



# NORDIC QAFORUM

Clarion Hotel Copenhagen Airport  
November 21st, 2024

## THE NATIONAL AUTHORITIES:



Peter Borgå



Emil Schwan



Eva Tollefsen



Thomas Noe Vestergaard Pedersen



Katja Belt



## Specially invited:



"European Shortages Monitoring Platform – how will it impact the industry?"

Véronique Davoust  
Pfizer

- GDP/GMP – the most common findings
- European Shortages Monitoring Platform
- Fiscal import
- Inspections of wholesalers
- The responsibility of MAH
- + much more

We offer live stream

## Speakers:

**Danish Medicines Agency:**  
Thomas Noe Vestergaard Pedersen  
Medicines Inspector & Team manager

**Finnish Medicines Agency:**  
Katja Belt  
Senior Pharmaceutical inspector

**Norwegian Medicines Agency:**  
Eva Tollefsen  
Pharmaceutical inspector

**Swedish Medical Products Agency:**  
Peter Borgå  
Pharmaceutical inspector

**Swedish Medical Products Agency:**  
Emil Schwan  
Pharmaceutical inspector

**The Danish Medicines Verification Organisation (DMVO):**  
Marie Louise Shee  
Managing Director

**Pfizer:**  
Véronique Davoust  
Global Supply Global Quality Operations, Regulatory Intelligence

**Novo Nordisk Denmark:**  
Irene Bøggild  
Quality Manager & RP

## Partners:





# NORDIC QA FORUM

Dear Colleague,

There has never been a more relevant time to have a unique platform dedicated to discuss the key challenges that you and your peers are facing in the field of Quality Assurance and Regulatory Affairs.

**The Nordic QAforum will give you the opportunity to :**

- Understand how the authorities in Sweden, Denmark, Norway and Finland will interpret the changes.
- Find out best practices.
- Network with other senior QA & RA professionals.
- Learn how to implement the new requirements.

Nordic QAforum is a must-attend conference dedicated to supporting your work in QA and RA by facilitating high-level networking opportunities with your peers and providing leading industry knowledge.

I look forward to meeting you in November!

Kind regards,

Niko Fastman  
Project Manager Nordic QAforum & Qaforum  
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niko.fastman@kompetensinstitutet.se

**Our latest Nordic Qaforum received the overall rating 4,51 out of 5.0.  
Register today to secure your ticket.**

## Testimonials of our previous QAforum:

*"Very good conference. Lots of opportunities to network with colleagues."*

Tomas Wahlgren, AstraZeneca

*"Fantastic selection of topics and speakers. This was one of the best events I have participated in."*

*"I appreciated the networking and the opportunity to listen to MPA. Excellent initiative!"*

### TARGET AUDIENCE:

The Nordic QAforum is of particular interest to management and personnel from pharmaceutical companies as well as from distributors and service providers involved in quality assurance and regulatory affairs.

### LIMITED TICKETS AVAILABLE:

Our latest QAforum was fully booked early and received the grade 4,51 out of 5.0.  
Register today to secure your ticket.

08.30 Registration, morning coffee & refreshments

09.15 Chairperson Anna Pontén Engelhardt  
welcome & opening remarks

Expert presentation:

09.20 European Shortages Monitoring Platform  
– how will it impact the industry?

- What is the ESMP and how will it work?
- How will this new initiative impact pharma industry and supply chain?
- When is the platform coming live and how can the industry start preparing? What parts of the company will it involve?
- Will the ESMP replace national initiatives? How can companies and national authorities interact around the ESMP?
- How can the ESMP be set up on time with the right quality?
- EFPIAs/MFE initiative to use EMVS (European Medicines Verification System) in connection to the ESMP

Véronique is responsible for the monitoring and analysis of global and European emerging regulations and guidelines. Furthermore, she ensures the communication and implementation of the guidelines and regulations within the firm, as well as the coordination of responses to the authorities.

She is deeply involved in Trade Associations activities (eg. European Federation of Pharmaceutical Industries and Associations (EFPIA) or the French LEEM), serving on leadership teams in quality or supply chain. She currently supports the development of EMA's European Shortages Monitoring Platform (ESMP) as Efpia's subject matter expert.

