

Feedback from Registration

*ECHA's Fourth Stakeholders Day
19 May 2010*

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ECHA – Registration and Dossier Submission Unit*

Feedback from Registration

- This information will help you to guarantee a successful registration
- All the tools and supporting information are in place
- Allow time to use these tools and to understand the submission process

Feedback from Registration

- **Registration: predictions**
- Overview of the registration process
- Latest registration statistics
- Key messages to ensure a successful registration

Predictions

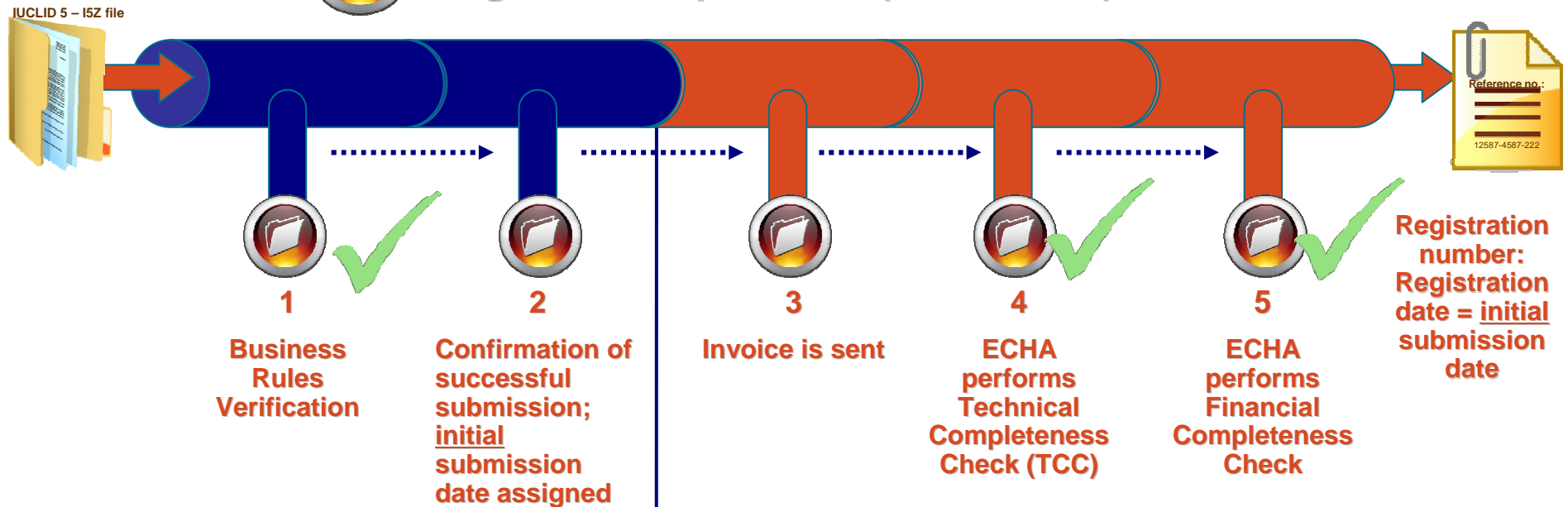
- Information from several sources:
 - Industry survey
 - ECHA survey
 - Lead Registrant Nominations
 - Historical data
- Expect registrations for ca. 5000 substances
 - List of 4515 identified to be registered published on our web site
- Based on the ratio of lead to member we would expect to receive ca. 38,000 dossiers

Feedback from Registration

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Registration process (REACH-IT)



‘Pre-processing’:

