

## Site Quality Self-Assessment

based on

## **Rx-360 Supplier Assessment Questionnaire**

Module 2, Site Specific Information

Relevant for

Sigma Aldrich Chimie S.a.r.l 80 Rue de Luzais, L'Isle D'Abeau Chesnes St. Quentin Fallavier Cedex 38297 France

The site self-assessment covers our quality management system for the following applications:

- distribution and warehouse



Merck KGaA, Darmstadt, Germany is an active member of the Rx 360 Consortium

As a trusted partner of our customers, we deliver quality - always.

Merck KGaA, Darmstadt, Germany Corporation with General Partners Frankfurter Str. 250 64293 Darmstadt, Germany Phone +49 6151 72-0 Sigma-Aldrich Corporation A subsidiary of Merck KGaA, Darmstadt, Germany 3050 Spruce Street St. Louis, MO 63103, USA Phone +1 (800) 521-8956 / +1 (314) 771-5765 EMD Millipore Corporation A subsidiary of Merck KGaA, Darmstadt, Germany 400 Summit Drive Burlington, MA 01803, USA Phone +1 (781) 533-6000



## **Information**

This document is based on the Rx-360 Consortium's Supplier Assessment Questionnaire template, Module 2. The contents of this questionnaire are built on the Rx-360 questionnaire version 2.0 intact with no question added or deleted.

Rx-360's CEO/COO gave permission to Life Science to use the Rx-360 logo.



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## Rx-360 Supplier Assessment Questionnaire : Site-Specific Information

		nts are attached
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	SECTION 1. General Site Information			
1.1	Site or Facility-Specific Name: Sigma Aldrich Chimie S.a.r.l.			
1.2	Address: Sigma Aldrich Chimie S.a.r.l 80 Rue de Luzais L' Isle D'Abeau Chesnes St. Quentin Fallavier Cedex 38297 France			
	GPS Coordinates: latitude 45.644640, Longitude 5.095650			
1.3	Phone: 0800 21 14 08			
1.4	Email: please contact your responsible Service partner			
1.5	Fax: please contact your responsible Service partner			
1.6	Website: www.sigmaaldrich.com			

	SECTION 2. General Site Operating Information					
2.1	What year did the site start operating? 1987					
2.2	What is the primary activity of the site? (e.g. manufacturing, distribution, etc.)  Distribution					

	SECTION 2. General Site Operating Information				
2.3	To which, if any, subdivision of the parent company does the site belong? Life Science business of Merck KGaA, Darmstadt Germany				
2.4	Size of site (in sq. ft. or m.): 7000 squaremeter warehouse				
2.5	Please list or attach the normal hours/schedule of the facilities, including shutdown dates (if applicable):  Warehouse operation hours: 0830H - 1730H  Security: 24 hours  Warehouse operation hours: 0830H - 1730H				
2.6	Total number of employees on site: 175				
2.7	Total number of employees in Quality: 2				
2.8	Total number of employees in Manufacturing: N/A				
2.9	What quality management system is utilized on site?  ☐ ISO 9001 ☐ ISO 13485 ☐ 21 CFR Part 210/211 ☐ 21 CFR Part 820 ☐ European GMP, Eudralex Volume 4 Part I ☐ European GMP, Eudralex Volume 4 Part II ☐ ICH Q7 ☐ HACCP ☐ ISO 22000 ☐ Other Please describe: GDP  Which Regulatory Initiatives does the site follow/comply with? ☐ REACH ☐ RoHs ☐ Ca Prop. 65 ☐ WEEE				

