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**QUESTION 1**

After submission to the regulatory authority, a substantial error was found in the application. In order to resolve this issue, what should be done FIRST?

- A. Resubmit the entire package.
- B. Inform upper management immediately.
- C. Contact the legal department and ask them how to proceed.
- D. Verify the procedure in the regulation for the corrections.

Correct Answer: D

QUESTION 2

According to WHO, what are the temperature and humidity conditions for a Zone IVb long-term stability study?

- A. 25: C and 60% RH
- B. 30°C and 35% RH
- C. 30°C and 65% RH
- D. 30: C and 75% RH

Correct Answer: D

QUESTION 3

A company is developing a device-drug combination product. Which of the following should be evaluated FIRST in order to determine the applicable guidance documents?

- A. Approved indications of the drug
- B. Determination of primary mode of action
- C. Determination of product design deliverables
- D. Guidance documents for the device

Correct Answer: C

QUESTION 4

Who has the PRIMARY responsibility for recall of products with quality defects?



- A. Consumer
- B. Distributor
- C. Manufacturer
- D. Regulatory authority

Correct Answer: C

QUESTION 5

A materials supplier informs a company that it intends to stop supplying a material critical to the manufacture of the company's products. What action should the company take FIRST?

- A. Review the company's existing Quality Management System
- B. Reformulate the products with a replacement material.
- C. Qualify another supplier and execute a supplier agreement.
- D. Complete a gap analysis to identify options.

Correct Answer: CD

QUESTION 6

SOPs for preventive and corrective actions MUST include the procedure to eliminate which of the following?

- A. Inadequate training
- B. Late and/or incorrect deliverables
- C. Causes of non-conformities
- D. Adverse environmental impacts

Correct Answer: C

QUESTION 7

What is the BEST approach to ensure that raw materials, services, and sub-contractors at the level of the vendors comply with GMP requirements?

- A. Ask the vendor to take responsibility.
- B. Document and perform audits.



- C. Request an inspection from a regulatory authority.
- D. Request documentation from the sub-contractor.

Correct Answer: B

QUESTION 8

Company X is planning to acquire the rights for a product marketed by Company Y. As part of due diligence, what is the MOST important information the Company X regulatory affairs professional should ask senior management to request from Company Y?

- A. Intellectual property
- B. Clinical trial data
- C. Safety issues
- D. Marketing materials

Correct Answer: C

QUESTION 9

A regulatory authority announces an inspection of a regulatory affairs professional's facility during a holiday season when most of the staff is not available. What is the MOST practical approach to this dilemma?

- A. Negotiate with colleagues and the authority to find a better time.
- B. Insist that key personnel be available for the inspection.
- C. Inform the authority that the time is not suitable and request a new time
- D. Arrange for an inspection without all intended personnel.

Correct Answer: A

QUESTION 10

A company is currently marketing an implantable orthopedic medical device. The RandD department is planning to change the material used for the implant. The RandD department states that the change does not impact the safety and effectiveness of the product.

What action should the regulatory affairs professional take FIRST?

- A. No action is needed in this situation.
- B. Prepare regulatory submissions that detail the medical device's change in materials.



- C. Review the content of change and supporting data for the equivalency with the current material.
- D. Write a memo to file since the change does not impact product safety and effectiveness.

Correct Answer: C

QUESTION 11

A drug product presents degradation during the manufacturing process. In addition to the amount, what information should be provided FIRST in order to use API overage?

- A. Specification
- B. Formulation
- C. Property
- D. Justification

Correct Answer: D

QUESTION 12

After numerous failed attempts to decrease an identified risk in a medical device to an acceptable level, the medical device continues to have unacceptable risks. However, the development team wants to continue development. Which is the BEST recommendation to make in this situation?

- A. Add a warning in the IFU.
- B. Discontinue the project.
- C. Perform another risk-benefit analysis.
- D. Redesign the device.

Correct Answer: D

QUESTION 13

Which of the following double-blind clinical trial designs would be MOST appropriate for a Phase III study with a new product intended to treat an acute life-threatening disease with less than optimal available therapy?

- A. Active-controlled
- B. Cross-over
- C. Dose-ranging
- D. Placebo-controlled

Correct Answer: B



QUESTION 14

Which question is pertinent to demonstrate a new pharmaceutical's effectiveness during evaluation by a reimbursement agency?

- A. "Is the product profitable for the manufacturer?"
- B. "Is the product better than currently available alternatives?"
- C. "Has the product been approved for more than 10 years?"
- D. "Is the product an established gold standard?"

Correct Answer: B

QUESTION 15

A regulation change is imminent and may require further non-clinical testing on a product currently in Phase III clinical trials. What is the most appropriate action to take FIRST?

- A. Obtain a copy of the proposed regulation and analyze the impact.
- B. Inform the company's senior management and arrange an emergency meeting
- C. Consult with the company's legal department regarding options.
- D. Arrange for additional testing of the product at the testing facility.

Correct Answer: A

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