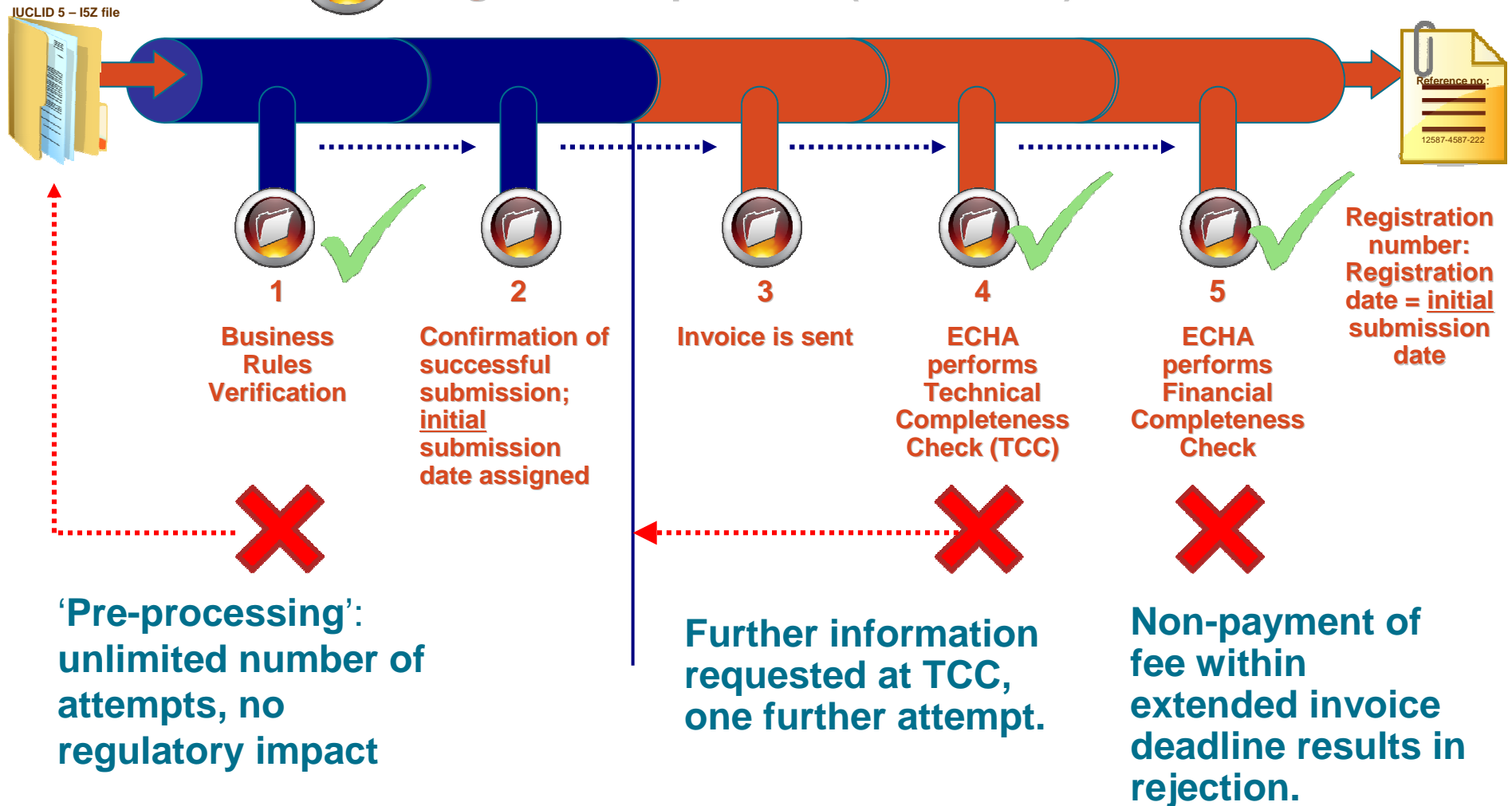
This step **must** be successfully completed before the deadline.

Earlier submission recommended.