	SECTION 2. General Site Operating Information					
2.10	Does the company/site  \( \sum \)	Tes	No [	N/A		
2.11	Is the site registered with any gove GMP certification, etc.)?  Yes No If yes, please specify.  ANSM	ernment regulat	ory agency (F	DA registration,		
2.12	By whom is the site inspected (reg the last three years: inspection visit of local ANSM	ulatory or third	party) and lis	et inspections within		
2.13	How often, as an annual average, i	s the site audite	ed by custome	ers or third parties?		
2.14	Has an Rx-360 audit been performed Please also state the date of the audithttp://rx-360.org/audit-programs/		Yes	⊠ No		
	interview of the second					
2.15	Are you willing to have Rx-360 con according to the Rx-360 audit program Yes No		•	ur customers		
2.16	Are you willing to have your custon X Yes  No	ners conduct au	dits on your s	ite?		
2.17	Please list regulatory sanctions impa warning letters, CEP suspension, im N/A			five years (i.e.		
2.18	Does the site outsource any quality-	related activity	?			
	☐ Yes ☐ No ☐	N/A				
	If answering yes, please specify the	activities:				
2.19	Please check the supplier controls in	place for this f	facility:			
2.19a	Quality Agreements with Suppliers	⊠ Yes	☐ No	□ N/A		
2.19b	Subcontractor Qualification/Audit Program	⊠ Yes	☐ No	□ N/A		

	<b>SECTION 2. General</b>	Site Opera	ting Info	ormatio	n
2.19c	Periodic Review of Supplier Performance	⊠ Yes		No	□ N/A
2.19d	Supplier Feedback Program	Yes		□ N/A	
2.19e	Approved Material Supplier List	∑ Yes		N/A	
2.19f	Approved Service Supplier List	⊠ Yes		No	□ N/A
Addit	ional comments:				
Suppl	iers are subcontractor for transportat	ion			
	SECTION 3. Object		<u>aterials (</u>	on Site	
3.1	Does the site or production plant p				
	process or store any of the following	ng:	<b>T</b> 7	<b>3</b> .7	Not
			Yes	No	Applicable
3.1a	Beta-Lactam Antibiotics				
3.1b	Steroids and/or hormones				
3.1c	High potency compounds				
3.1d	Materials of animal origin/Biolog	ics			
3.1e	Live virus or micro-organism				
3.1f	Allergens				
3.1g	Genetically Modified Organisms (	(GMO)			
3.1h	Agrochemicals (Pesticides, Herbid	cides,			$\boxtimes$
	Fungicides, etc.)				
3.1i	Other (Please specify):				
	storage only				
	SECTION 4. Cross	Contamin	nation Co	ontrol	
4.1	Are any of the following cross-		Yes	No	Not
	contamination controls in place	?	168	110	Applicable
4.1a	Dedicated Facilities				

Additional Comments:

Access Controls

Dedicated Personnel

Dedicated Gowning

Procedural Controls
Other (please specify):

