Certificate number:	
Application received date (YYYY/MM/DD):	



APPLICATION FORM FOR INDIVIDUAL CERTIFICATION AS

Auditor / Lead Auditor

PHARMACEUTICALS / MEDICAL DEVICE

This form is used for registration for the certification test for Personal Certification as Auditor / Lead Auditor - Pharmaceuticals / Medical Devices organized by SBQ Certification AB.

Send the completed and signed application (scanned pdf or digitally signed) by e-mail to info@sbq.nu

Fill out the form on the following pages with your experience and training in the subject.

The application will be processed by SBQ within 2-3 weeks. You will recieve feedback from us whether your application is properly filled out and if your experience in the subject is sufficient for certification. If you are approved, you will recive instructions on how to proceed and book your exam.

Note that you can perfom the certification test even if your experience is not enough (additional experience can be completed within one year).

The registration is binding upon receipt and approval by SBQ. The fee for certification is 530 EURO.

APPLICATION FORM

To be completed by person applying for certification as Auditor/Lead Auditor

Name:	Social security no (or other identification no):
Company:	Phone (work):
Address:	Phone (mobile):
Post code and City:	Email:
Invoice address (if other):	
I am applying for individual certification:	☐ Auditor Pharmaceuticals
	Lead Auditor Pharmaceuticals
	☐ Auditor Medical Devices
	☐ Lead Auditor Medical Devices

To ensure the link between application and certification (with respect to the same person), we need your social security number or other identification number. The personal identification number is used by SBQ for traceability to your certificate. Your personal information will not be used for other purposes. In accordance with the GDPR, this application for certification is the legal basis and the agreement underlying our handling of your personal data.

Continue your application by filling out the form on the following pages, as well as approve and sign your application on the last page.

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Work experience (Describe the experience five years back)



1.	Company/organisation:		Reference:
	From- until (YYYY/MM):		Name:
	Position:		E-mail:
	Industry:	Pharmaceuticals Medical Device Other	Relation: (for example manager, co-worker)
2.	Company/organisation:		Reference:
	From- until (YYYY/MM):		Name:
	Position:		E-mail:
	Industry:	Pharmaceuticals Medical Device Other	Relation: (for example manager, co-worker)
3.	Company/organisation:		Reference:
	From- until (YYYY/MM):		Name:
	Position:		E-mail:
	Industry:	Pharmaceuticals Medical Device Other	Relation: (for example manager, co-worker)
4.	Company/organisation:		Reference:
	From- until (YYYY/MM):		Name:
	Position:		E-mail:
	Industry:	Pharmaceuticals Medical Device Other	Relation: (for example manager, co-worker)
5.	Company/organisation:		Reference:
	From- until (YYYY/MM):		Name:
	Position:		E-mail:
	Industry:	Pharmaceuticals Medical Device Other	Relation: (for example manager, co-worker)
6.	Company/organisation:		Reference:
	From- until (YYYY/MM):		Name:
	Position:		E-mail:
	Industry:	Pharmaceuticals Medical Device Other	Relation: (for example manager, co-worker)

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Training related to audits, regulations and GMP/quality for pharmaceutical / medical device (last 5 years *)



	Course / training title:	Training organisation:	Date (YYYY/MM):	Length of training:
1.				
2.				
3.				
4.				
5.				
6.				
7.				
8.				
9.				
10.				
11.				
12.				
13.				

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1.	Date (YYYY/MM):		Contact information for the	company/organisation re-vised:	Contact data for the person confirming the audit execution*** (if other than contact)
	Total time of the audit* (hours):		Company/org.:		
	Which total time on the premises of the audited site ** (hours):		Contact person:		
	Your role in the audit:	Auditor Lead auditor	Address:		
	Internal or External audit:	Internal External	Phone no:		
	No of persons in the audit team:		E-mail:		
	Audit criteria: (eg. ISO13485, GMP Part 1, GMP Part 2, IVD, MDD)				
2.					
	Date (YYYY/MM):		Contact information for the	company/organisation re-vised:	Contact data for the person confirming the audit execution*** (if other than contact)
	Total time of the audit* (hours):		Company/org.:		
	Which total time on the premises of the audited site ** (hours):		Contact person:		
	Your role in the audit:	Auditor Lead auditor	Address:		
	Internal or External audit:	Internal External	Phone no:		
	No of persons in the audit team:		E-mail:		
	Audit criteria: (eg. ISO13485, GMP Part 1, GMP Part 2, IVD, MDD)				
3.	Data (WWW/NANA).		Contact information for the	company/organisation re-vised:	Contact data for the person confirming the
	Date (YYYY/MM):		contact mornation for the	company, organisation re-visea.	audit execution*** (if other than contact)
	Total time of the audit* (hours):		Company/org.:		
	Which total time on the premises of the audited site ** (hours):		Contact person:		
	Your role in the audit:	Auditor Lead auditor	Address:		
	Internal or External audit:	Internal External	Phone no:		
	No of persons in the audit team:		E-mail:		
	Audit criteria: (eg. ISO13485, GMP Part 1, GMP Part 2, IVD, MDD)				