**Véronique Davoust**, Global Supply Global Quality Operations, Regulatory Intelligence, Pfizer

Case study:

10.00 Handling counterfeit in an affiliate

**Irene Bøggild**, Quality Manager & RP, Novo Nordisk Denmark

10.25 Questions to the speakers

10.40 Coffee break & networking

Norwegian Medicines Agency:

11.05 Inspections of wholesalers

- Inspections of wholesalers with common Nordic quality organizations
- The most common deviations, 2021-2024
- Deviations/findings that surprised me

**Eva Tollefsen**, Pharmaceutical inspector, Norwegian Medicines Agency

Finnish Medicines Agency:

11.35 Common issues in supplier management in GMDP environment

- The most common deviations, 2021-2024
- Deviations/findings that surprised me

**Katja Belt**, Senior Pharmaceutical Inspector, Finnish Medicines Agency

FMD:

12.05 Falsified Medicines Directive

- How we deal with alerts
- How we work with improving quality on both the industry and end-user side
- What are the learnings from implementing the FMD in Denmark?
- What is the current status working with the FMD and what is currently in focus for the Danish Medicine Verification Organization?
- Future changes and development?

**Marie Louise Shee**, Managing Director, The Danish Medicines Verification Organisation (DMVO)

12.30 Questions to the speakers

12.40 Lunch

Danish Medicines Agency:

13.40 Dicillin-recall case Part 2: Next steps from an authority perspective

- Impact on inspection practices
- Coming changes in guidance documents to prevent the issue to reoccur
- The most common deviations, 2021-2024
- Deviations/findings that surprised me

**Thomas Noe Vestergaard Pedersen**, Medicines Inspector & Team Manager, Danish Medicines Agency

Icelandic Medicines Agency:

14.15 Topic to be announced

Speaker to be announced

14.45 Questions to the speakers

15.00 Coffee break & networking

Swedish Medical Products Agency:

15.30 Import of medicinal products into the EES – which transactions are acceptable and which are not?

The responsibility of MAH: requirements for GMP and GDP oversight

**Emil Schwan**, Pharmaceutical Inspector, Swedish Medical Products Agency

**Peter Borgå**, Pharmaceutical Inspector, Swedish Medical Products Agency

16.45 Final Questions to the speakers

17.15 Chairpersons closing remarks

17.20 Reception & networking

It has been a long day filled with information. Time to relax and enjoy great conversations over drinks.

Pre-conference networking:

If you arrive on Wednesday - welcome to the informal pre-conference networking at Clarions Bar starting at 20.00.



# NORDIC QA FORUM

## 3 ways to register:

**Website:** [www.nordicqaforum.com/register](http://www.nordicqaforum.com/register)

**E-mail:** [kundtjanst@kompetensinstitutet.se](mailto:kundtjanst@kompetensinstitutet.se)

**Phone:** +46 (0) 8 20 03 73

## VENUE:

Clarion Hotel Copenhagen Airport  
Ellehammersvej 20  
Copenhagen  
Phone nr to the venue:  
+45 32 50 15 01

Transport from Copenhagen Airport:  
2 min walk through a covered  
walk-way

## Price:

**SAVE 200 DKK**  
**- register before 20th of August**

Nordic Qaforum November 21st 2024:

7990 DKK

**7790 DKK**

Lunch, coffee, networking reception evening before the conference, networking reception after the conference, documentation and certificate of participation is included in the delegate fee.

## Interested in a Live stream?

Please contact: [kundtjanst@kompetensinstitutet.se](mailto:kundtjanst@kompetensinstitutet.se)



KOMPETENS  
INSTITUTET

## About QAforum:

QAforum, a division of Kompetensinstitutet, provides a forum to address the critical issues facing the Pharmaceutical Industry today. QAforum utilizes workshops and conference formats to facilitate a learning environment for pharmaceutical professionals working within the areas of quality assurance and regulatory affairs.

## Terms and Conditions

In the event that Kompetensinstitutet cancels an event for any reason, you will receive a complete refund.

You can pay by invoice or by creditcard. Payment is required 30 days from the date of invoice. All prices are excluding VAT. The booking is binding after you received a confirmation of the booking. You may substitute delegates at any time by informing QAforum. For any cancellations received in writing not less than fourteen (14) days prior to the conference, you will receive a credit of the invoiced amount to be used at another Kompetensinstitutet conference which must occur within two years from the date of issuance of such credit. No credit will be issued for any cancellations occurring within thirteen (13) days of the conference. Kompetensinstitutet reserves the right to change date, format or to cancel an event. Complete terms & conditions: [www.kompetensinstitutet.se/villkor/](http://www.kompetensinstitutet.se/villkor/)