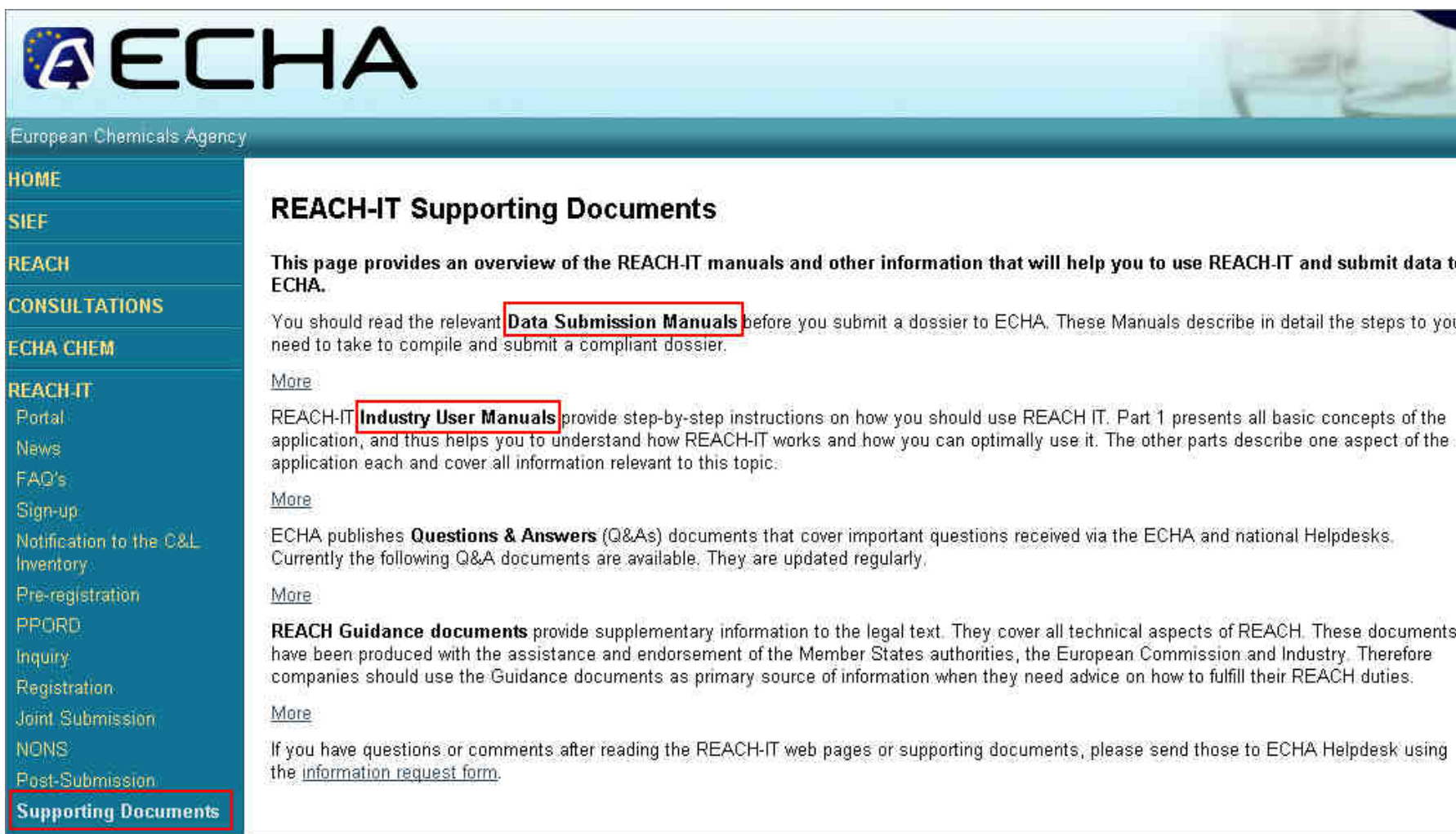
These steps can take place after the deadline, reasonable deadline set in case of TCC failure.