4.1b

4.1c

4.1d

4.1e

4.1f

Site Specific:		<b>SECTION 5. Site Operating Policies</b>							
Site Specific:   Site			Vas	No					
Site Specific:   5.1a	5.1	Does the site utilize the following written		110	Аррисавіе				
Site Specific:	3.1	=							
S.1a   Environmental, Health, and Safety   S.1b   Facility Environmental Control Policy   S.1c   General Facility Cleaning Procedures   S.1c   Hygiene and Sterilization Procedures   S.1d   Validated Equipment Cleaning Procedures   S.1e   Preventative Maintenance Program/Procedures   S.1f   Pest Control Program   S.1g   Master Production Procedure   Master Validation Plan   S.1j   Periodic Product Quality Review   S.1k   Master Validation Plan   Supplier Approval Procedure   Supplier Approval Procedure   S.1n   Monitoring and Review of Approved Suppliers   S.1n   Monitoring and Review of Approved Suppliers   S.1o   Mechanism to Reduce Testing   Master Validation Procedure   S.1d   Change Control Procedures   S.1r   Document Management Policy   S.1s   Document Retention Policy   S.1s   Sampling Procedure   S.1w   Out of Specification Procedure   S.1w   Deviation/Investigation Procedure   S.1w   Sampling Procedure   S.1s   Sampling Procedure   S.1s   Sampling Procedure   S.1s   Raw Material Retention Program   S.1c   CAPA Procedure   S.1s   CAPA Procedure   S.1s   CAPA Procedure   S.1s   Sampling Procedure   S.1s   Capa Product Release Procedure   S.1c   Employee Training Program   S.1cd   Stability, Expiration, and Shelf-Life Program   S.1dd   Stability, Expiration, and Shelf-Life Program   S.1ff   Recall Procedure	Sita Sn								
S.1b   Facility Environmental Control Policy   S.1c   General Facility Cleaning Procedures   S.1d   Validated Equipment Cleaning Procedures   S.1d   Validated Equipment Cleaning Procedures   S.1e   Preventative Maintenance Program/Procedures   S.1f   Pest Control Program   S.1g   Master Production Procedure   S.1g   Master Production Procedure   S.1h   Quality Control/Quality Management Policy   S.1i   Quality Manual   S.1j   Periodic Product Quality Review   S.1k   Master Validation Plan   S.1l   Risk Assessment Program   S.1l   Risk Assessment Program   S.1m   Supplier Approval Procedure   S.1n   Monitoring and Review of Approved Suppliers   S.1o   Mechanism to Reduce Testing   S.1p   Receiving Incoming Inspection   S.1q   Change Control Procedures   S.1r   Document Management Policy   S.1s   Document Management Policy   S.1s   Document Retention Policy   S.1t   Change Notification Procedure   S.1v   Deviation/Investigation Procedure   S.1v   Deviation/Investigation Procedure   S.1v   Deviation/Investigation Procedure   S.1v   Sampling Procedure/Sampling Plan   S.1v   Raw Material Retention Program   S.1v   CAPA Procedure   S.1a   Label Control and Accountability   S.1bb   Product Release Procedure   S.1cc   Employee Training Program   S.1dd   Stability, Expiration, and Shelf-Life Program   S.1ec   Product Retention Program									
S.1c   General Facility Cleaning Procedures   S.1c   Hygiene and Sterilization Procedures   S.1d   Validated Equipment Cleaning Procedures   S.1e   Preventative Maintenance Program/Procedures   S.1e   Preventative Maintenance Program/Procedures   S.1f   Pest Control Program   S.1g   Master Production Procedure   S.1g   Master Production Procedure   S.1h   Quality Control/Quality Management Policy   S.1i   Quality Manual   S.1j   Periodic Product Quality Review   S.1k   Master Validation Plan   S.1l   Risk Assessment Program   S.1m   Supplier Approval Procedure   S.1n   Monitoring and Review of Approved Suppliers   S.1o   Mechanism to Reduce Testing   S.1o   Mechanism to Reduce Testing   S.1q   Change Control Procedures   S.1r   Document Management Policy   S.1s   Document Retention Policy   S.1t   Change Notification Procedures   S.1t   Change Notification Procedure   S.1t   Control of Nonconforming Material   S.1v   Deviation/Investigation Procedure   S.1x   Sampling Procedure/Sampling Plan   S.1y   Raw Material Retention Program   S.1y   Raw Material Retention Program   S.1z   CAPA Procedure   S.1aa   Label Control and Accountability   S.1bb   Product Release Procedure   S.1dd   Stability, Expiration, and Shelf-Life Program   S.1dd   Stability, Expiration, and Shelf-Life Program   S.1ec   Product Retention Program   S.1ec   Product Rete									
S.1c									
S.1d									
S.1e   Preventative Maintenance Program/Procedures   S.1f   Pest Control Program   S.1g   Master Production Procedure   S.1g   Master Production Procedure   S.1h   Quality Control/Quality Management Policy   S.1i   Quality Manual   S.1i   Periodic Product Quality Review   S.1k   Master Validation Plan   S.1l   Risk Assessment Program   S.1m   Supplier Approval Procedure   S.1n   Monitoring and Review of Approved Suppliers   S.1o   Mechanism to Reduce Testing   S.1p   Receiving Incoming Inspection   S.1p   Receiving Incoming Inspection   S.1q   Change Control Procedures   S.1r   Document Management Policy   S.1s   Document Retention Policy   S.1t   Change Notification Procedures   S.1v   Deviation/Investigation Procedure   S.1v   Deviation/Investigation Procedure   S.1v   Deviation/Investigation Procedure   S.1x   Sampling Procedure/Sampling Plan   S.1y   Raw Material Retention Program   S.1z   CAPA Procedure   S.1z   CAPA Procedure   S.1z   Capa   Capa			<u> </u>						
S.1f   Pest Control Program				$\Box$					
S.1g   Master Production Procedure									
Quality:         5.1h       Quality Manual       □       □       □       □       5.1i       Quality Manual       □			$\boxtimes$						