- * Total number of hours of active audit work, ie including time for preparation, document review, on-site audit, report-writing and follow-up
- ** Number of hours that the audit is in progress from the opening session to the closing meeting
- *** This person should confirm that the audit has been conducted correctly and that the information you provided is accurate

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4.	Date (YYYY/MM):		Contact information for the	company/organisation re-vised:	Contact data for the person confirming the audit execution*** (if other than contact)
	Total time of the audit* (hours):		Company/org.:		
	Which total time on the premises of the audited site ** (hours):		Contact person:		
	Your role in the audit:	Auditor Lead auditor	Address:		
	Internal or External audit:	Internal External	Phone no:		
	No of persons in the audit team:		E-mail:		
	Audit criteria: (eg. ISO13485, GMP Part 1, GMP Part 2, IVD, MDD)				
5.					
	Date (YYYY/MM):		Contact information for the	company/organisation re-vised:	Contact data for the person confirming the audit execution*** (if other than contact)
	Total time of the audit* (hours):		Company/org.:		
	Which total time on the premises of the audited site ** (hours):		Contact person:		
	Your role in the audit:	Auditor Lead auditor	Address:		
	Internal or External audit:	Internal External	Phone no:		
	No of persons in the audit team:		E-mail:		
	Audit criteria: (eg. ISO13485, GMP Part 1, GMP Part 2, IVD, MDD)				
6.	Date (YYYY/MM):		Contact information for the	company/organisation re-vised:	Contact data for the person confirming the audit execution*** (if other than contact)
	Total time of the audit* (hours):		Company/org.:		
	Which total time on the premises of the audited site ** (hours):		Contact person:		
	Your role in the audit:	Auditor Lead auditor	Address:		
	Internal or External audit:	Internal External	Phone no:		
	No of persons in the audit team:		E-mail:		
	Audit criteria: (eg. ISO13485, GMP Part 1, GMP Part 2, IVD, MDD)				

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7.	Date (YYYY/MM):		Contact information for the	company/organisation re-vised:	Contact data for the person confirming the audit execution*** (if other than contact)
	Total time of the audit* (hours):		Company/org.:		
	Which total time on the premises of the audited site ** (hours):		Contact person:		
	Your role in the audit:	Auditor Lead auditor	Address:		
	Internal or External audit:	Internal External	Phone no:		
	No of persons in the audit team:		E-mail:		
	Audit criteria: (eg. ISO13485, GMP Part 1, GMP Part 2, IVD, MDD)				
8.	D . (\0004/2444)		Contact information for the	company/organisation re-vised:	Contact data for the nerson confirming the
	Date (YYYY/MM):		Contact information for the	company/organisation re-viseu.	Contact data for the person confirming the audit execution*** (if other than contact)
	Total time of the audit* (hours):		Company/org.:		
	Which total time on the premises of the audited site ** (hours):		Contact person:		
	Your role in the audit:	Auditor Lead auditor	Address:		
	Internal or External audit:	Internal External	Phone no:		
	No of persons in the audit team:		E-mail:		
	Audit criteria: (eg. ISO13485, GMP Part 1, GMP Part 2, IVD, MDD)				
€.	Date (YYYY/MM):		Contact information for the	company/organisation re-vised:	Contact data for the person confirming the
	Date (1111/www).				audit execution*** (if other than contact)
	Total time of the audit* (hours):		Company/org.:		
	Which total time on the premises of the audited site ** (hours):		Contact person:		
	Your role in the audit:	Auditor Lead auditor	Address:		
	Internal or External audit:	Internal External	Phone no:		
	No of persons in the audit team:		E-mail:		
	Audit criteria: (eg. ISO13485, GMP Part 1, GMP Part 2, IVD, MDD)				