Registration process (REACH-IT)



Further information: Manuals



ECHA
European Chemicals Agency

HOME
SIEF
REACH
CONSULTATIONS
ECHA CHEM
REACH-IT
Portal
News
FAQ's
Sign-up
Notification to the C&L Inventory
Pre-registration
PPORD
Inquiry
Registration
Joint Submission
NONS
Post-Submission
Supporting Documents

REACH-IT Supporting Documents

This page provides an overview of the REACH-IT manuals and other information that will help you to use REACH-IT and submit data to ECHA.

You should read the relevant **Data Submission Manuals** before you submit a dossier to ECHA. These Manuals describe in detail the steps to you need to take to compile and submit a compliant dossier.

[More](#)

REACH-IT **Industry User Manuals** provide step-by-step instructions on how you should use REACH IT. Part 1 presents all basic concepts of the application, and thus helps you to understand how REACH-IT works and how you can optimally use it. The other parts describe one aspect of the application each and cover all information relevant to this topic.

[More](#)

ECHA publishes **Questions & Answers** (Q&As) documents that cover important questions received via the ECHA and national Helpdesks. Currently the following Q&A documents are available. They are updated regularly.

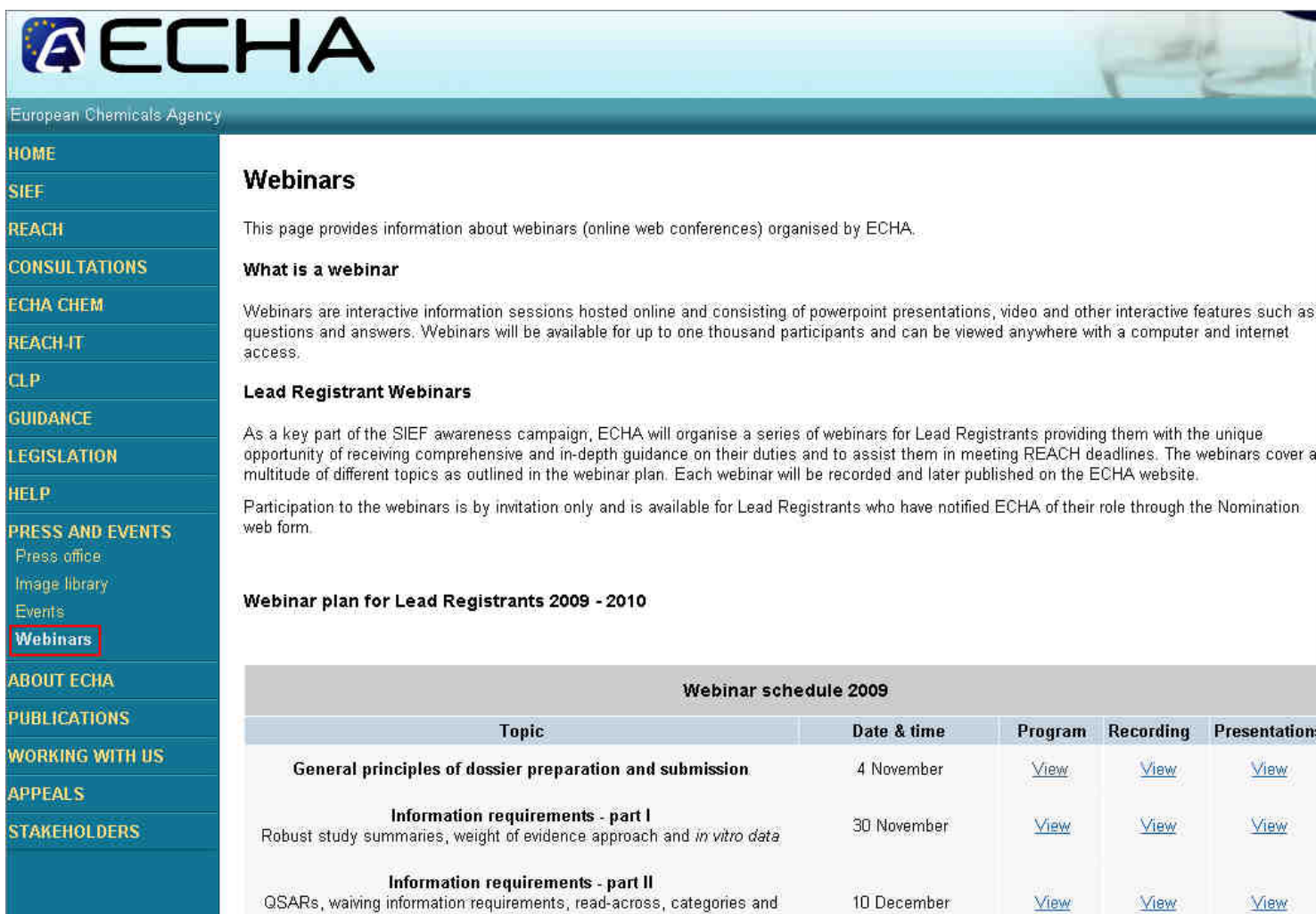
[More](#)