5.1h   Quality Control/Quality Management Policy   S.1i   Quality Manual   S.1j   Periodic Product Quality Review   S.1k   Master Validation Plan   S.1l   Risk Assessment Program   S.1m   Supplier Approval Procedure   S.1n   Monitoring and Review of Approved Suppliers   S.1o   Mechanism to Reduce Testing   S.1p   Receiving Incoming Inspection   S.1q   Change Control Procedures   S.1r   Document Management Policy   S.1s   Document Management Policy   S.1t   Change Notification Procedures   S.1t   Change Notification Procedures   S.1u   Control of Nonconforming Material   S.1v   Deviation/Investigation Procedure   S.1w   Out of Specification Policy and Procedure   S.1x   Sampling Procedure/Sampling Plan   S.1z   CAPA Procedure   S.1a   Label Control and Accountability   S.1a   Label Control and Accountability   S.1a   Label Control and Accountability   S.1bb   Product Release Procedure   S.1cc   Employee Training Program   S.1cc   Employee Training Program   S.1cc   Product Retention Program   Product Retention Program   Product Retention Program   Product R	5.1g	Master Production Procedure							
S.1i   Quality Manual   S.1j   Periodic Product Quality Review   S.1k   Master Validation Plan   S.1l   Risk Assessment Program   S.1l   Risk Assessment Program   S.1m   Supplier Approval Procedure   S.1n   Monitoring and Review of Approved Suppliers   S.1o   Mechanism to Reduce Testing   S.1p   Receiving Incoming Inspection   S.1q   Change Control Procedures   S.1r   Document Management Policy   S.1s   Document Management Policy   S.1t   Change Notification Procedures   S.1u   Control of Nonconforming Material   S.1u   Control of Nonconforming Material   S.1v   Deviation/Investigation Procedure   S.1v   Deviation/Investigation Procedure   S.1x   Sampling Procedure/Sampling Plan   S.1y   Raw Material Retention Program   S.1z   CAPA Procedure   S.1a   Label Control and Accountability   S.1b   Product Release Procedure   S.1cc   Employee Training Program   S.1cc   Employee Training Program   S.1cc   Employee Training Program   S.1ce   Product Retention	Quality	<b>y:</b>							
5.1i   Quality Manual	5.1h	Quality Control/Quality Management Policy	$\square$						
S.1j	5.1i		$\overline{\boxtimes}$						
5.1k       Master Validation Plan         5.11       Risk Assessment Program         5.1m       Supplier Approval Procedure         5.1n       Monitoring and Review of Approved Suppliers         5.1o       Mechanism to Reduce Testing         5.1p       Receiving Incoming Inspection         5.1q       Change Control Procedures         5.1r       Document Management Policy         5.1s       Document Retention Policy         5.1t       Change Notification Procedures for Clients         5.1u       Control of Nonconforming Material         5.1v       Deviation/Investigation Procedure         5.1w       Out of Specification Policy and Procedure         5.1x       Sampling Procedure/Sampling Plan         5.1y       Raw Material Retention Program         5.1z       CAPA Procedure         5.1aa       Label Control and Accountability         5.1bb       Product Release Procedure         5.1cc       Employee Training Program         5.1dd       Stability, Expiration, and Shelf-Life Program         5.1ff       Recall Procedure									
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5.1p       Receiving Incoming Inspection         5.1q       Change Control Procedures         5.1r       Document Management Policy         5.1s       Document Retention Policy         5.1t       Change Notification Procedures for Clients         5.1u       Control of Nonconforming Material         5.1v       Deviation/Investigation Procedure         5.1w       Out of Specification Policy and Procedure         5.1x       Sampling Procedure/Sampling Plan         5.1y       Raw Material Retention Program         5.1z       CAPA Procedure         5.1aa       Label Control and Accountability         5.1bb       Product Release Procedure         5.1cc       Employee Training Program         5.1dd       Stability, Expiration, and Shelf-Life Program         5.1ee       Product Retention Program         5.1ff       Recall Procedure									
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5.1u       Control of Nonconforming Material       Image: Control of Nonconforming Material Procedure       Image: Control of Nonconforming Material Procedure       Image: Control of Nonconforming Material Procedure       Image: Control of Nonconforming Program       Image: Control of									
5.1v Deviation/Investigation Procedure   5.1w Out of Specification Policy and Procedure   5.1x Sampling Procedure/Sampling Plan   5.1y Raw Material Retention Program   5.1z CAPA Procedure   5.1aa Label Control and Accountability   5.1bb Product Release Procedure   5.1cc Employee Training Program   5.1dd Stability, Expiration, and Shelf-Life Program   5.1ee Product Retention Program   5.1ff Recall Procedure									
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5.1x       Sampling Procedure/Sampling Plan       Image: Comparison of the procedure of the p									
5.1y       Raw Material Retention Program       Image: Capa of the program of the product Release Procedure of the product Release Procedure of the product Retention Program of the product									
5.1z       CAPA Procedure       Image: CAPA Procedure of the control and Accountability of th				$\dashv \vdash \vdash \vdash$					
5.1aa       Label Control and Accountability		Č							
5.1bb     Product Release Procedure       5.1cc     Employee Training Program       5.1dd     Stability, Expiration, and Shelf-Life Program       5.1ee     Product Retention Program       5.1ff     Recall Procedure					+				
5.1cc       Employee Training Program       Image: Comparison of the program of the program of the program of the product Retention Program of the product Retention Program of the product Recall Procedure of the product of the program of the product Recall Procedure of the product Recall Procedure of the program of the program of the product Recall Procedure of the program of the product Recall Procedure of the program					<del>                                     </del>				
5.1dd     Stability, Expiration, and Shelf-Life Program     \( \) \( \)       5.1ee     Product Retention Program     \( \) \( \)       5.1ff     Recall Procedure     \( \) \( \)									
5.1ee Product Retention Program   5.1ff Recall Procedure					+ 💆				
5.1ff Recall Procedure				$+ \vdash$					
CALGG LEUSTOMER COMPISINT HANGLING LYT LITE LY LITE	5.1gg	Customer Complaint Handling		<del>                                     </del>					

