

MANAGEMENT SYSTEMS QUOTE REQUEST FORM



INSTRUCTIONS FOR COMPLETION:

Please ensure when completing this form it is downloaded and saved locally before completing. This interactive PDF should be opened and completed in Adobe Reader/Acrobat before resaving and returning to NQA.

IF YOU ARE A MULTI-SITE CLIENT PLEASE DOWNLOAD, COMPLETE AND RETURN THE NQA MULTI-SITE SUPPLEMENT QUESTIONNAIRE.

	THE ROA MOET-OFFE COTT ELIMENT GOLOTIONNAME.							
1. Organisatio	n details:							
Company name (Leg requiring certification				Con	untry:			
Main office address:								
Postcode:		We	bsite:					
Contact name:								
Job title:								
Email:								
Direct dial:			Mobile:					
2. Which mana	agement syste	ms standards are y	ou requiring	certificati	on for? (Tick a	ill that apply)		
ISO 9001:2015 (Qua Complete Section A	37		ISO 14001:2015 (Environmental) Complete Section B			ISO 45001:2018 (H&S) Complete Section C		
ISO 50001:2018 (End Complete Section D		ISO 13485:2016 Complete Section	(Medical Devices) on E		SSIP (Safety Syst	tems in Procurement on C	:)	
ISO 27001:2013 (Info	3 /	ISO 44001:2017 Complete Section			ISO 55001:2014 (Complete Section	,		
ISO 27701 (Privacy In Complete Section F		Please note; you must have this certification. If you are provider, then please apply	certified to ISO 27001 w	ith another	Transferring your Complete Section			
3. Integrated n	nanagement s	ystems:			Yes - full	Yes - partial	No	
Is your management	system integrated wi	th other standards and to wh	at extent?					
For further detail on	integration approac	ches within management sy	/stem standards, p	olease <u>click he</u>	ere.			
4. Please prov	4. Please provide details of the breakdown of your employees at this location:							
	Core hours	Shift 1	Shift 2		Shift 3	Total no. of emp	loyees	
No. of staff:								
Please detail the prod	cesses and activities	at this site:						

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Please detail the activities your employees conduct and the number involved in each task (e.g. maintenance, office based, production):

Task	Employees	Task	Employees	Task	Employees
Sales		Operations/Delivery – office/site based		R&D	
Marketing		Operations/Delivery – field based		Management	
Finance		Compliance		Other	
HR		Maintenance			
Total no. of employees:					
_	-	omplete and return an NQ		Questionnaire.	
5. Are you?	Yes	No		Yes No	
A new client?		Expar	nding your scope of certifi	cation?	
An existing NQA client?		Addin	g a standard to your certi	fication?	
A transferring client?		Addin	g a site to your certification	on?	
6. Requested sco	-				
		pose and output covered by al design services, or Inform			he organisation does,
7. Do vou provide	installation, cor	ntract site works o	· undertake vour l	business activity a	Yes No
client locations				, a a a a a a a a a a a a a a a a a a a	
					Yes No
R Do you have o	iteourood or euk	contracted activiti	ne?		Tes No
		rocesses, products and ser			
. ,	,, [, []			

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9. Does the organisation have a simple structure with vertical lines of management communication and few decision makers?									
Yes No 10. Does the organisation have staff speaking in more than one language and/or use an									
interpreter?									
If yes, please specify which language/s:									
11. Do you have a target ass	11. Do you have a target assessment date?								
12. At what stage of implement	entation are you in?								
Researching Implementing	System in place Already co	ertified							
13. Consultant use:			Yes	No					
Are you using a consultant to help you im	plement/manage the management system?								
Consultancy name/contact info:									
14. Where did you hear abou	ut NQA's service? (Tick all that apply	<u>n</u>							
Existing client	Event (exhibition or virtual)	Social media							
Consultant recommendation	Promotional email	Advertising campaign							
Professional recommendation	Professional recommendation NQA website Search engine (Google)								
Other (please specify)									
Please ensure that the following sections of this form are also completed (as appropriate). PLEASE CLICK BELOW TO GO DIRECTLY TO THE RELEVANT SECTION:									

If you have any problems completing this form please call +31 6 27034643 or email infobenelux@nqa.com

If you choose to give us any personal information (for example your e-mail address) we will treat this information in line with our privacy notice which can be located here: https://www.nqa.com/privacy. We will only use the information provided to respond to your enquiry and provide you with any information or materials requested. By submitting this information you are requesting a quote for services from NQA and a subsequent quote letter will be issued to you based on the information provided within this form.



