



Anti-CCP Fast Test Kit (Immunofluorescence Assay)

Instructions for Use

INTENDED USE

Anti-CCP Fast Test Kit (Immunofluorescence Assay) is intended for *in vitro* quantitative determination of Anti-CCP in serum, plasma or whole blood samples. The test is designed to aid in the diagnosis of autoimmune diseases, such as rheumatoid arthritis (RA). For professional and laboratory use only.

SUMMARY

Anti-CCP is an autoantibody that synthesizes cyclic citrulline polypeptide (CCP) as antigens. Anti-CCP mainly consists of IgG and it can be used for the early diagnosis of RA due to its high sensitivity and specificity to the disease. Currently, it is believed that the sensitivity of Anti-CCP to RA diagnosis is 50%-78% with a specificity of 96%. The positive rate of early RA patients is 80%. Meanwhile, radiological examination results show that patients with positive Anti-CCP have significantly more severe joint injuries than those with negative Anti-CCP. Therefore, Anti-CCP as a prognostic indicator can play an important role in monitoring the development and progression of rheumatoid arthritis disease. In addition, Anti-CCP is a sensitive indicator for differentiating invasive and non-invasive RA.

PRINCIPLE

Anti-CCP Fast Test Kit (Immunofluorescence Assay) adopts an indirect sandwich method to quantitatively detect the concentration of Anti-CCP in human serum, plasma and whole blood samples. After the sample has been added to the test card, the fluorescently labelled CCP antigen combined with Anti-CCP in sample and forms a marked antigen-antibody complex. The complex moves to the detection area by capillary action. Then it is captured by Anti-human IgG antibody coated on the detection area of nitrocellulose membrane, forming a CCP-Anti-CCP-Anti-human IgG complex. The fluorescence intensity of the test line increases in proportion to the amount of Anti-CCP in sample. Fluorescent signals intensity can be analyzed by applicable device thus the Anti-CCP in sample be detected quantitatively.

APPLICABLE DEVICE

Getein 1100 Immunofluorescence Quantitative Analyzer
Getein 1160 Immunofluorescence Quantitative Analyzer
Getein 1180 Immunofluorescence Quantitative Analyzer
Getein 1600 Immunofluorescence Quantitative Analyzer
Getein 1200 Immunofluorescence Quantitative Analyzer

CONTENTS

| Materials provided | Getein 1100/Getein 1160/ Getein 1180 | | Getein 1200/ Getein 1600 | |
|----------------------|---|----------|---|---|
| | 10 T/kit | 25 T/kit | 2*24 T/kit | 2*48 T/kit |
| Anti-CCP test card* | 10 pcs | 25 pcs | 24 test cards in 1 cartridge, and 2 cartridges in 1 box | 48 test cards in 1 cartridge, and 2 cartridges in 1 box |
| Disposable pipet | 10 pcs | 25 pcs | / | / |
| Sample diluent** | 10 tubes | 25 tubes | 1 box | 1 box |
| Instructions for use | 1 pc | 1 pc | 1 pc | 1 pc |
| SD card | 1 pc | 1 pc | 1 pc in each cartridge | 1 pc in each cartridge |

* Anti-CCP test card

A test card consists of: Fluorescently labelled CCP antigen, Anti-human IgG antibody.

** Sample diluent

(1) Sample diluent for Getein 1100/ Getein 1160/ Getein 1180 in each tube mainly consists of: phosphate buffer (20 mmol/L), NaH_2PO_4 (<0.1%).

(2) Sample diluent for Getein 1200/ Getein 1600 is an independent packing box mainly consists of:

-Phosphate buffer (20 mmol/L), NaH_2PO_4 (<0.1%) (25 mL/bottle for Getein 1200, 30 mL/bottle for Getein 1600),

-Box with pipette tips (96 tips/box),

-Mixing plate (1 piece/box).

Note:

- The standard curve data can be written to RFID card in the kit. According to the function of RFID card, we define it as "Standard Curve Data Card", short for "SD Card".
- Do not mix or interchange different batches of kits.

STORAGE AND STABILITY

Realtime stability:

Store the kit at 4-30°C with a valid period of 24 months. The test kits are stable until the expiry date printed on the labels.

In-use stability:

-For the test card of Getein 1100/Getein 1160/Getein 1180: Use the test card within 1 hour once the foil pouch is opened.

-For test card of Getein 1200/Getein 1600: if the cartridge is opened, it could be stable within 24 hours once exposed to air.

If the test cards can't be used up at a time, please use the test cards back to the foil pouch and reseal along the entire edge of zip-seal. The remaining test cards should be used up within 7 days.