Equipment validation/qualification procedure					
Internal audit/self-inspection program					
procedure					
Site Security/Site Access Control Policies	$\boxtimes$				
New Hire Program/Induction Program					
Continuity/Contingency Plan:					
Disaster Recovery Plan					
Pandemic Preparedness Plan					
Supply Chain Emergency Preparedness Plan	$\boxtimes$				
Business Continuity/Contingency Plan	$\boxtimes$				
Can the company provide a plan upon request? C	R provide	a short o	description		
below:					
5.1mm the company has an organisation in place	that hand	les specif	ically the		
pandemic topics; measures are put in place to avoid any contamination on site					
and to perenise the business. A referent person ha	as been ide	entified to	ensure the		
safety on site but also to follow upon governmental directives and confinement/					
isolation rules/sanitary pass requirements.					
5.1.u non conforming relates to product appearance and not to quality testing					
5.1x control of containers after reception, not testing related					
	Internal audit/self-inspection program procedure  Site Security/Site Access Control Policies New Hire Program/Induction Program  Continuity/Contingency Plan:  Disaster Recovery Plan  Pandemic Preparedness Plan  Supply Chain Emergency Preparedness Plan  Business Continuity/Contingency Plan  Can the company provide a plan upon request? Continuity below:  5.1mm the company has an organisation in place pandemic topics; measures are put in place to avoid and to perenise the business. A referent person has safety on site but also to follow upon government isolation rules/sanitary pass requirements.  5.1.u non conforming relates to product appearance.	Internal audit/self-inspection program procedure  Site Security/Site Access Control Policies  New Hire Program/Induction Program  Continuity/Contingency Plan:  Disaster Recovery Plan  Pandemic Preparedness Plan  Supply Chain Emergency Preparedness Plan  Business Continuity/Contingency Plan  Can the company provide a plan upon request? OR provide below:  5.1mm the company has an organisation in place that handle pandemic topics; measures are put in place to avoid any contained and to perenise the business. A referent person has been ideas afety on site but also to follow upon governmental directive isolation rules/sanitary pass requirements.  5.1.u non conforming relates to product appearance and not	Internal audit/self-inspection program procedure  Site Security/Site Access Control Policies  New Hire Program/Induction Program  Continuity/Contingency Plan:  Disaster Recovery Plan  Pandemic Preparedness Plan  Supply Chain Emergency Preparedness Plan  Business Continuity/Contingency Plan  Can the company provide a plan upon request? OR provide a short obelow:  5.1 mm the company has an organisation in place that handles specific pandemic topics; measures are put in place to avoid any contamination and to perenise the business. A referent person has been identified to safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon governmental directives and contamination is safety on site but also to follow upon gov		