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SECTION A - ISO 9001:2015

ONLY COMPLETE THIS SECTION IF APPLYING FOR CERTIFICATION AGAINST THIS STANDARD

	Yes	No
1. Do you undertake design and development of products and services?		
If yes, please detail the number of staff engaged in design activities:		

SECTION B - ISO 14001:2015

ONLY COMPLETE THIS SECTION IF APPLYING FOR CERTIFICATION AGAINST THIS STANDARD

Please complete the following questions considering ALL locations applying for certification. Yes No 1. Are your operations subject to an authorisation/permit/licence/registration from a regulatory body? (e.g. environmental permit, hazardous waste producer registration, abstraction licences, registered waste or water discharge exemptions, etc.) If yes, please provide details (including permit/licence/registration numbers): 2. Discharges to water/sewer: Do you produce any industrial effluent (other than domestic sewage and surface water)? Frequently Occasionally Neve 3. Waste: Do you produce hazardous, special or clinical waste? Neve Frequently Occasionally 4. Noise and nuisance: Have you had complaints with respect to noise or other nuisances (smoke, dust, fumes, Frequently Occasionally odours or other escapes) from your premises? Details, including which location(s) this applies to: 5. Incidents/prosecutions: Have you had, including significant stakeholder complaints any environmental incidents leading to high clean-up costs or a breach of legislation (including prosecution)? If you answered yes to any of the above questions, please provide details, including which location(s) this applies to: 6. Are any of the following site specific issues relevant? Are there any surface waters (rivers, lakes, streams, etc.) or boreholes within or adjacent to the site boundaries? No Yes Is your site overlying groundwater of significance (e.g. major/minor aquifer)? Yes No Do you have listed buildings (Grade I, Grade II*, Grade II) or archaeological sites (tumuli, burial mounds etc.) on site? Yes No Is the site within or adjacent to any designated nature conservation sites including Site of Special Scientific Interest (SSSI), Yes Nο National Park, or Special Areas of Conservation? Are there any other conservation issues at the site? Yes Is there evidence to suggest land contamination requiring clean-up is present at the site? Yes If you answered yes to any of the above questions, please provide details, including which location(s) this applies to:

SECTION C - ISO 45001:2018

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1. If you are applying for SSIP please identify						whicl	n role(s) you woul	d like	approving again	nst:
Designer		Principal Designer		Contra	ctor		Principal Contractor		Non-construction	
2. Plea	se pro	ovide details of th	ne ha	azards as	soci	iated	with your activitie	es:		
Hazards				Pl	ease t	tick	Please detail whic	h proces	sses these hazards re	late to?
Working w	ith asbe	stos								
Working w	ith explo	osives								
Working w	ith and s	storage of flammable sub	stance	es						
Transport	of dange	erous goods								
Underwate	er diving	at work								
Working w	ith mate	rials at extreme temperat	ures							
Working w	ith dang	erous animals								
Working ir	n proximi	ty to water (risk of drown	ing)							
Working w	ith gas									
Working w	ith ionisi	ng radiation								
Working w	ith lifting	equipment and lifting or	peration	าร						
Working w	ith biolo	gical hazards								
Working ir	n proximi	ty to moving vehicles								
Food prep	aration f	or other parties								
Working ir	n compre	essed air (risk of decomp	ressior	illness)						
Working a	t heights									
Working ir	n confine	d spaces								
Working w	ith press	sure systems								
Use of lea	Use of lead and heavy metals at work									
Working w	ith fume	s/gasses/dust								
Working w	ith chem	nical hazards								
Use of wo	rk equipı	ment (PUWER)								
Other (ple	ase spec	cify)								

