

Interpretation of GMP Annex 1 2022 (Rev. 1)

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1 Purpose and scope

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This technical interpretation focuses on some of the most important main changes of the revision 2022 of Annex 1 and also covers aspects that were already included in the previous version of this guideline and that repeatedly gave rise to questions. This technical interpretation is intended to reflect the general opinion of the Swiss Inspectorates on these topics and to serve as a support during the inspection of manufacturers of sterile medicinal products.

2 Basics

The Revised Annex 1 to the PIC/S GMP Guide (PE 009), about manufacture of sterile medicinal products, adopted on 9 September 2022 by the PIC/S Committee and came into force on 25 August 2023 (with the exception of point 8.123, which will become binding from 25 August 2024).

3 Definitions and abbreviations

n.a.

4 Interpretation: Questions and Answers

4.1 Scope (Annex 1, Chapter 1)

Q&A	Paragraph	Questions	Answers
No.	No.		
1	Chapter 1	Does Annex 1 apply without re-	It is recognised that ATMPs cover a very hetero-
	(Scope)	striction also to ATMPs or are	geneous range of products and that for some of
		only defined aspects of Annex 1	these products, due to their nature and manufac-
		to be followed for some specific	turing technology, specific considerations are re-
		product types, such as for exam-	quired. This certainly applies, for example, to the
		ple the allogeneic and autolo-	allogeneic and autologous cell therapy pro-ducts,
		gous cell therapy products?	which are to be manufactured under conditions
			suitable to avoid microbial contaminations, but
			which usually cannot be terminally sterilised or
			sterile filtered. In addition, such products are
			made from unsterile patient material. Specifically
			with regard to cellular therapy, Annex 2A, para-
			graph 5.29(b), requires that aseptic processing
			be maintained from the time of procurement of
			cells through manufacturing and administration
			back into the patient. Annex 2A refers to Annex 1
			several times (e.g., in connection with the re-
			quirements for the provision of systems for
			closed processing), but implies the possibility of
			exceptions from applying the requirements of An-
			nex 1. It must also be taken into account that An-
			nex 2A became valid in May 2021, i.e., more
			than one year before the publication of the new
			Annex 1 version.



	It is exp	pected that ATMP manufacturers, based
	on the	knowledge of their manufacturing pro-
	cesses	and the execution of detailed risk anal-
	yses co	overing all process steps, materials and
	system	s, develop and implement Contamination
	Control	Strategies suitable to avoid or largely
	minimis	se risks of product contaminations. Justifi-
	cation r	must be given for any exceptions to the re-
	quirem	ents of Annex 1.

4.2 Premises (Annex 1, Chapter 4)

Q&A No.	Paragraph No.	Questions	Answers
2	4.1, 4.11 & 4.12	Are grade A and B cleanrooms required to have separate air- locks for material and personnel and must the flows in such air- locks be strictly unidirectional?	In general, it is expected that new facilities have for grade A and B zones segregated airlocks for personnel and material and that the flows in such airlocks are unidirectional (i.e., separate MALs for transport into and out of the cleanroom and separate PALs for personnel entry and exit). Existing facilities that do not have such airlock separation must ensure that, as a minimum, tem- porary separation of the flows in the airlocks is guaranteed and that the situation is covered by scientifically sound risk analysis also assessing the need of additional technical or organizational measures. The rationale for not applying physi- cal separation of the above-mentioned flows through segregated airlocks and the risk assess- ment on which it is based must be integrated in the overall contamination control strategy.
3	4.12	Paragraph 4.12, point ii, states that only materials and equip- ment that are on an approved list and that have been assessed during validations of the transfer process, should be transferred into grade A or grade B areas via airlocks or pass-through hatches. What does "validation of the transfer process" mean, for example for the material transfer into an isolator?	As mentioned in paragraph 4.10 of Annex 1, the transfer of materials, equipment or components into and out of a cleanroom (incl. the critical zone within a grade A environment), represents one of the greatest potential sources and risks of contamination. In order to minimise such risks, great care must be taken in particular when defining the technical and procedural measures associated with the transfer of materials/equipment into an aseptic processing area. Only in relatively rare cases it is possible to bring materials into an isolator before it is sealed and bio-decontaminate them together with the isolator using a validated VHP treatment



