



COVID-19 Antigen Rapid Test Kit (Saliva)

Manufacturer disposal procedure

Hunan Dear Sens Biotechnology Co., Ltd.

Control and management of nonconforming products

1. General

a. Inspect and test raw materials, semi-finished products and finished products according to inspection and test management procedure and relevant technical and quality documents, and determine whether they are qualified or unqualified.

b. The nonconforming products shall be identified and isolated according to the nonconforming product management procedure, and records shall be made to determine the scope of nonconformity (when applicable); - Time, batch, quantity, etc.) shall not be transferred to the next process, warehousing and delivery before treatment.

c. The quality control department is responsible for organizing relevant departments to investigate and review nonconforming products, Analyze the causes and authorize relevant departments to rework, reject or scrap; The evaluation of nonconformities shall include determining whether it is necessary to investigate and notify all external parties responsible for nonconformities.

Keep records of the nature of nonconformity and any subsequent measures taken, including the reasons for evaluation, any investigation and decision-making.

2. Response measures for nonconforming products found before delivery.

The company disposes of nonconforming products through one or more of the following ways:

- a. Take measures to eliminate the found nonconformities;
- b. Take measures to prevent its original intended use or application;
- c. Authorize concession use, release or acceptance.

The quality control department shall ensure that nonconforming products are only provided with reasons. Concession acceptance can be made only after obtaining approval and meeting the requirements of applicable regulations. And keep records of concession acceptance and the identity of concession authorized personnel.

3. Response measures for nonconforming products found after delivery

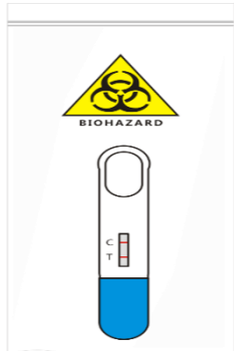
When nonconforming products are found after delivery or use, the company shall take measures appropriate to the impact or potential impact of the nonconformity.

Records of actions taken shall be maintained. If the determined response measures for nonconforming products are found after delivery, they shall be carried out in accordance with the applicable regulations.

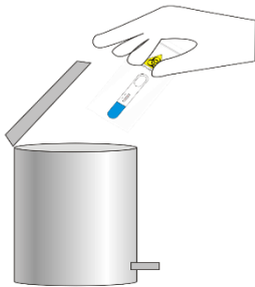
Regulations require that the procedures for issuing Advisory notices be documented, according to the company's management procedure for issuance and implementation of advisory notice and management.

4. Disposal the sample and clean-up

After testing, place the test card in the biohazard waste bag and seal the bag.



Throw away the remaining sample kit items.



Re-apply hand sanitizer.



5. Precautions

- 5.1. Please equilibrate the test card to room temperature (above 20min) before testing.
- 5.2. The inspection should be carried out in strict accordance with the instructions.
- 5.3. The result must be interpreted within 15-20min, and the result read after 20min is invalid.
- 5.4. The test sample should be regarded as an infectious substance, take protective measures and pay attention to biosafety operations.
- 5.5. The used test cards are treated as biomedical waste after the test, and wash your hands in time.
- 5.6. Do not use the kit with obvious damage, and test card with damaged package.
- 5.7. This product is a one-time use product, please do not reuse it, and do not use expired products.
- 5.8. Avoid direct sunlight and direct blowing from electric fans during testing.

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5.9. Tap water, distilled water or deionized water and beverages cannot be used for testing.

5.10. Due to the difference of the samples, some test lines may be lighter or grayish in color. As a qualitative product, as long as there is a band at the position of the T line, it can be judged as positive.

5.11. If the test is positive, it is recommended to use this test card to recheck once to avoid small probability events.

5.12. There is a desiccant in the aluminum foil bag, do not take it orally.