

CORRECTION FACTOR:

DOSE REQUIRED:

SIGNED:

## Microbiological Validation Services Ltd **Method 1 Submission Form** CUSTOMER NAME: PURCHASE ORDER NO: NUMBER OF SAMPLES SUBMITTED: **DESCRIPTION OF SAMPLES:** IF PRODUCT IS FOR ANY OTHER METHOD THAN GAMMA, PLEASE SPECIFY: Please select one of the 3 options below: 1. JUSTIFICATION / SUBSTANTIATION FOR MULTIPLE BATCHES OF PRODUCT YES / NO NOTE: Follows ISO 11137-2 'Procedure for Method 1 Dose Setting for multiple production batches'. Requires non-sterile samples, at least 10 samples from each of 3 production batches for bioburden testing and 100 samples for tests of sterility, 130 samples in total. 2. RE-JUSTIFICATION / AUDIT OF DOSE SUBSTANTIATED IN ORIGINAL EXERCISE YES / NO TARGET DOSE (kGy): NOTE: Follows ISO 11137-2 'Procedure for auditing a sterilization dose substantiated using Method 1 Dose Audit'. Requires non-sterile samples, at least 10 samples from a single production batch for bioburden testing and 100 samples for tests of sterility, 110 samples in total. IF OPTION 2 IS SELECTED PLEASE PROVIDE REPORT REFERENCE OF ORIGINAL JUSTIFICATION / SUBSTANTIATION: If the original dose setting was carried out by a different laboratory, the full report must be provided before work commences. 3. SINGLE PRODUCTION BATCH VALIDATION YES / NO NOTE: Follows ISO 11137-2 'Procedure for Method 1 Dose Setting for a single production batch'. Requires non-sterile samples, at least 10 samples from a single production batch for bioburden testing and 100 samples for tests of sterility, 110 samples in total. Valid only for the sterilization of the batch of product submitted. PRODUCT REFERENCE: BATCH NUMBER(S); FOR OFFICE USE ONLY: ANY OTHER INFORMATION:

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STASIS REQUIRED: BIOBURDEN / STERILITY

DOSE RANGE:

DATE:

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