			(only possible for small batches and if materials
			are resistant to VHP). In the majority of cases,
			however, it is necessary to transfer materials to
			an isolator that has already been decontami-
			nated. For this, all materials must first be steri-
			lised and then moved through the physical bar-
			rier of the isolator in such a way that the sterility
			of the goods and the integrity of the isolator are
			maintained. Regardless of the technology used
			(e.g., usage of double-door sterilisers upstream
			of the isolator, use of transfer isolators or of rapid
			transfer port technology), the entire transfer pro-
			cess must be considered within detailed risk
			analysis and be part of the overall contamination
			control strategy. In addition, appropriate control
			mechanisms must be defined to monitor the
			maintenance of the integrity and functionality of
			the systems (e.g., measurement of differential
			pressure and control of door interlocks between
			adjacent zones of the transfer system). The tech-
			nical solutions must be covered by appropriate
			equipment/system qualifications (incl. smoke
			studies if applicable) and sterilisation validations
			and the suitability of the entire transfer process
			must be verified through APS (validations and
			also regular APS). Appropriate qualification
			measures and APS must also be used to
			demonstrate that the egress of materials from
			the isolator does not affect the maintenance of
			the grade A zone requirements.
			The arrangement of the installations, the pro-
			cesses carried out in it and the material move-
			ments must also be considered when defining
			the points to be sampled during PQ activities or
			during routine or event based environmental
4	4 1 2	le it always required to strictly	monitoring.
4	4.12	Is it always required to strictly adhere to the area cleanliness	Compliance with the cleanroom sequence for the
			transfer of materials via airlocks or pass-through
		cascade (i.e., respecting the se-	hatches is expected to be fulfilled for zones A
		quential order of cleanroom clas-	and B (exceptions from this rule are possible for
		ses) for material transfer through	sterility test rooms). For cleanroom areas with
		airlocks or pass-through hatches	lower classification, it is principally feasible for
		or is it possible to skip a grade	materials to be transferred from one low zone
		(e.g., moving from CNC directly	(CNC) through an airlock or pass-through hatch



	-		
		to class C) under certain circum- stances?	directly into an area with two grades higher clas- sification (grade C area), provided that suitable technical and/or procedural measures are estab- lished ensuring fulfilment of the cleanroom speci- fications in the respective areas. The adequacy of the established systems/procedures needs to be demonstrated by appropriate qualification ac- tivities and the results of regular environmental monitoring. The defined measures and the risk analyses on which they are based must be part of the CCS.
5	4.20	What are the expectations for older barrier technology systems that do not meet all the require- ments according to the new An- nex 1? By when do they have to be replaced or upgraded?	The company has to perform an in-depth internal evaluation of the current barrier technology and assess whether the installation, its cleanroom background and all related systems/procedures meet the requirements of the new Annex 1 or whether technical measures are required. If nec- essary, a project has to be initiated for example to upgrade the cleanroom used as background and install additional airlocks. From August 25 th 2023, all barrier technology equipment not com- plying with the revised Annex 1 are considered deficient and deviations will be issued upon find- ings during inspections. Depending on the CAPA plan and interim risk reducing measures defined, an additional implementation timeline of approx. one year may be acceptable. This refers to the inclusion in such CCS risk as-
0	4.20	by the need to take into consid- eration, among others, the "ex- tent of automation" when carry- ing out CCS related risk assess- ments of an isolator?	sessments of an evaluation of all automated functionalities and processes associated with the use of the isolator and the activities taking place in it (from cleaning and disinfection of the equip- ment, to the transport of materials into the isola- tor, their handling and the product filling, to the capping and removal of the filled containers). The use of a well-designed, automated, recipe- controlled and possibly robotised system, equipped with appropriate control and alarm sys- tems, can increase the reproducibility of the op- erations and minimise both errors and manual in- terventions. Ideally, such risk analyses should al- ready be carried out as part of the design or se- lection of the isolator system and should be re- vised or supplemented during the lifecycle of the

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	I		aquipment on knowledge and everying in a
			equipment, as knowledge and experience in-
			creases or in the event of changes to be imple-
			mented.
7	4.22	Are manual operations accepted	The decontamination process for an isolator
		for bio-decontamination?	should always be an automatic process. How-
			ever, paragraph 4.22 refers to both RABS and
			isolators, and manual decontamination pro-
			cesses are most commonly encountered for
			RABS. Such manual processes must be de-
			signed in such a way to be reproducible and to
			cover the entire surface area of the equipment,
			and their robustness and effectiveness must be
			demonstrated by appropriate validation and by
			regular monitoring.
8	4.22	"Evidence should also be availa-	The cleaning or bio-decontamination procedure
U	1.22	ble to demonstrate that the	should include steps designed to effectively re-
		cleaning and bio-decontamina-	move cleaning agent or disinfectant (including
		tion agents used do not have ad-	sporicidal agent) residues from direct and indi-
		verse impact on the product pro-	rect product contact surfaces within the
		duced within the RABS or isola-	
		tor". How should this be demon-	RABS/isolator. The effectiveness of these steps should be demonstrated based on validation
		strated?	data. For isolators, validation data should be
			available to demonstrate that the residual
			amount of sporicidal agent is below the concen-
			tration that could be detrimental to the product
			quality and stability at the end of the sporicidal
			cycle. Control mechanisms must be defined to
			ensure that the conditions prevailing during vali-
			dation are also maintained during routine pro-
			duction (e.g., compliance with the time after
			completion of the sporicidal cycle, resp. meas-
			urement of the peroxide concentration).
			The extent to which samples of the surfaces are
			to be taken during validation and analysed for
			disinfectant/cleaning agent residues can be de-
			fined based on a risk assessment, taking into ac-
			count the risk of transferring such residues to the
			product or a product contacting surface of a
			packaging component.
9	4.30	Is it acceptable that for barrier	The most important requirement for barrier tech-
	4.00	technology systems with unidi-	nology systems stated in paragraph 4.30 is that
		rectional air flow other air speed	the air velocity in unidirectional airflow systems
			-
		and speed measurement posi-	must be defined in such a way that unidirectional
		tions are defined than those	and uniform airflow conditions prevail at the
		mentioned in Annex 1?	working positions where high-risk operations

take place, suitable to protect the product and
open components (e.g., containers) from con-
tamination.
The air speed range of 0.36 - 0.54 m/s is, as
stated in the above paragraph itself, merely a
guideline value that has been encountered in the
pharmaceutical industry for decades.
Annex 1, however, clearly allows for the estab-
lishment of alternative air speed ranges or meas-
urements at different heights in the system than
the working position, provided this is "scientifi-
cally justified in the CCS". It is important that the
suitability of the defined airflow conditions is
proven by airflow visualisation studies (part of
the system qualification) covering the entire sys-
tem and that these are correlated with the re-
spective defined air speed range at specified
height/position. The air speed must be measured
continuously during operations and kept within
this defined range.