3. Please identity the n	nain nazaro	ious materi	ais ass	socialed Wi	ur your proces	sses and pr	ovide	uetall	5 :
4. Radioactive and dar	ngerous sul	ostances:							
Do you keep, use, accumulate or	dispose of radio	active substance	es?				Yes	No	
Does your business handle, produce, use or store dangerous substances (including toxic, oxidising, explosive, flammable, etc.) in large quantities and could therefore be subjected to COMAH (Control of Major Accident Hazards)?									
If you have answered yes to any o	you have answered yes to any of the above questions, please provide details, including which location(s) this applies to:								
5. Are there members	of the publi	c present a	ıt your	organisatio	on's sites?		Yes	No	
If yes, please specify which sites:									
6. Please provide deta business:	ils of legisl	ation, regul	ations	, obligation	s and guidanc	e notes app	olicabl	e to t	he
E.g. Construction Design and Ma	nagement Regul	ations, Control o	of Major A	ccident Hazards	Regulations, etc.				
7. Please provide a de	scription of	any forma	l involv	ement with	n a competent	regulatory	author	ity:	
E.g. HSE in the UK					-				
8. Have you had any in enforcement notices If yes, please provide details:				g prosecut	ion/insurance	claims/	Yes	No	
, yee, preude provide detaile.									
9. Please state accurate months:	tely all inju	ies, diseas	es and	dangerous	s occurrences	(RIDDOR)	or the	past	12
Number of reportable injuries:	Fatal		Major		Over seven days				
Number of reportable dangerous	occurrences:		Numbe	r of reportable a	ccidents involving a r	member of the p	ublic:		
Details of reportable diseases:									
Details of reportable injuries:									
Note: Disclosure of information	n is a requireme	ent for contracti	ual obliga	ition. The applic	cant may be contac	ted before issue	of a qu	otation.	
personnel number (10. Are there any additional personnel that are not detailed in your employed personnel number (e.g. contractors/subcontractors personnel) performing work or work-related activities under the control or influence of the organisation's system?								
If yes, please state how many:									

SECTION D - ISO 50001:2018

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IF YOU ARE A MULTI-SITE CLIENT PLEASE DOWNLOAD, COMPLETE AND RETURN

THE NQA MULTI-SITE SUPPLEMENT QUESTIONNAIRE.

Number of EnMS effective personnel on site:	
Role(s) of EnMS personnel:	
Processes/activities of site:	
Annual energy consumption (Terajoules):	
Energy sources:	
Significant energy uses:	
Energy regulations applicable to site:	

For additional guidance on how to complete this section please <u>click here</u>.

SECTION E - ISO 13485:2016

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2. Are your products sterile?							Yes	No
If yes, please provide details of sterilization method:								
When/how was the sterilization conducted?	During production			Outso	ource	Inter	nd for end-use	r sterilizatio
Sterilization methods	Р	lease ti	ck			Details	6	
Ethylene oxide gas, (e.g. ethylene oxide gas sterilization	n):							
Moist heat (e.g. pressure steam sterilizer):								
Aseptic processing (e.g. sterilization by boiling; disinfed	tion; ozone disinfection):							
Radiation sterilization (e.g. gamma, x-ray, electron bean	n):							
Sterilization method other than specified above								
3. Is software used in the product?	Yes		No					
If yes, please provide details for software:								
As an independent medical used software?	Yes		No					
As a component part of the finished medical device?	Yes		No					
As an embedded part of the finished medical device?	Yes		No					
4. Have you had any incidents leading enforcement notices in the last year		secu	ition	ı/insı	ırance cl	aims/	Yes	No
If yes, please provide details:								
5. Please list below legal obligations	relevant to the pro	pose	d sc	ope	of certific	cation:		