REACH Guidance documents provide supplementary information to the legal text. They cover all technical aspects of REACH. These documents have been produced with the assistance and endorsement of the Member States authorities, the European Commission and Industry. Therefore companies should use the Guidance documents as primary source of information when they need advice on how to fulfill their REACH duties.

[More](#)

If you have questions or comments after reading the REACH-IT web pages or supporting documents, please send those to ECHA Helpdesk using the [information request form](#).

Further information: Webinars



The screenshot shows the ECHA website's 'Webinars' page. The left sidebar contains a navigation menu with 'Webinars' highlighted. The main content area includes a title 'Webinars', a brief description, a definition of a webinar, information about Lead Registrant Webinars, and a table titled 'Webinar schedule 2009'.

Webinars

This page provides information about webinars (online web conferences) organised by ECHA.

What is a webinar

Webinars are interactive information sessions hosted online and consisting of powerpoint presentations, video and other interactive features such as questions and answers. Webinars will be available for up to one thousand participants and can be viewed anywhere with a computer and internet access.

Lead Registrant Webinars

As a key part of the SIEF awareness campaign, ECHA will organise a series of webinars for Lead Registrants providing them with the unique opportunity of receiving comprehensive and in-depth guidance on their duties and to assist them in meeting REACH deadlines. The webinars cover a multitude of different topics as outlined in the webinar plan. Each webinar will be recorded and later published on the ECHA website.

Participation to the webinars is by invitation only and is available for Lead Registrants who have notified ECHA of their role through the Nomination web form.