4.3 Utilities (Annex 1, Chapter 6)

Q&A No.	Paragraph No.	Questions	Answers
10	6.12	Water generation - reverse os- mosis system: What are the re- quirements regarding the saniti- zation (disinfection) of the sys- tem?	Paragraph 6.12 gives detailed guidance on the requirements. It is important that the system is designed to allow for routine sanitization / disinfection and a procedure is in place defining this regular preventive sanitization or disinfection of the RO-system. It should also include a regular change of membranes. The frequency of sanitization should be determined based on quality risk management principles and on the data gathered during the qualification of the system, and it should be reviewed at least annually taking into consideration the routine monitoring data. The system must continuously be maintained meaning that the sanitization also has be performed when no production is running or when no water is used for production.
11	6.13	What are the sampling require- ments for regular ongoing moni- toring of Water for Injection?	A suitable sampling schedule should be in place to ensure that representative water samples are obtained for analysis on a regular basis. For



	WFI distribution system sampling plans are
	more important because microbial control must
	be much more stringent. In general, water sam-
	pling for microbial and bacterial endotoxin test-
	ing is expected to occur daily somewhere in the
	system, with each outlet being sampled periodi-
	cally to characterize the quality of the water.
	The use of cold loops requires a much closer
	microbiological monitoring and special sanitiza-
	tion measures.
	Quality control sampling locations in the main
	distribution system should include all POUs,
	having also process control sampling be located
	before the first and after the last POU and at
	other specified worst-case locations. POU sam-
	pling plans should rotate through all use points
	on the system, with the expectation that sam-
	ples are collected on a daily basis from various
	use points, and that all use points are sampled
	on a rotational basis. The loop return should be
	sampled each day of use of the system in order
	to provide additional assurance of the quality of
	water utilized in the manufacturing processes.
	For WFI, it is an expectation that water samples
	should be taken daily from a minimum of one
	POU, with all point of use tested weekly during
	the qualification phase. The final phase of quali-
	fication may form the basis for the ongoing sam-
	pling frequencies with the goal of ensuring that
	the system is maintained in a validated state.
	However, it has become good industry practice
	to continue to utilize the same sampling fre-
	quency beyond the completion of the perfor-
	mance qualification to collect sufficient historical
	data in order to justify adjusting the sampling
	frequency. The use of risk analysis tools cou-
	pled with stringent periodic data review may be
	used to alter the frequency of sampling. Any de-
	crease of the sampling frequency for routine
	monitoring should be based on historical data
	and should only occur when a large number of
	historical data is available to allow statistical
	analysis. Based on the outcome of analysis of
	data and on the regular review of the perfor-
	mance of the point of use or the system, and if

			operational SOPs are in place which ensure also an increase of the sampling frequency if in- dicated and regular maintenance activities, e.g. for all outlets, a less frequent sampling can be justified. The risk assessment should consider the fact that decreased sampling frequencies also results in a higher number of batches that will be put at risk and a problem may have a se- rious impact on supply of products for patients.
12	6.19	Where should process gas be monitored?	The monitoring of process gas should be per- formed as close as possible before the steriliza- tion filter (the level of contamination before steri- lization should be under control to ensure the ef- ficiency of the gas sterilization process).

4.4 Personnel/Training (Annex 1, Chapter 7)

	Paragraph No.	Questions	Answers
13 7	7.4	This paragraph requires that all personnel accessing grade A and B areas be trained in asep- tic gowning and aseptic behav- iors. It also stipulates that com- pliance with the gowning proce- dure must be confirmed by means of assessments and peri- odic reassessments on an an- nual basis, covering both visual and microbiological checks (monitoring of gloved fingers, forearms, etc.). Are these assessments to be covered by staff participation in APS? Paragraphs 9.38 and 9.39 mention staff participation in APS only in the context of staff requalification. Do APSs also have to take place during the ini- tial qualification of employees? Does every employee have to perform every manual interven- tion in APS in order to be quali- fied or requalified?	To ensure product quality, adequate training of employees working in grade B and A areas or involved in aseptic processes (incl. the necessa- ry preparatory activities) is essential. The qualifi- cation must be adapted to the respective activi- ties of the single employee and, after initial train- ing (initial qualification), must also include regu- lar requalification / retraining. Annex 1 requires that each employee qualified and involved in aseptic processes participates in a successful APS at least annually (or every six months if the aseptic processes are manual) as part of his requalification. However, Annex 1 is not specific about the scope of the initial employee qualification, but in- dicates in paragraph 7.4 that the relevant train- ing must cover theory and knowledge transmis- sion as well as practical aspects and that evi- dence of training effectiveness is required through assessments using both visual and per- sonnel monitoring examinations. Although not explicitly required in Annex 1, the expectation for the initial qualification of an em- ployee for the aseptic area is that practical pro- cess simulations, including manual interven- tions, are carried out under the supervision of