Clause	Reason			
7. Organisational and proces	es complexity:			
	t range and/or complexity of medical device?	Yes	No	
	ply processes or parts that are critical to the function of the medical device			
and/or the safety of the user or finished pro		Yes	No	
Does the organisation install products on the	ne customer's premises?	Yes	No	
Does the organisation have poor regulatory	y compliance?	Yes	No	
Does the organisation have multiple shifts/a	a number of production lines?	Yes	No	
Does the organisation have no production	(e.g. wholesale, retail, transportation or maintenance of equipment?	Yes	No	
Does the organisation reduce the production	on range since last audit?	Yes	No	
boes the organisation reduce the production	office last addit:	162	NO	
If you answered yes to any of the above qu	estions, please provide details below:			

SECTION F - ISO 27001:2013

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.egal (e.g. Data Prote	ction Act, Computer Misuse Act etc):				
Regulatory (e.g. PCI [OSS, Information Governance Statement of Comp	liance (IG SoC)):			
2. Risk level ar	nd complexity - if you answer yes	to any of the below	v you	must	provide details:
Гуре	Criteria	Examples	Yes	No	Comments
Government classification	Do you handle Government information classified at or above secret?	e.g. military bases, defence supply chain, government departments.			
Nature of nformation managed	Could the nature of information held result in a breach or loss; having material financial, personal or reputational impact to any interested party? Information handled includes: Customers, end users, staff contractors or others sensitive personal information e.g. health records or financial information Intellectual property (e.g. designs, software source code)	e.g. Solicitors, law firms, banks, insurers, credit agencies (regulated by FCA), organisations providing payroll services or pension administration etc.			
Volume of data managed - aggregated data sets	Does the information held include a large set of sensitive personal information that could be used for identity theft or fraud? e.g. This could include individuals' usernames and passwords used to access web portals or other systems.	e.g. E-commerce websites, utility companies, online payment websites, organisations collecting individual's data via web portals, organisations processing and analysing customer data.			
Complexity of lechnology used	Does the technology used include a diverse or complex infrastructure? e.g. Many servers (>100 physical or virtual servers) AND/OR "Bring your own device" (BYOD) is permitted.	e.g. Large IT infrastructure, many servers, multiple different platforms, any organisation permitting BYOD ("bring your own device") is included in this criterion, regardless of size.			
Regulation	Is your organisation regulated? e.g. Regulated by Financial Conduct Authority, Ofcom, Ofsted, Oftel, Solicitors Regulatory Authority, Law Society, GMC). AND/OR Subject to sector specific rules e.g. Cheque Printers Accreditation Scheme C &CCC Standard 55, UK Health Service's Information Governance Statement of Compliance (IG SoC), ADISA (Asset	e.g. Banking, cheque printers, hospitals, education.			

Does your organisation develop software?

Complex tasks

Туре	Criteria	Examples	Yes No	Comments			
National importance of products/services & high availability requirements	Are your services: Part of critical national infrastructure (e.g. emergency services, communications, financial services, health, transport, utilities) AND/OR: An essential part of national infrastructure supply chain (e.g. data centre hosting national infrastructure systems) AND/OR: Potential terrorist target AND/OR: Non-availability of your services or product may severely affect the health, well-being, safety or security of people.	e.g. broadcasting support providers, utilities (power, water, gas), internet and mobile service providers, air traffic control, examination boards Or banking services, borders and immigration controls, health management systems.					
Supply Chain	Do you share sensitive information with third parties? e.g. Customers'/end users'/staff or others personal information. Including outsourced payroll, third party vetting services (criminal records, credit checks) AND/OR: Intellectual property (designs, source code or other sensitive proprietary information).	e.g. Criminal records, credit checks, outsourced payroll etc.					
Importance of integrity of information	If the information produced by your company is incorrect or incomplete, could there be a threat to individual or collective health / wellbeing / safety / security / miscarriage of justice or risk of fraud?	e.g. Organisations such as secure printers (passport/ visa printers/prescription/ medical instruction printers), health providers (clinical information/ medical record systems), gambling service providers.					
Susceptibility to fraud or targeted disruption	Could the theft of information (by staff / contractors or others) managed by your organisation result in fraud or targeted disruption? e.g. Theft of personal information by staff working in finance / insurance, call centres, clinics, pharmacies. AND/OR: Hacking of software/website/IT systems.	e.g. Organisations susceptible to fraud (e.g. by theft or misuse of data) or heightened risk of attempted fraud.					
Information not available to audit	Do you hold any ISMS related information that cannot be made available for review by the audit team because it contains confidential or sensitive information?	N/A					
Clearance	Does the audit team require security clearance to attend the site?						
	ISO 27701:2013 (PRIVACY INFORMATION MANAGEMENT) 1. Please detail below the data protection/privacy legislation applicable to your organisation: (e.g. GDPR)						
2. Are you currently or has your business ever been under investigation/fined by a data enforcement agency? (e.g. ICO) f yes, please provide details below:							
3. Please confi	rm whether your organisation is a	a data processor, d	lata control	ler or both:			
Data Controller Both Data Processor and Data Controller							