PRECAUTIONS

- For *in vitro* diagnostic use only.
- For professional and laboratory use only, not for near-patient test and self-testing.
- Do not use the test card if the foil pouch or the cartridge is damaged.
- Do not open pouches until performing the test.
- Handle all specimens as potentially infectious. The foil bag is non-degradable. Proper handling and disposal methods should be followed in accordance with local regulations.
- It is recommended that operators take necessary self-protection measures (work clothes, goggles and disposable gloves, etc) when touching kits or samples.

SPECIMEN COLLECTION AND PREPARATION

- Serum, plasma or whole blood samples** can be used for the test. Other body fluids and samples may not give accurate results. Samples should be free of hemolysis.
- Venous blood should be collected under aseptic conditions; serum or plasma is preferred for testing.
- Heparin, sodium citrate or EDTA can be used as the anticoagulant for plasma and whole blood samples.
- It is recommended to test the sample within 4 hours after collection. Stable in serum and plasma for 7 days at 2-8°C and 3 months at -20°C. Stable in whole blood for 3 days when stored at 2-8°C.
- Refrigerated or frozen sample should be reached to room temperature before testing. Frozen samples must be completely thawed, rewarmed and evenly mixed. Serum and plasma samples can be freeze and thawed twice at most.
- Sample volume (**for Getein1100/Getein1160/Getein1180**): **10 µL**

TEST PROCEDURE

- User must carefully read and operate in strict accordance with the instructions for use before testing, otherwise reliable results cannot be guaranteed.
- Test kit and sample should be brought to room temperature before testing.

For Getein 1100:

- Confirm SD card lot No. in accordance with test kit lot No. It is required to perform "SD card" calibration when using a new batch of kits.
- Select the corresponding "Sample" on the analyzer according to the sample type (see the user manual of analyzer for details).
- Remove the test card from the sealed pouch before use. Horizontally place the test card.

(4) Deliver 10 µL of sample into one tube of sample diluent using disposable pipet or pipette, mix gently and thoroughly. Then drop 100 µL of sample mixture into the sample well on the test card.

(5) **Reaction time: 15 minutes.** After reaction time is elapsed, insert the test card into Getein 1100 and press "ENT" button (click on "Start" icon for Android Getein 1100). The result will be shown on the screen and printed automatically.

For Getein 1160/Getein 1180:

- Confirm SD card lot No. in accordance with test kit lot No. It is required to perform "SD card" calibration when using a new batch of kits.
- Select the corresponding "Sample" on the analyzer according to the sample type (see the user manual of analyzer for details).
- Remove the test card from the sealed pouch before use. Horizontally place the test card.
- Deliver 10 µL of sample into one tube of sample diluent using disposable pipet or pipette, mix gently and thoroughly. Then drop 100 µL of sample mixture into the sample well on the test card.
- Insert the test card into Getein 1160/Getein 1180 **immediately** after sample loading. The analyzer will count down the reaction time (15 minutes) and automatically test the card after reaction time is elapsed. The result will be shown on the screen and printed automatically.

For Getein 1200/Getein 1600:

- Each cartridge for Getein 1200/Getein 1600 contains a specific RFID card (SD card) which can calibrate automatically.
- Place the sample diluent at the correct position in Getein 1200/Getein 1600.
- Place samples in the designed area of the sample holder, insert the holder, set parameters (more operational details refer to the user manual of analyzer) and run the instrument, Getein 1200/ Getein 1600 will do the testing and print the result automatically.

Note:

- It is required to perform "SD card" calibration when using a new batch of kits for Getein 1100/Getein 1160/Getein 1180.
- The directions for using disposable pipet are as follows:

Directions to use disposable pipet

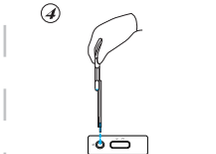
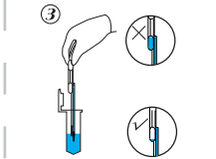
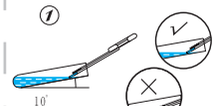
Insert the disposable pipet into the sample tube, gently touch the liquid surface with the capillary tip, and draw the sample.

Note: Do not immerse the exhaust pipe below the liquid level.

Insert the disposable pipet (including the exhaust tube) into the dilution liquid, gently squeeze the suction bulb to perform 2-3 aspiration washing cycles, then mix the dilution manually.

Insert the disposable pipet (including the exhaust tube) into the dilution liquid, firmly squeeze the suction bulb to aspirate the mixed sample.

Squeeze the suction bulb and drop the mixed sample vertically into the sample well on the test card.



symptoms.