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qualified trainers/QA and are followed by per-
sonnel monitoring as training verification.
It is at the discretion of the respective pharma-
ceutical company to define and justify whether
these process simulations need to be conducted
separately (but under similar conditions as an
APS) or can be integrated within an APS. It is
important that the representativeness of the ac-
tivities to be performed by the trainee for the ac-
tual processes is justified and that the simula-
tions cover each critical activity to be carried out
by the respective employee. Equivalent repre-
sentative interventions can be grouped for staff
qualification. All operators should perform one
intervention per year from each group of equiva-
lent representative interventions.

4.5 Production and Specific Technologies (Annex 1, Chapter 8)

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Q&A No.	Paragraph No.	Questions	Answers
14	8.36 & 8.38 / 8.39	Sterilisation: what are the requi- red loading patterns for initial and periodic autoclave (re-) vali- dations?	Initially each loading pattern must be validated. Re-validation of each loading pattern must be done annually. If a suitable worst-case load (the same material, same loading pattern, same cycle) for re-valida- tion (backed up with data) can be identified, not every load of this material needs to be re-vali- dated. A theoretical reference load is not ac- ceptable, as 8.36 states that "each type of load"
15	8.63	Moist heat sterilisation: is it ex- pected that routine re-validation includes a temperature mapping for systems where steam in place is used for sterilization?	needs to be validated. Yes, routine (or periodic) validation should in- clude tests providing evidence that the positions used for temperature monitoring throughout the sterilization process are still representative of and correspond to the slowest to heat locations during sterilisation.
16	8.128	Is the sterility of the product-con- tacting surface of a closed sys- tem ensured if the system is opened in a cabinet with laminar airflow (LAF)?	Opening a sterile, closed system should be avoided whenever possible. In general, a closed system that needs to be opened should be re- turned to the sterile state by carrying out a vali- dated sterilization process (if required, preceded by cleaning).

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			If a sterilisation of the system after opening is not possible, the system's opening could be per- formed in a decontaminated isolator (provided that the introduction of the closed system to be opened or relevant parts of it into the isolator does not compromise the isolator's decontami- nation status and can be considered covered by successful APS). Opening the system in an LAF with classification A and background B could possibly be an alter- native to the isolator but rather to be envisaged in exceptional cases only as the risks of intro- ducing contamination from the environment are higher and require appropriate consideration
17	8.128	Are non-aseptic connections al- lowed to be carried out for cou- pling closed systems if a sterili- zation cycle (SIP) occurs prior to use?	and risk mitigating measures. Yes, such an approach is possible, provided that the SIP process used is appropriately validated.
18	8.128	Is the use of sterile aseptic con- nectors purchased from qualified suppliers permitted as suitable strategy to connect sterile equip- ment to each other and may the end-user rely on the sterility doc- umentation (sterilisation valida- tion) provided by the respective supplier?	Provided the supplier of the aseptic connector in question was covered by comprehensive qualifi- cation activities and the validation package/data provided by the supplier for the connector (e.g., validation of the gamma irradiation process, data on microbial challenge tests, etc.) have been checked and found to be sound, the end-user can rely on such data, but must cover this equip- ment and its handling during his manufacturing process APS activities. See also paragraph 4.6.4 on single-use systems. If single-use connectors that are sterilised by the end-user (e.g., by autoclaving) are used for cou- pling sterile systems, this sterilisation process must be validated. It must also be ensured that single-use connectors are suitable for sterilisa- tion and that the latter does not impact their functionality or integrity (e.g., by causing the ma- terial of construction to become more porous).

19	8.128	Is it considered acceptable to in- troduce small amounts of prod- uct or cells into a sterile closed	Such a practice should be avoided for aseptic steps, as piercing a septum with a needle is to be regarded as a breach of the sterile barrier. In
		system with a syringe fitted with a needle through a septum? Can the system be considered closed after piercing of the sep- tum?	aseptic processes where the above approach is used, measures must be taken to re-design and optimise the procedure accordingly. If the process concerned cannot be improved and adapted immediately, consideration should be given, as a temporary measure, to minimise the risk of contamination, as to whether the sy- ringe should be left with the needle inserted in the septum after completion of the material addi- tion and appropriately secured in this position. In addition, it should be considered that the top of the septum (prior to piercing) be protected by a sterile film, which is removed just prior to inser- tion of the syringe needle to reduce the risk of contaminants on the septum surface entering the process and avoiding treating the septum with a disinfectant, which may also pose a risk of contamination of the product (by disinfectant
20	8.128	Is tubing welding considered a suitable strategy for aseptically connecting equipment parts maintaining the closed status of a system?	residues). Welding equipment and processes must be qualified/validated. If such processes are used in sterile or aseptic filling processes, they must be covered also by APS. However, as tubing welding processes are both less monitorable and entail risks of undetected integrity deficiencies, such practices should be avoided and more reliable systems should be used, which should be taken into account when- ever possible already during facility and process design.