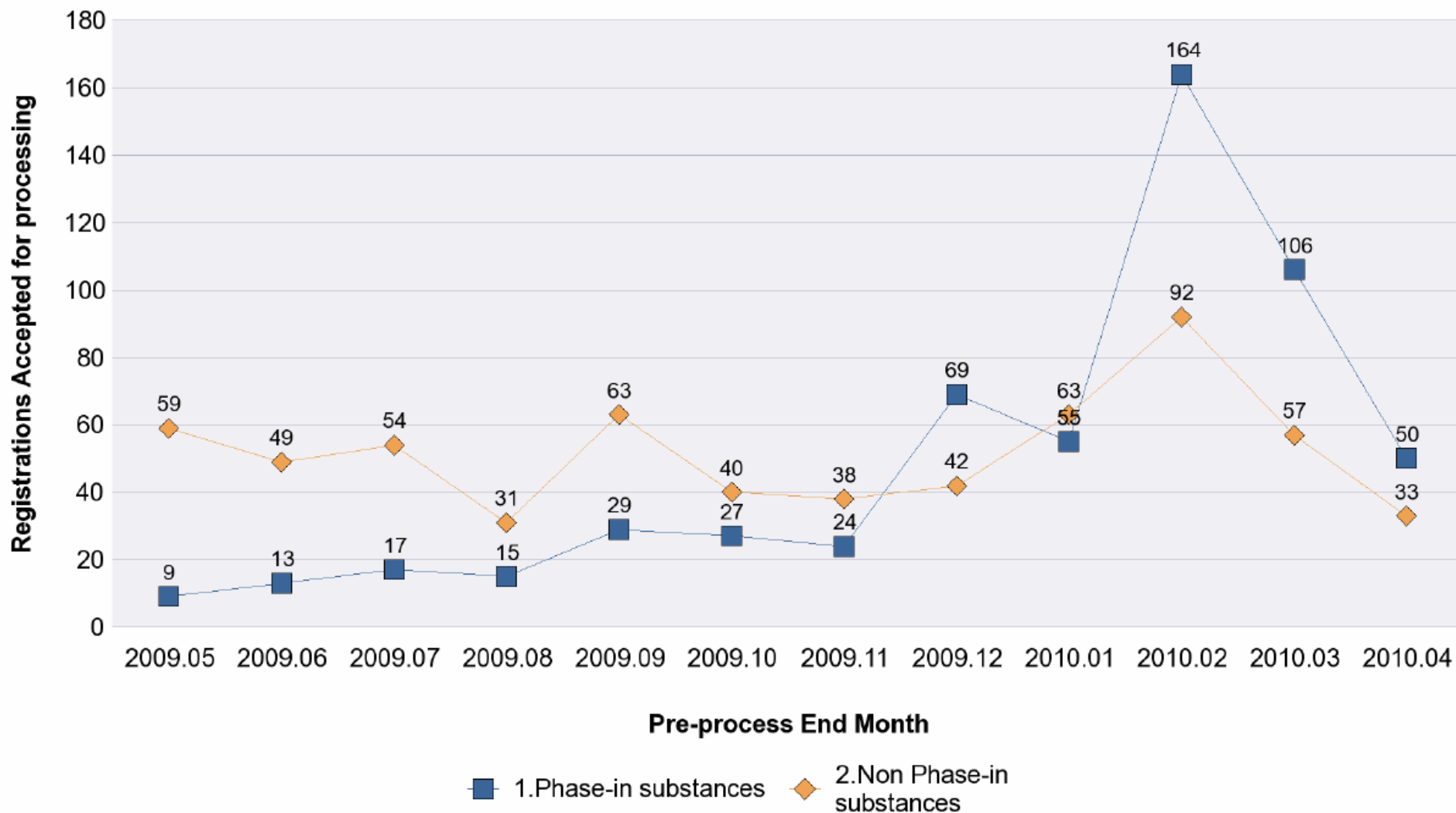
Webinar plan for Lead Registrants 2009 - 2010

Webinar schedule 2009				
Topic	Date & time	Program	Recording	Presentations
General principles of dossier preparation and submission	4 November	View	View	View
Information requirements - part I Robust study summaries, weight of evidence approach and <i>in vitro</i> data	30 November	View	View	View
Information requirements - part II QSARs, waiving information requirements, read-across, categories and	10 December	View	View	View

