

# Periodic surveillance of rDNA activities by the IBC

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## Human gene therapy trials

- A. Changes in vectors or inserts
- B. Serious adverse events (SAEs)
- C. IBC responses to violations
- D. Flaws in program of surveillance

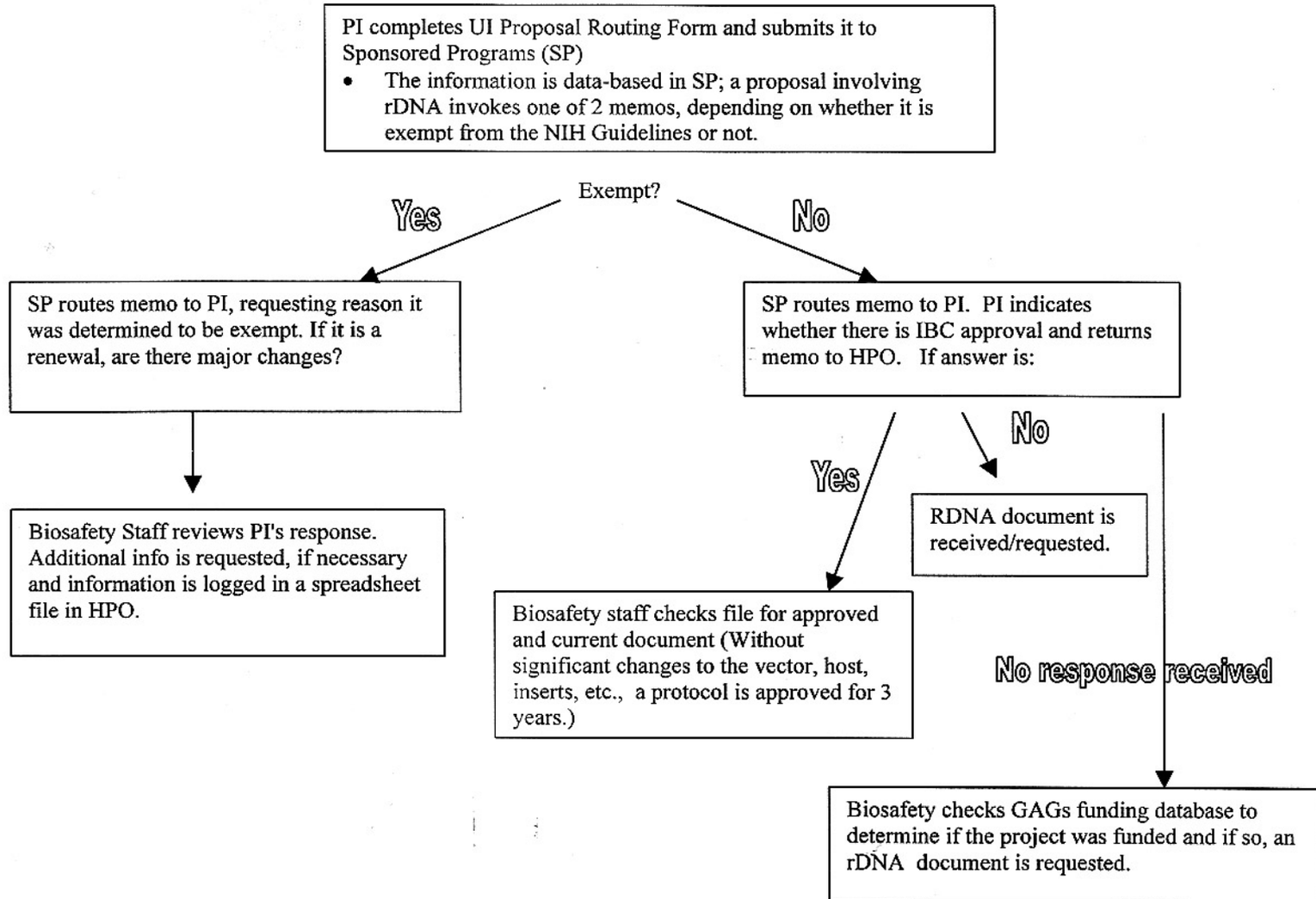
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rDNA work not involving humans

- A. Flow chart
- B. IBC responses to incidents and violations
- C. Flaws in program of surveillance

# rDNA Annual Review Process



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The need for regulation and surveillance

- A. Human gene therapy trials (Group I)
- B. rDNA work involving genetic alterations of animals, plants, and microorganisms, or toxin-encoding genes (Group II)
- C. None of the above (Group III)