21	8.129	Closed systems: in case of us- ing single-use systems, can sys- tem integrity tests performed by the respective suppliers be lev- eraged without having to carry out own tests?	Whenever possible, the integrity of critical sin- gle-use systems should be tested by the end- user on site (i.e., before use in production). It is acknowledged that such an integrity test, e.g., by means of a pressure hold test using an inert gas, is difficult to establish for small single-use bags/containers and is also only reliable to a lim- ited extent. However, the decision in this respect must be justified by well-founded measures and considerations, be verified by risk assessments and must be included in the CCS. The possibility of relying for single-use materials (such as bags) on integrity test results provided by the respective suppliers requires a detailed assessment of the situation, taking into account, among other things, the criticality of possible in- tegrity deficiencies on the manufacturing pro- cesses/product quality and their detection proba- bility during the process. The adoption of integrity results from the vendor requires an in-depth qualification of the supplier and must also take into account the risks of sub- sequent damage to the single-use material dur- ing its delivery and installation in production.
22	8.134	Single-use systems: what are the expectations placed on the assessment of such suppliers and what must it comprise?	The supplier assessment should be understood as a comprehensive qualification of the single- use systems (SUS) supplier. This assess- ment/qualification should cover not only the sup- plier delivering the SUS but in particular the SUS manufacturer (or each relevant manufacturing site, if the SUS in question is produced at sev- eral sites) as well as any sub-contractors in- volved in critical services or processes (e.g., sterilisation of the SUS). The supplier assess- ment/qualification should be carried out in paral- lel with the evaluation of the SUS material and should play a crucial role in the SUS selection decision. For all SUS that the end-user intends to use in his manufacturing process and that will have di- rect contact with the product, intermediates, pro- cess solutions or starting materials/raw materi- als, a Quality Agreement should be concluded with the respective supplier. This Quality Agree- ment should cover the SUS specifications as



			well as quality relevant service conditions (e.g., requirement to manufacture SUS in cleanrooms) and regulate, among other things, the terms rel- evant for the notification of planned changes and their approval by customers, the procedures in the event of major/critical deviations impacting delivered SUS, the terms in case of customer complaints and the oversight responsibility for sub-contractors. Supplier qualifications must include an assess- ment of the supplier's quality systems, a com- prehensive review of all relevant technical docu- mentation received (incl. for example drawings, documentation of components used such as fil- ters, aseptic connectors, tubings etc., certificates and validation/study packages), and audits. It is expected that audits cover all systems, relevant processes and control strategies (e.g., sterilisa- tion process and its validation, subcontractor qualification, etc.) considered critical for the re- spective SUS and these contents must be com- prehensibly documented in the respective audit
23	8.138	What aspects should be taken into account by the end-user when determining the acceptan- ce criteria of the respective SUS and in which form should they be specified?	report. Acceptance criteria should be defined taking into account the intended use of the particular SUS in the manufacturing process, the criticality of its use/impacted process, existing process knowledge, as well as available SUS experi- ence. Acceptance criteria should encompass quality aspects (e.g., sterility, biocompatibility, visible particles testing by compendial method, integrity tests, certificates, etc.), functionality (e.g., inserts and components required, temper- ature resistance in operating range, autoclaving or freezing resistance, chemical compatibility, substainable pressure, packaging requirements, etc.) as well as validation/qualification require- ments to be fulfilled by the SUS and its supplier. According to Annex 1, paragraph 8.132, the use of SUS and the associated risks should be also assessed as part of the Contamination Control Strategy, taking into account the fragile nature and potential complexity of the SUS in question, possible interactions of the SUS surfaces with



			the product, risks associated with manual opera-
			tions/connections and risks of holes or particle
			contaminations. The resulting conclusions from
			-
			these assessments and any risk mitigating
			measures should be taken into account, if ap-
			propriate, when establishing the SUS ac-
			ceptance criteria and the expectations placed on
			the SUS suppliers. To comply with the require-
			ments of paragraph 8.138, according to which
			the conformity of the SUS with the approved
			specification has to be checked upon good re-
			ceipt, the quality requirements should be defined
			in a written specification (including or referenc-
			ing a technical drawing of the material). The
			other expectations regarding the functionality of
			the SUS or the expected validation / qualifica-
			tion/study package to be made available by the
			supplier and to be agreed with the supplier, can
			be defined in another document, such as a SUS
			user requirement document, an annex to the
			Quality Agreement, or similar.
24	8.138 &	What should the incoming goods	Due to the special nature of SUS and their deliv-
	8.134	inspection at the end-user inclu-	ery in packaging that serves to protect them
		de to comply with paragraph	from damage but does not allow for a full visual
		8.138 & 8.134?	inspection of the materials, the scope of a feasi-
			ble routine incoming inspection program upon
			receipt is generally very limited.
			Immediately upon receipt, in accordance with
			paragraph 8.138, an initial documented inspec-
			tion of the shipment should be performed, con-
			sisting primarily of a review of the documents
			provided by the supplier, a visual check of the
			integrity of the outer packaging, label printing,
			and a reasonably cursory verification of the con-
			tents of the shipment (without complete unpack-
			ing of the SUS to avoid the risk of damages).
			On the basis of a positive result of this first in-
			coming check, the SUS can be released, allow-
			ing its transfer to the production area to be sub-
			jected to a more thorough examination. This ex-
			amination must consist at least of a deep visual
			inspection of the SUS by qualified employees
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			according to an established procedure, and the
			according to an established procedure, and the results of which must be documented as part of the batch record. The visual inspection should