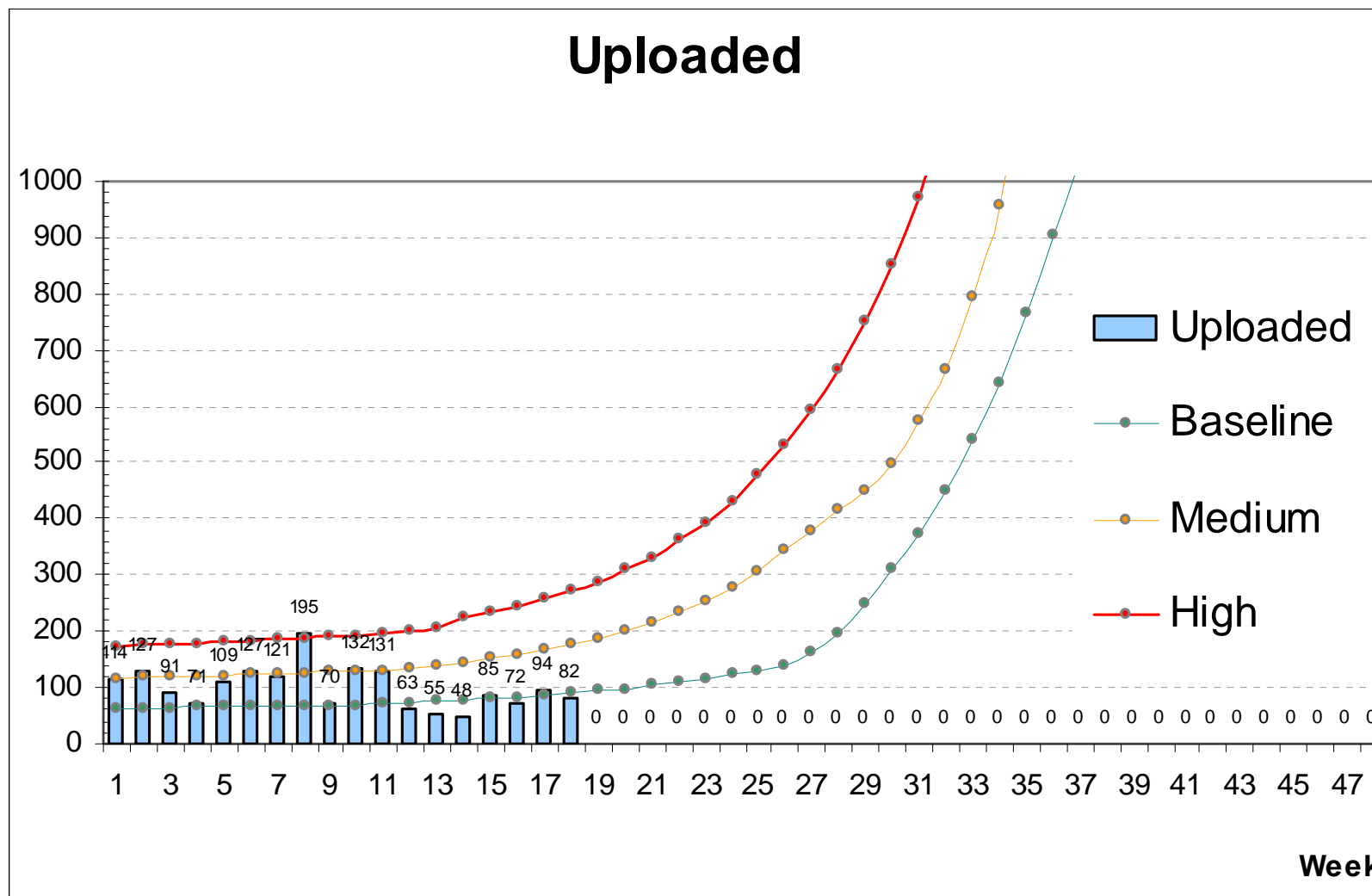
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Registration statistics I



Registration statistics II



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Key messages for successful registration I



- Passing the Business Rules verification:
 - Read the main section of Data Submission Manual 4 (five pages)
 - Watch the Lead Registrant Webinars
 - Submit as early as possible. This step **must** be completed before the deadline

- Passing the Technical Completeness Check (TCC)
 - Use the TCC tool
 - If failures occur, Data Submission Manual 5 provides screenshots and detailed information
 - Ample time will be given to correct missing information, and you remain legally on the market – contact relevant helpdesk for assistance if needed

Key messages for successful registration I



- Ensure fee payment is made in time:
 - Check payment mechanism with your accounting dept. Can be arranged in advance.
 - Ensure that payment is made within the extended deadline (44 days from invoicing date), or the dossier will be rejected
- Familiarise yourself with the submission process:
 - REACH-IT Industry User Manuals
 - Data Submission Manuals
 - Lead Registrant Webinars
- ECHA is fully committed to providing the best service possible; let's make this work!

Feedback from Registration



<http://echa.europa.eu>

CLP