2. Some substances in blood as listed below may interfere with the test and cause erroneous results. The maximum allowance concentration of each is as follows:

| Interferent | Concentration (Max) |
|--------------|---------------------|
| Triglyceride | 10g/L |
| Bilirubin | 0.2g/L |

EXPECTED VALUE

The expected normal value for Anti-CCP is determined by testing samples from 282 apparently healthy individuals. The upper 99.0th percentile value is 25.0 U/mL. It is recommended that each laboratory determine the applicability of the reference value through experiments and establish its own reference ranges if necessary.

PERFORMANCE CHARACTERISTICS

| | |
|-----------------------|-----------------|
| Measuring Range | 10.0-400.0 U/mL |
| Limit of Detection | ≤10.0 U/mL |
| Within-Run Precision | ≤10% |
| Between-Lot Precision | ≤15% |

REFERENCES

1. Ishigooka, Nozomi, Fujii Takao, et al. Predicting factors for disappearance of anti-mutated citrullinated vimentin antibodies in sera of patients with rheumatoid arthritis [J]. *Modern rheumatology*, 2020, 30(3):450-457.
2. Tülay Kars Fertelli. Effects of Education About Rheumatoid Arthritis and Sexuality on the Sexual Problems of Women With Rheumatoid Arthritis [J]. *Clinical Nursing Research*, 2020, 29(3):189-199.
3. Verheul M K, Bhringer S, Mam V D, et al. The combination of three autoantibodies, ACPA, RF and anti-CarP antibodies is highly specific for rheumatoid arthritis: implications for very early identification of individuals at risk to develop rheumatoid arthritis [J]. *Arthritis and Rheumatism*, 2018, 70(11):1721-1731.
4. Shakiba Yadollah, Koopah Susan, Jamshidi Ahmad Reza, et al. Anti-cyclic citrullinated peptide antibody and rheumatoid factor isotypes in Iranian patients with rheumatoid arthritis: evaluation of clinical value and association with disease activity [J]. *Iranian Journal of Allergy, Asthma and Immunology*, 2014, 13(3):147-156.

DESCRIPTION OF SYMBOLS USED

The following graphical symbols used in or found on the test kit are the most common ones appearing on medical devices and their packaging. They are explained in more details in the European Standard EN ISO 15223-1:2021.

| Key to symbols used | | | |
|---------------------|---|--|---|
| | Manufacturer | | Use-by date |
| | Do not re-use | | Date of manufacture |
| | Consult instructions for use or consult electronic instructions for use | | Batch code |
| | Temperature limit | | In vitro diagnostic medical device |
| | Contains sufficient for <n> tests | | Authorized representative in the European Community/ European Union |
| | CE mark | | Do not use if package is damaged and consult instructions for use |
| | Catalogue number | | Caution |

Thank you for using Anti-CCP Fast Test Kit (Immunofluorescence Assay). Please read the instructions for use carefully before operating to ensure proper use.

Version: WIF106-S-06

Getein Biotech, Inc.
 Add.: No.9 Bofu Road, Luhe District, Nanjing, 211505, China
 Tel: +86-25-68568508
 Fax: +86-25-68568500
 E-mail: tech@getein.com.cn
 overseas@getein.com.cn
 Website: www.getein.com

CMC Medical Devices & Drugs S.L.
 Add.: C/ Horacio Lengo Nº 18, CP 29006, Málaga, Spain
 Tel: +34951214054

| Catalogue number | Applicable analyzer | Package specification |
|------------------|---------------------|-----------------------|
| IF1029-10T | Getein 1100 | 10 T/kit |
| IF1029 | Getein 1100 | 25 T/kit |
| IF5029-10T | Getein 1160 | 10 T/kit |
| IF5029 | Getein 1160 | 25 T/kit |
| IF3029-10T | Getein 1180 | 10 T/kit |
| IF3029 | Getein 1180 | 25 T/kit |
| IF4029 | Getein 1200 | 2*24 T/kit |
| IF4029-96T | Getein 1200 | 2*48 T/kit |
| IF2029 | Getein 1600 | 2*24 T/kit |
| IF2029-96T | Getein 1600 | 2*48 T/kit |

RESULTS

Getein 1100/Getein 1160/Getein 1180/Getein 1200/Getein 1600 can scan the test card automatically and display the result on the screen. For additional information, please refer to the user manual of Getein 1100/Getein 1160/Getein 1180/Getein 1200/Getein 1600.

Others: Samples whose concentration exceeds the upper limit should be diluted no more than 4 times.

LIMITATIONS

1. As with all diagnostic tests, a definitive clinical diagnosis should not be made based on the result of a single test. The test results should be interpreted considering all other test results and clinical information such as clinical signs and