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Interpretation GMP Annex 1

			include verification of the compliance of the SUS
			with approved technical drawings, the presence
			of gamma irradiation points (if applicable and as
			evidence of the sterilization), and a visual exami-
			nation for integrity (e.g., inspection of the welds,
			connectors, absence of critical scratches, etc.)
			as well as for the absence of particulates. If
			technically practicable and indicated, the integ-
			rity of the single-use system should be verified
	0.404		by means of a pressure hold test.
25	8.134	Is it considered acceptable that	When looking at the wording in the paragraphs
		some study or validation data	on single-use systems (SUS), it can be con-
		provided by SUS suppliers (e.g.,	cluded that Annex 1 allows principally the adop-
		validation packages incl. sterili-	tion by the end-user of data received from quali-
		zation or material chemical and	fied suppliers. However, this requires that the
		biological compatibility data) are	end-user confirms through a detailed review of
		incorporated by the end-user	the respective documentation that its contents
		into his own assessments with-	meet the user's standards and that the condi-
		out the need to carry full stud-	tions used by the suppliers when generating the
		ies/validations on his own?	data are representative (or worst-cases) for its
			own actual production conditions. The extent of
			the end-user's own studies/verifications or vali-
			dations activities depends on the representative-
			ness of the supplier data, the criticality of the in-
			tended SUS use in the process, established
			control strategies at the end-user which allow
			detection of possible SUS deficiencies (e.g.,
			pressure hold test, extensive program of micro-
			biological testing during the process), etc.
			For example, an end-user operating in the bio-
			tech area must verify with own studies or as-
			-
			sessments whether possible leachables emitted
			from the films of a single-use bioreactor can
			negatively affect the growth of cells and it is also
			the responsibility of the end-user to verify or pro-
			vide data to prove whether absorption effects of
			its product on the SUS surface are possible, re-
			sulting in an impact on product quality (see para-
			graph 8.132, « <i>These risks include but are not</i>
			limited to: i. the interaction between the product
			and product contact surface (such as adsorp-
			tion, or leachables and extractables)»).
			When SUS are used in the sterile production, it
			is mandatory that they are covered by APS, as
			required in paragraph 8.139. For cell culture (or

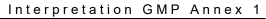


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		fermentative) processes, the end-user should
		evaluate the need to perform a process simula-
		tion before starting routine production to confirm
		the suitability, integrity, and handling of the SUS
		equipment.
26	Do end-users have to carry out	Most SUS suppliers provide comprehensive ex-
	their own extractable studies or	tractable studies packages to end-users. In re-
	can they use the supplier's data	cent years, efforts have been undertaken by in-
	and when is it necessary to exe-	ternational industry working groups to harmonize
	cute leachable studies?	and standardize the conditions for extractions
		and analysis of extractables. Additionally, many
		suppliers also provide certificates covering the
		product contact films, e.g., Biological Reactivity
		Test in Vivo per USP <88>, Class VI.
		It is expected that based on the package of ex-
		tractables data obtained, the end-users evaluate
		the adequacy of the data provided, potentially
		"add together" data from different SUS compo-
		nents, and define the need for additional ex-
		tractables studies to simulate process-specific
		worst-case conditions and perform health safety
		assessments, as appropriate.
		The decision to carry out leachable studies with
		the respective product is usually based on a
		comprehensive evaluation of the possible risks
		of administering leachables in doses that may
		be of concern when using the respective drug
		product. In addition to the results of the extracta-
		ble data and any resulting safety assessments,
		the decision regarding the need for a leachable
		study should take into account the route of ad-
		ministration of the respective drug (e.g., oral,
		parenteral, subcutaneous), the dosing frequency
		to a patient, the use of the respective SUS in the
		process (e.g., use in early or late manufacturing
		step) and the contact time of the process solu-
		tion/product with SUS surface.
		Based on an assessment of such aspects, the
		company may justify that no leachable studies
		are performed for products that are still in the
		clinical phase and/or are only administered infre-
		quently (e.g., vaccines). Since extractable/leach-
		able studies are part of the filing dossier for mar-
		keting authorization, it is necessary to align the
		strategy with the requirements of the marketing

			authorization or, if necessary, with the respec- tive regulatory authority.
27	8.80	Is it expected that there are two redundant sterilizing filtration steps in the process before aseptic filling?	Annex 1 encourages an additional filtration through a sterile sterilising grade filter, as close to the point of fill as possible. The installation of such a redundant sterile filter significantly re- duces the risk of a product quality impact in the event of a failed filter integrity test, which is why this risk-minimising measure is to be considered state of the art process and should be used es- pecially for new processes. The risks and impacts of filter integrity failures of pre-fill point sterile filters should be assessed as part of the CCS evaluations and the decision not to install a redundant sterilising filter be justified in the CCS.
			Even in the case of using two sterilising filters, any filter integrity failure that may have occurred should be investigated.
28	8.83 / 8.84	Paragraph 8.83 makes referen- ce to the relevant Pharmacopeia requirements in relation to the validation of sterile filtration of fluids. Which paragraph of the Pharmacopeia should be con- sidered for this purpose?	Relevant guidance can be found in Ph. Eur. 5.1.1, paragraph "Membrane filtration/Filtration effectiveness" and in guideline EMA/CHMP/CVMP/QWP/850374/2015. Useful details and expected data/methods can also be found in PDA TR26 "Sterilizing filtration of liquids".
29	8.91 / 8.92 / 8.93	Is a pre-use / post-sterilisation integrity testing ("PUPSIT") of sterilising grade filters used in aseptically processes manda- tory?	The expectation is that PUPSIT be applied to verify the integrity of the sterilized filter assem- bly. However, paragraph 8.87 allows some flexibility in justified cases supported by risk analysis and covered in the CCS.

