8
	3	25.81	8.9
	1	5.42	8.4
7	2	4.53	24.3
	3	4.98	16.1

Inter-assay

Assay to assay reproducibility within one laboratory was evaluated in 3 independent experiments. Each assay was carried out with 6 replicates of 7 serum samples) containing different concentrations of rat IL-17A. Two standard curves were run on each plate. Data below show the mean rat IL-17A concentration and the coefficient of

variation calculated on 18 determinations of each sample (see Table 4). The calculated overall inter-assay coefficient of variation was 7.6%.

Table 4 The mean rat IL-17A concentration and the coefficient of variation of each sample

Sample Mean rat IL-17A Concentration (pg/mL)		Coefficient of Variation (%)
1	75.50	6.4
2	43.45	10.2
3	20.41	6.7
4	12.23	11.0
5	75.05	7.6
6	25.19	2.4
7	4.98	9.0

Spike recovery

The spike recovery was evaluated by spiking 3 levels of rat IL-17A into serum, plasma (EDTA, citrate, heparin) and cell culture supernatant samples. Recoveries were determined with 4 replicates each. For recovery data see Table 5.

The unspiked serum, plasma, cell culture supernatant was used as blank in these experiments.

Recoveries were shown to depend on the serum used.

Table 5 Spike recovery

Sample matrix	Spike high (%)	Spike medium (%)	Spike low (%)
Serum	44	34	39
Plasma (EDTA)	31	20	26
Plasma (citrate)	59	37	29
Plasma (heparin)	64	45	37
Cell culture supernatant	112	92	103

Dilution parallelism

Serum, plasma cell culture supernatant samples with different levels of rat IL-17A were analyzed at serial 2-fold dilutions with 4 replicates each. For recovery data see Table 6.

Table 6 Dilution parallelism

Comple metric	Recovery of Exp. Val.		
Sample matrix	Range (%)	Mean (%)	
Serum	85 - 122	102	
Plasma (EDTA)	93 - 125	107	
Plasma (citrate)	95 - 114	106	
Plasma (heparin)	64 - 126	97	
Cell culture supernatant	79 - 89	86	

Sample stability

Freeze-Thaw stability

Aliquots of spiked serum samples were stored at -20°C and thawed 5 times and the rat IL-17A levels determined. There was no significant loss of rat IL-17A immunoreactivity detected by freezing and thawing.

Storage stability

Aliquots of spiked serum samples were stored at -20° C, 2° C to 8° C room temperature, and at 37° C, and the rat IL-17A level determined after 24 hours. There was no significant loss of rat IL-17A immunoreactivity detected during storage at -20° C and 2° C to 8° C. A significant loss of rat IL-17A immunoreactivity was detected during storage at room temperature and at 37° C after 24 hours.

Specificity

Crossreactivity and interference of circulating factors of the immune system were evaluated by spiking these proteins at physiologically relevant concentrations into a rat IL-17A positive sample.

There was no crossreactivity detected, notably not with rat IFN γ , rat TNF α , rat IL-1 α , rat IL-4, rat MCP-1, rat GM-CSF.

Expected values

There were no detectable rat IL-17A levels found. Elevated rat IL-17A levels depend on the type of immunological disorder.

Reagent preparation summary

Wash buffer (1x)

Add Wash Buffer Concentrate 20x (50 mL) to 950 mL distilled water.

Number of Strips Wash Buffer Concentrate (ml		Distilled Water (mL)
1 - 6	25	475
1 - 12	50	950

Assay buffer (1x)

Add Assay Buffer Concentrate 20x (5 mL) to 95 mL distilled water.

Number of Strips	Assay Buffer Concentrate (mL)	Distilled Water (mL)
1 - 6	2.5	47.5
1 - 12	5.0	95.0

Biotin-Conjugate

Make a 1:100 dilution of Biotin-Conjugate in Assay Buffer (1x):

Number of Strips	Biotin-Conjugate (mL)	Assay Buffer (1x) (mL)
1 - 6	0.03	2.97
1 - 12	0.06	5.94

Streptavidin-HRP

Make a 1:200 dilution of Streptavidin-HRP in Assay Buffer (1x):

Number of Strips	Streptavidin-HRP (mL)	Assay Buffer (1x) (mL)
1 - 6	0.03	5.97
1 - 12	0.06	11.94

Rat IL-17A standard

Reconstitute lyophilized rat IL-17A standard with distilled water. (Reconstitution volume is stated on the label of the standard vial.)

Test protocol summary

- 1. Determine the number of microwell strips required.
- 2. Wash microwell strips twice with Wash Buffer.
- 3. Standard dilution on the microwell plate: Add 100 μ L Sample Diluent, in duplicate, to all standard wells Pipette 100 μ L prepared standard into the first wells and create standard dilutions by transferring 100 μ L from well to well. Discard 100 μ L from the last wells.

Alternatively external standard dilution in tubes (see "External standard dilution" on page 3): Pipette 100 μL of these standard dilutions in the microwell strips.

- **4.** Add 100 μL Sample Diluent, in duplicate, to the blank wells.
- 5. Add 50 μL Sample Diluent to sample wells.
- 6. Add $50 \mu L$ sample in duplicate, to designated sample wells.
- 7. Prepare Biotin-Conjugate.
- 8. Add 50 μL Biotin-Conjugate to all wells.
- Cover microwell strips and incubate 2 hours at room temperature (18°C to 25°C).
- 10. Prepare Streptavidin-HRP.
- 11. Empty and wash microwell strips 4 times with Wash Buffer.
- 12. Add 100 µL diluted Streptavidin-HRP to all wells.

- **13.** Cover microwell strips and incubate 1 hour at room temperature.
- 14. Empty and wash microwell strips 4 times with Wash Buffer.
- 15. Add 100 µL of TMB Substrate Solution to all wells.
- Incubate the microwell strips for 30 minutes at room temperature (18°C to 25°C).
- 17. Add 100 μL Stop Solution to all wells.
- 18. Blank microwell reader and measure color intensity at 450 nm.

Note: If instructions in this protocol have been followed, samples have been diluted 1: 2 (50 μ L sample + 50 μ L Sample Diluent) and the concentration read from the standard curve must be multiplied by the dilution factor (x 2).

Customer and technical support

Visit **thermofisher.com/support** for the latest service and support information.

- Worldwide contact telephone numbers
- Product support information
 - Product FAQs

- Software, patches, and updates
- Training for many applications and instruments
- Order and web support
- Product documentation
 - User guides, manuals, and protocols
 - Certificates of Analysis
 - Safety Data Sheets (SDSs; also known as MSDSs)

Note: For SDSs for reagents and chemicals from other manufacturers, contact the manufacturer.

Limited product warranty

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For descriptions of symbols on product labels or product documents, go to thermofisher.com/symbols-definition.

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