	SECTION 6. Quality Assurance and Production				
		Yes	No	Not Applicable	
6.1	Does the site have an independent and defined Quality Assurance/Quality Management Division?				
6.2	Does QA/QM have authority over the following:				
6.2a	Policies and procedures?				
6.2b	Review of documentation for release?				
6.2c	Release or rejection of incoming materials?				
6.3	Does QA/QM investigate and resolve quality complaints?	$\boxtimes$			
6.4	Does QA/QM investigate and resolve internal deviations?				
6.5	Does the QA/QM have the authority to assign a disposition to materials?				
6.6	Does the QA/QM review manufacturing and testing records prior to release?				
6.7	Does the facility utilize computerized systems for managing GxP activities or data?				
6.8	Are relevant computerized systems 21 CFR part 11 and EU GMP annex 11 compliant?				
6.9	Does the site use statistical methods for consistency and uniformity?				

SECTION 6. Quality Assurance and Production						
		Yes	No	Not Applicable		
6.10	Does the site use controlled documents for following and recording manufacturing instructions?					
6.11	Does the company qualify and/or validate manufacturing procedures?			$\boxtimes$		
6.12	Is any environmental monitoring conducted in production/finishing areas?			$\boxtimes$		
6.13	Does the site supply BSE/TSE declarations?	П	П	$\square$		
6.14	Does the site supply a declaration of Elemental Impurities?	П	П	X		
6.15	Are ICH Q3C(R4) solvents used in the manufacturing process of supplied materials?					
6.16	Are stability studies carried out according to ICH guidance?	П		$\boxtimes$		
6.17	Are solvents and mother liquor reused/recycled?	П	П			
6.18	Does the site have a process water treatment system?	П	П			
	Distilled water Dionized water Water for injection (WFI) Reverse Osmosis Clean steam Ultra-filtrated water (purified water) Other:					
6.19	Does the plant have a batch/lot system?					
6.19a	Is the system traceable?	$\boxtimes$				
6.19b	Is it unique?			$\boxtimes$		
6.19c	Is batch/lot manufacturing continuous?			$\boxtimes$		
6.19d	Is manufacturing batch by batch?					
6.20	Does the site perform on-plant audits prior to approving critical GxP suppliers?			$\boxtimes$		
6.21	Does the site audit critical GxP suppliers after initial approval?			$\boxtimes$		
6.22	Does the site inspect incoming materials?	$\boxtimes$				
6.23	Does the site test incoming materials to defined specifications?					
6.24	Does the site establish purchase specifications for raw materials?					
6.25	Is the equipment multi-use?			$\boxtimes$		
6.26	Does the site qualify equipment installation?					