4.6 Environmental & Process monitoring (Annex 1, Chapter 9)

Q&A No.	Paragraph No.	Questions	Answers
30	9.4	What is deemed regularly ("These risk assessments should be reviewed regularly")? How often does the risk assess- ment need to be reviewed?	It is not possible to give definitive guidance here, because, as ICH Q9 (R1) states, the fre- quency of Risk Review should be based on the level of risk. The frequency or timing of a Risk Review exercise may be based on the type and number of risks identified during an earlier Risk Assessment exercise, and on the extent of risk control that was required to mitigate risks. It



			may also depend on the level of uncertainty (i.e. lack of knowledge) that was present during an earlier risk assessment. The higher the level of uncertainty in relation to risk estimates and the related risk-based decisions, the greater the need to review those estimates and decisions at an early timepoint once such uncertainties have been reduced. An environmental monitoring trend report will be compiled every year and depending on the re- sults, the risk assessment might be reviewed. It should be assessed annually if the review of the risk assessment is required. Swissmedic recom- mends to review the risk assessment regularly, e.g. for a new plant it is recommended to reas- sess the RA after one year when more experi- ence and knowledge have been gained.
31	9.9	When do we expect more strin- gent action limits?	More stringent action limits might be necessary if the trend data shows very low levels of detec- tion of total particles and viable particles with no action limit excursions over a longer time period (e.g. one year).
32	9.10	What statistics do we expect for establishing alert levels?	For a new process where limited data and expe- rience for environmental monitoring data is available, it is acceptable e.g., to calculate the alert level limit based as 50% of the action level limit. When more data becomes available the alert level limits should be statistically from the environmental monitoring data to ensure that the alert setting takes into account its own re- cent historical behaviour. Traditionally the "2 or 3 standard deviation rule" (alert level = Average value + 2 x SD) which as- sumes the data is normally distributed, has been applied. As environmental monitoring data are usually not normally distributed, other statistical ap- proaches such as a nonparametric approach based on 99.9 Th or 99.99 th percentiles, a non- parametric tolerance limit approach, or a cut-off Value approach (e.g., at 59 th or 99 th percentiles) should be used. Alert level limits should be reviewed regularly by the company and be adapted, if necessary, based on the actual performance. Performance

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			limits should be considered as a confirmation of
		la Garage de la Activitation de la Collection de la Colle	
33	9.22 & 9.23 9.28 & 9.29 9.31	Is it possible to fully replace mi- crobiological monitoring using e.g. settle plates and volumetric air sampling systems, by other integrated sampling and testing systems (e.g. Rapid Microbio- logical Methods, RMM)	 based alert levels that are well below the action limits should be considered as a confirmation of solid microbial control of the environment. Paragraph 9.22 requires a microbial monitoring using a combination of methods such as settle plates, volumetric air sampling, glove, gown and surface sampling. New technologies are available as continuous active air sampling and rapid microbial testing system, e.g., based on digital imaging technology to detect and count growing microbes. Equivalence of methods should be demonstrated and the effectiveness of the chosen method should be proven, including for in-house germs. Validation data of these new methods should include recovery studies of the sampling method. The exposure time should not have any negative effect on the suitability of the media used. For the use of real-time viable particle counting and given the non-equivalency AFUs versus CFUs and current GMP / Pharmacopeial limits are in CFU, the company need to collect data on their process for the real-time viable particle counting to compare it to the standard environmental monitoring data. A full understanding of what triggers signals and
			what is a normal AFU signal in the process and the development of appropriate alert and action limits based on this data, together with appropri- ate procedures that should define the actions to be taken in response to alarms including the consideration of additional microbial monitoring must be established. A scientific justification for
			 the limits applied is required. Data must be available for at least 12 months. Required elements the company needs to have: Primary Validation Package of the vendor for the system User specific Validation Package / Data for
			 Verification of Validation Parallel phase grade A (RRM/traditional active air sampling method) in operation to gain experience with new technology under grade A