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10.	Date (YYYY/MM):		Contact information for the	company/organisation re-vised:	Contact data for the person confirming the audit execution*** (if other than contact)
	Total time of the audit* (hours):		Company/org.:		
	Which total time on the premises of the audited site ** (hours):		Contact person:		
	Your role in the audit:	Auditor Lead auditor	Address:		
	Internal or External audit:	Internal External	Phone no:		
	No of persons in the audit team:		E-mail:		
	Audit criteria: (eg. ISO13485, GMP Part 1, GMP Part 2, IVD, MDD)				
11.	Date (YYYY/MM):		Contact information for the	company/organisation re-vised:	Contact data for the person confirming the audit execution*** (if other than contact)
	Total time of the audit* (hours):		Company/org.:		
	Which total time on the premises of the audited site ** (hours):		Contact person:		
	Your role in the audit:	Auditor Lead auditor	Address:		
	Internal or External audit:	Internal External	Phone no:		
	No of persons in the audit team:		E-mail:		
	Audit criteria: (eg. ISO13485, GMP Part 1, GMP Part 2, IVD, MDD)				
12				,	
12.	Date (YYYY/MM):		Contact information for the	company/organisation re-vised:	Contact data for the person confirming the audit execution*** (if other than contact)
	Total time of the audit* (hours):		Company/org.:		
	Which total time on the premises of the audited site ** (hours):		Contact person:		
	Your role in the audit:	Auditor Lead auditor	Address:		
	Internal or External audit:	Internal External	Phone no:		
	No of persons in the audit team:		E-mail:		
	Audit criteria: (eg. ISO13485, GMP Part 1, GMP Part 2, IVD, MDD)				

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- *** This person should confirm that the audit has been conducted correctly and that the information you provided is accurate

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13.	Date (YYYY/MM):		Contact information for the	company/organisation re-vised:	Contact data for the person confirming the audit execution*** (if other than contact)
	Total time of the audit* (hours):		Company/org.:		
	Which total time on the premises of the audited site ** (hours):		Contact person:		
	Your role in the audit:	Auditor Lead auditor	Address:		
	Internal or External audit:	Internal External	Phone no:		
	No of persons in the audit team:		E-mail:		
	Audit criteria: (eg. ISO13485, GMP Part 1, GMP Part 2, IVD, MDD)				
14.	Date (YYYY/MM):		Contact information for the	company/organisation re-vised:	Contact data for the person confirming the audit execution*** (if other than contact)
	Total time of the audit* (hours):		Company/org.:		
	Which total time on the premises of the audited site ** (hours):		Contact person:		
	Your role in the audit:	Auditor Lead auditor	Address:		
	Internal or External audit:	Internal External	Phone no:		
	No of persons in the audit team:		E-mail:		
	Audit criteria: (eg. ISO13485, GMP Part 1, GMP Part 2, IVD, MDD)				
15.	Date (YYYY/MM):		Contact information for the	company/organisation re-vised:	Contact data for the person confirming the audit execution*** (if other than contact)
	Total time of the audit* (hours):		Company/org.:		
	Which total time on the premises of the audited site ** (hours):		Contact person:		
	Your role in the audit:	Auditor Lead auditor	Address:		
	Internal or External audit:	Internal External	Phone no:		
	No of persons in the audit team:		E-mail:		
	Audit criteria: (eg. ISO13485, GMP Part 1, GMP Part 2, IVD, MDD)				

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APPLICATION FORM FOR INDIVIDUAL CERTIFICATION AS

Auditor / Lead Auditor

PHARMACEUTICALS / MEDICAL DEVICE

Declaration and signature	
I hereby apply for individual certification as Auditor / L document, I certify that the information provided is co binding upon receipt and approved by SBQ.	
I accept that the fee of 530 EUR will be charged (25% V companies within the EU, the "Reverse charge" tax rule $^{\circ}$	
My signature also means that I allow the certification be accordance with GDPR.	oody SBQ to save my personal data in
Date: S Name in block letters:	ignature:

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