		Yes	No	Not Applicable
6.27	Does the site qualify equipment operation?			
6.28	Does the site qualify equipment performance?			
6.29	Are production critical use instruments calibrated regularly?			
6.30	Is rework allowed?			
6.31	Is reprocessing allowed?			
6.32	Are manufacturing and packaging activities traceable to the equipment, areas, and materials used?			$\boxtimes$
6.33	Are production materials handled and stored in a manner to prevent degradation, contamination and cross-contamination?			$\boxtimes$
6.34	If answering 'not applicable' for any of the above, please elabo Any manufacturing related questions are not applicable.	rate:		
Additi	onal Comments:			

SECTION 7. Laboratory Procedures			⊠ N/A	for this Site
		Yes	No	Not Applicable
7.1	Does the site have standard procedures for sample handling/tracking?			
7.1a	Does the site have standard procedures for retaining samples?			
7.1b	Does the site have standard procedures for retesting samples?			
7.2	Does the site have written and approved specifications and test methods?			
7.3	Are laboratory instruments calibrated regularly?			
7.4	Is there a standard procedure in place for analytical method development?			
7.5	Does the site qualify and/or validate analytical test procedures?			
7.6	Does the site perform stability testing on materials and/or products?			
7.7	Are retention samples of key raw materials maintained?			
7.8	Are standards traceable to their preparation and reagents used?			
7.9	Are retention samples of finished product maintained?			
7.10	Are shelf life/retest/expiration dates available and standardized?			

	<b>SECTION 7. Laboratory Procedures</b>		$\times$ N/A	for this Site
		Yes	No	Not Applicable
7.11	Does the company provide a certificate of analysis (CoA) and/or a Certificate of Conformation/Compliance (CoC) for each lot or batch?			
7.12	Does the CoA/CoC contain the manufacture name and location?			
7.13	Does the CoA/CoC signed/e-signed by a Quality representative?			
7.14	If a repacker performs analyses, will the CoA reflect both the original manufacturing site data as well as the repacking site data?			
7.15	If answering 'not applicable' for any of the above,	please elab	orate:	
7.16	Additional Comments:			
SECTION 8. Packaging, Storage, and Transport		sport	$\square$ N/A	A for this Site
		Yes	No	Not Applicable
8.1	Does the site have a validated or qualified labeling system?			
8.2	Are batch production records retained and available?			
8.3	Are packaging and labeling areas separate from production?			
8.4	Are barcode readers in use and challenged regularly?			
8.5	Are vision systems in use?			
8.6	Is product ever packaged without a label being initially applied (i.e. bright stocking)?			
8.7	Do labels include shelf life/expiration dates?			
8.8	Do labels include lot/batch number?			
8.9	Do labels include requirements for storage conditions?			
8.10	Is tamper evident seal used for each container of supplied materials?			
8.11	Does the company use a First-In-First-Out or First-Expiration-First-Out system?	$\boxtimes$		
8.12	Does the company maintain appropriate storage conditions?	$\boxtimes$		
8.12a	Are those storage conditions monitored and documented?	$\boxtimes$		

SECTION 8. Packaging, Storage, and Transport			☐ N/A for this Site	
		Yes	No	Not Applicable
8.13	Does the site make available a description of storage and/or warehouse conditions?	$\boxtimes$		
8.14	Does the site distribute products via a third party?	$\boxtimes$		
8.15	Are good distribution policies implemented?	$\boxtimes$		
8.16	Are transport mechanisms dedicated?	$\boxtimes$		
8.17	Does the company validate shipping method?			
8.18	Does the company validate packaging methods?			
Additi	onal Comments:			
Shippi	ng validation is not required as site currently handles re	esearch u	se produ	ets only, but it
is plan	ned for handling of gmp-related products		-	-
	e does not have labeling or packing operations. Only d	istributio	n.	

I (Supplier) confirm that the information provided in this questionnaire is correct and can be verified.

Date:20.January 2022

Title:Site Quality manager