			 Test for interferences to be addressed, e.g., by disinfectants, materials or product Operational Approach: Implementation / Alarm Handling Concept Supportive data: Collection and evaluation of data in a production area grade C and D Identification of microorganisms must be per- formed as it is essential to determine the (possi- ble) root cause of a contamination and evaluate the risk of the contamination for the drug prod- uct. If an action limit exceedance occurs, the agar plate from the system must be incubated in order to isolate associated CFUs and to allow identification of species for further investigation and impact assessment of product quality.
34	9.34	What means "frequency" here? Absolute number of interven- tions or how often they occur during a certain time interval?	Frequency means that the absolute number of interventions that occur during the routine asep- tic process should be included in the APS.
35	9.34	What interventions should be included in the APS for the an- nual operator's requalification?	Each operator should perform each intervention. The worst case must be covered, which means that the interventions are independent of the lot size and the duration of the production. See in more detail in the Chapter above about Person- nel (Annex 1, paragraph 7.4).
36	9.36ii	What does "same container/clo- sure configuration" mean?	"Same container/closure configuration" refers to the dimensions, (e.g. diameter of opening) shape and material of the container and closure, like e.g. vial/stopper. E.g. has a stopper for ly- ophilisation a different form than a stopper for liquid products, this is therefore considered a different container/closure system.
37	9.36ii	There is a filling line with subse- quent lyophilisation. Can the liq- uid filling with subsequent ly- ophilization be considered as worst case, so that APS of liq- uid filling with lyophilization would cover as well liquid filling process without lyophilization?	No, liquid filling and lyophilisation are different processes on the same line, with different paths.
38	9.36ii	When can a bracketing or ma- trix approach be applied?	If equivalence can be shown between e.g. glassware and stoppers a bracketing approach can be applied for the APS. New materials must



			be validated and an APS is part of the valida- tion.
39	9.36xii & xiii	Does the APS of campaign manufacturing require the simu- lation of the maximal number of batches and duration of a cam- paign?	This is a complex question and the scenario de- pends on many factors. Consideration should be given to designing and performing the pro- cess simulation so that it simulates the risks as- sociated with both the beginning and the end of the campaign and demonstrating the campaign duration does not pose any risk. The start-of- campaign (including aseptic assemblies if the case) AND end-of-campaign studies should be conducted in any case.
40	9.46	Can we differentiate between 1 CFU and >1 CFU like in ISO 13408-1 (2015) for 5'000 10'000 and > 10'000 units filled: if 1 CFU is detected, investiga- tion, and consideration of one APS, if > 1 CFU, investigation, corrective measures and repeti- tion of validation with 3 APS runs?	No. The new Annex 1 is stricter than the ISO 13408-1 (2015). Any contaminated unit with a contamination > 0 CFU results in a failed APS and actions according to chapter 9.46 should be followed.
41	9.46	With identification of root cause and corrective actions imple- mented, would it be acceptable to resume production (with batches at risk if a positive) prior to the 14-day reads off test and completion of successful growth promotion? Release of batches could only resume after completion of successful revali- dation (3x APS)?	No, as it clearly states, that PRODUCTION should resume only after completion of success- ful revalidation (9.46 vii)

4.7 Quality Control (QC) (Annex 1, Chapter 10)

Q&A No.	Paragraph No.	Questions	Answers
42	10.1	To support the design of manu- facturing activities, environ- mental monitoring regime etc., Annex 1 requires that person- nel with appropriate training and experience in microbiology and sterility assurance should	Microbiological knowledge (incl. sterility assur- ance) can be acquired by education, training and experience. The best prerequisite for the involvement of a person as an expert, for example in CCS as- sessments, the definition of resulting measures or in investigations on microbial contaminations,



	be available. What is consid-	is an education (in particular a university degree
	ered appropriate training and experience?	or an equivalent diploma e.g. an institution of higher technical education) in the field of micro- biology (or possibly other natural sciences, or
		medicine).
		However, a good understanding of the manufac-
		turing processes concerned is also required.
10.2	What limits do we expect for specifications for raw materials, components and products? What is typical?	The need for microbiological testing of raw ma- terials and the limits to be defined for such test- ing should take into account the nature of the raw materials (e.g., if of biological origin and whether they can be considered growth promot- ing) and their use in the respective process. The relevant chapters and monographs in the Phar- macopeia, the requirements as defined in the marketing authorisation and other regulations should be considered. Raw materials, components and products and their handling should be assessed as part of the CCS. The specifications should be justified.
10.3	Should bioburden be tested on	Yes
	each batch of raw material as	
	incoming control AND on the	
	compounding solution in which	
	it is formulated before sterile fil-	
	tration?	
10.6 iii		Lyophilization load means loads for each lyophi-
	•	lizer for each batch, if e.g. more than one lyophi-
	• •	lizer is used.
	•	
	lizer?	
10.10	How should situations for prod- ucts with short shelf life be han- dled when data exceeds the established limits (including OOS for sterility, see 10.) only after product batch certifica-	A procedure should be in place in case a post- release OOS should be obtained to inform phy- sicians, patient and health authorities, to assess the risk for the patient and to define remediation steps as needed. See also Annex 3: 45 and 46
	10.3 10.6 iii	10.2What limits do we expect for specifications for raw materials, components and products? What is typical?10.3Should bioburden be tested on each batch of raw material as incoming control AND on the compounding solution in which it is formulated before sterile fil- tration?10.6 iiiWhat does "different lyophiliza- tion loads" actually mean? First and last? Different lyophilizers? Worst cases? Does that mean each sample from a different ly- ophilizer or samples from differ- ent batches in the same lyophi- lizer?10.10How should situations for prod- ucts with short shelf life be han- dled when data exceeds the established limits (including OOS for sterility, see 10.) only

5 Changes to the previous version



• None