

Audit #: _____ Auditor(s): ___

GILES CHEMICAL ~ PREMIER MAGNESIA

Company Form

Number: Q12-PR-100-F008g

Title: Internal Audit Checklist - Control of

Documents, Records and Reports

Owner: Deborah Durbin Revision: 0
Effective Date: 05/04/16 Page: 1 of 3

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| | 110 |

Date: _____

| Subpart J | | |
|---|--|--------------------------|
| Records and Record Keeping - General | Document(s) Reviewed/Person(s) Interviewed/Objective Evidence/Comments: | Conforms to Requirements |
| All documents related to the quality system are prepared, reviewed, approved and distributed according to written procedure. | | Yes No |
| The issuance, revision, superseding and withdrawal of all documents are controlled and records of these activities are maintained in revision histories or equivalent. | | Yes No |
| Procedures exist describing cGMP-recordkeeping practices. E.g. permanent ink, identification of "who" and "when" for all entries and procedures for correcting entries (sign, date, explain and not obliterate original entry). | | ☐ Yes ☐ No |
| Procedures have been established that describe the requirements for record retention. | | Yes No |
| Records will be maintained for 1 year after the expiration date or 3 years beyond the date of distribution of the last batch associated with those records. | | ☐ Yes ☐ No |
| All records are maintained as original record, as true copies or electronic records. | | ☐ Yes ☐ No |
| Annual Product Review and Inspection of Reserve Samples | Document(s) Reviewed/Person(s) Interviewed/Objective Evidence/Comments: | Conforms to Requirements |
| The Quality Unit performs an Annual Product Review [21 CFR 211.180(e)]. | | ☐ Yes ☐ No |
| The Quality Unit perform an annual physical inspection of a representative number of reserve samples [21 CFR 211.170(b)]. | | ☐ Yes ☐ No |



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| Product Complaints and Complaint Files: 21 CFR | Document(s) Reviewed/Person(s) Interviewed/Objective | Conforms to |
|---|--|--------------|
| 211.198 | Evidence/Comments: | Requirements |
| Procedures have been established describing how product | | Yes No |
| complaints will be received, investigated and documented. | | |
| All product complaints have been reviewed to determine if | | |
| the complaint was the result of a product specification failure | | Yes No |
| or quality. | | |
| The decision to investigate a complaint as well as the final | | |
| decision as a result of the investigation, including corrective | | ☐ Yes ☐ No |
| action, has been approved by the Quality Unit. | | |
| The investigation for a product complaint was approximately | | ☐ Yes ☐ No |
| extended to other batches, products, processes, etc. | | |
| Product complaint information has included adequate | | ☐ Yes ☐ No |
| information. | | |
| Procedures for handling complaints include provisions for | | |
| investigation and, if necessary, reporting of serious adverse | | Yes No |
| events as outlined in 21 CFR 310.305. | | |
| Procedures have been established to define the recall of a | | |
| product and a mock recall exercise is conducted at least | | Yes No |
| annually. | | |
| Control of Documents and Records – Electronic Records | Document(s) Reviewed/Person(s) Interviewed/Objective | Conforms to |
| and Electronic Signatures | Evidence/Comments: | Requirements |
| Any GMP related computerized systems have been validated. | | Yes No |
| Procedures and controls have been established for electronic | | |
| closed systems used to create, modify, maintain or transmit | | |
| electronic records in order to ensure the authenticity, | | |
| integrity and confidentiality of the records [Closed System] | | Yes No |
| (21 CFR Part 11). | | |
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| Procedures and controls have been established for use of open electronic systems. Areas of control have been identified, as necessary, per the requirements of 21 CFR Part 11. | ☐ Yes ☐ No |
|--|------------|
| Electronic signatures conform to requirements. | Yes No |
| Passwords and codes have been established. | Yes No |
| Backup electronic files have been maintained of the following: current software programs, outdated software programs that may be necessary to retrieve past records and data that was entered. | ☐ Yes ☐ No |
| Backup files are an exact and complete record and are secure from alterations, erasures or loss and damage. | ☐ Yes ☐ No |