SECTION G - ISO 44001:2017

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1. Please provide the details below of the relationships you would like certificating:

	Collaborative Business Relationship to be certified	Number of employees involved in the Collaborative Business Relationship	Details of the Collaborative Business Relationship
1			
2			
3			
4			
5			
6			
7			
8			
9			
10			

SECTION H - ISO 55001:2014

ONLY COMPLETE THIS SECTION IF APPLYING FOR CERTIFICATION AGAINST THIS STANDARD

IF YOU ARE A MULTI-SITE CLIENT PLEASE DETAIL ON A SEPARATE SHEET THE ASSET GROUPS PERTAINING TO EACH SITE, UNLESS THESE ARE UNIFORM ACROSS ALL SITES

1. Please detail the business activities covered by your Asset Management System (AMS):							
2. Please list the different categories of Asset Groups below (use a separate sheet if necessary):							
	Asset group name	Asset group description	Company asset?	Client asset?			
e.g.	Vehicle Fleet	Lorries within vehicle fleet					
1							
2							
3							
4							
5							
3. Please select the most appropriate description applicable to your scope of AMS:							
The asset portfolio is a complex networked system of assets. It is a highly interdependent system.							
The asset portfolio is complex, but has discrete locations with partially interdependent systems.							
The asset portfolio is at a discrete location with independent functional systems.							
4. Please select the most appropriate description applicable to the criticality of your business assets within the scope of your AMS:							
High impact on business and stakeholders of asset failure.							
Medium impact on business and stakeholders of asset failure.							
Low impact on business and stakeholders of asset failure.							
5. Are there significant business continuity and supply chain risks? Yes No							
If yes, please provide details:							
6. Are there any statutory requirements for recording financial and non-financial information relevant to asset management, risk management, management of change, complexity of the outsourced processes etc.							
If yes, please provide details:							

SECTION I - TRANSFERRING

ANSWER THE FOLLOWING QUESTIONS IF YOU WISH TO TRANSFER YOUR CERTIFICATION FROM YOUR CURRENT CERTIFICATION BODY.

Please complete one transfer set of questions per certificate you wish to transfer to NQA.

1. Certificate details:							
Certificate number	Standard	Valid until date	Certification Body				
2. Reason for transferring:							
3. Are your certifications currently active?							
4. Have any complaints been raised against your organisation to your certification body, or is a regulatory body currently engaged with or investigating you in relation to activities you are certificated for? (e.g. HSE for health and safety breaches)							
If yes, please provide more information	n:						
5. Please detail the number	er of open major and/or m	inor No. of minors	No. of majors				
non-conformities on thi	s certificate:						
If one or more, please provide details:							
6. How frequently do you current certification boo		Annually	6 monthly Other				
7. Please detail your last audits up to and including the latest recertification or stage 2 audit:							
Audit type (Surveillance/Recert/Stage	2/Special)	Audit duration	Audit date				

To support your transfer please provide the following:

- Copies of your certificates
- · Audit reports for all audits conducted up to and including your last Recertification or Stage 2 audit
- Corrective action plan(s) for any non-conformances

If the required supporting documents are not provided a transfer may not be possible. NQA will contact your existing certification body to verify the validity of your certification. **Please note:** Do not cancel your certification with your existing certification body until the transfer process has been completed by NQA and you have received an NQA Certificate.