WHAT HAS CHANGED AND WHY?
Lukas Bruckner
Ph.Eur. expert group 15V

2011: Ph.Eur 7th edition
0062 Vaccines for Veterinary Use

2-1-3-2-6. Absence of extraneous viruses. ...

- ... neutralising antibody to the virus of the seed lot ... other methods are used to remove or neutralise the seed virus specifically.
- ... at least the virus content of 10 doses of vaccine per 0.1 mL for avian vaccines and per millilitre for other vaccines.
- ... inoculated onto cultures of at least 70 cm² of the required cell types ... at any suitable stage of growth up to 70 per cent confluency. At least 1 monolayer of each type must be retained as a control. ... monitored daily for a week. ... freeze thawed 3 times, centrifuged ... re-inoculated onto the same cell type ... repeated twice.
  - Cytopathic and haemadsorbing agents are tested
  - immuno-fluorescence
- inoculated onto:
  - primary cells of the species of origin of the virus,
  - cells sensitive to viruses pathogenic for the species for which the vaccine is intended,
  - cells sensitive to pestiviruses.
2.6.24/2.6.25 AVIAN LIVE VIRUS VACCINES: TESTS FOR EXTRANEOUS AGENTS

1. TEST FOR EXTRANEOUS AGENTS USING EMBRYONATED HENS’ EGS
   - group 1: 0.2 mL into the allantoic cavity of each 9- to 11-day-old embryonated egg
   - group 2: 0.2 mL onto the chorio-allantoic membrane ...

2. TEST IN CHICKEN KIDNEY CELLS
   - Prepare 7 monolayers of primary or secondary chick embryo fibroblasts ...

3. TEST FOR AVIAN LEUCOSIS VIRUSES

4. TEST FOR AVIAN RETICULOENDOTHELIOSIS VIRUS

5. TEST FOR CHICKEN ANAEMIA VIRUS

6. TEST FOR EXTRANEOUS AGENTS USING CHICKS
   - Inoculate each of at least 10 chicks with the equivalent of 100 doses of vaccine ...

Lukas Bruckner, What has changed and why?
Training on the Management of Extraneous Agents in IVMPs, Strasbourg (F), 1 April 2020
<table>
<thead>
<tr>
<th>Agent</th>
<th>Type of test</th>
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<tr>
<td><strong>Standard tests</strong></td>
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<tr>
<td>Avian adenoviruses, group 1</td>
<td>SN, EIA, AGP</td>
</tr>
<tr>
<td>Avian encephalomyelitis virus</td>
<td>AGP, EIA</td>
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<td>Avian infectious bronchitis virus</td>
<td>EIA, HI</td>
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<td>Avian infectious laryngotracheitis virus</td>
<td>SN, EIA, IS</td>
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<td>Avian leucosis viruses</td>
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<tr>
<td>Avian nephritis virus</td>
<td>IS</td>
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<tr>
<td>Avian orthoreoviruses</td>
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<tr>
<td>Avian reticuloendotheliosis virus</td>
<td>AGP, IS, EIA</td>
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<td>Chicken anaemia virus</td>
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<tr>
<td>Egg drop syndrome virus</td>
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<tr>
<td>Avian infectious bursal disease virus</td>
<td>Serotype 1: AGP, EIA, SN Serotype 2: SN</td>
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<tr>
<td>Influenza A virus</td>
<td>AGP, EIA, HI</td>
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<td>Marek’s disease virus</td>
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<td>Newcastle disease virus</td>
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<td><strong>Additional tests for turkey extraneous agents</strong></td>
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<tr>
<td>Chlamydia spp.</td>
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<tr>
<td>Salmonella pullorum</td>
<td>Agg</td>
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<tr>
<td>Avian infectious haemorrhagic enteritis virus</td>
<td>AGP</td>
</tr>
<tr>
<td>Avian paramyxovirus 3</td>
<td>HI</td>
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<tr>
<td>Avian infectious bursal disease virus type 2</td>
<td>SN</td>
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<td><strong>Additional tests for duck extraneous agents</strong></td>
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<tr>
<td>Chlamydia spp.</td>
<td>EIA</td>
</tr>
<tr>
<td>Duck and goose parvoviruses</td>
<td>SN, EIA</td>
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<tr>
<td>Duck enteritis virus</td>
<td>SN</td>
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<tr>
<td>Duck hepatitis virus type I</td>
<td>SN</td>
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<tr>
<td><strong>Additional tests for goose extraneous agents</strong></td>
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<tr>
<td>Duck and goose parvovirus SN</td>
<td>EIA</td>
</tr>
<tr>
<td>Duck enteritis virus</td>
<td>SN</td>
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<tr>
<td>Goose haemorrhagic polyomavirus</td>
<td>test in goslings shown ...</td>
</tr>
</tbody>
</table>
5.2.4 CELL CULTURES FOR THE PRODUCTION OF VETERINARY VACCINES

- Absence of contaminating viruses.
  - The cells must not be contaminated by viruses; suitably sensitive tests, including those prescribed below, are carried out. The monolayers tested shall have an area of at least 70 cm², ... maintained ... for a total of at least 28 days. Subcultures are made at 7-day intervals, ...

- Detection of cytopathic viruses.
  - Two monolayers of at least 6 cm² each are stained with an appropriate cytological stain, ... examined for any inclusion bodies, abnormal numbers of giant cells or any other lesion indicative of a cellular abnormality ...

- Detection of haemadsorbent viruses.
  - Monolayers totalling at least 70 cm² are washed several times with an appropriate buffer and a sufficient volume of a suspension of suitable red blood cells added to cover the surface of the monolayer evenly. After different incubation times cells are examined for the presence of haemadsorption.

- Detection of specified viruses.
  - Tests are carried out for freedom from contaminants specific for the species of origin of the cell line and for the species for which the product is intended.

- Tests in other cell cultures.
  - Monolayers totalling at least 140 cm² are required ...
5.2.5 SUBSTANCES OF ANIMAL ORIGIN FOR THE PRODUCTION OF IMMUNOLOGICAL VETERINARY MEDICINAL PRODUCTS

5.2.5 SUBSTANCES OF ANIMAL ORIGIN FOR THE PRODUCTION OF IMMUNOLOGICAL VETERINARY MEDICINAL PRODUCTS

Freedom from living extraneous viruses.

- A sample from each batch of the substance is tested for extraneous viruses by general and specific tests. These tests are validated with respect to sensitivity and specificity for detection of a suitable range of potential extraneous viruses. Suitably sensitive cell cultures are used for the tests for extraneous viruses ....

- General test. The inoculated cell cultures are observed regularly for 21 days for cytopathic effects, ... and a proportion is tested for haemadsorbing agents.

- Specific tests. ... The specific viruses to be tested for are potential extraneous viruses that are identified through the risk assessment and that would not be detected by the general test. A test for pestiviruses ...

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Specific Monographs for Veterinary Vaccines

- Inactivated viral vaccines for pigs
  - *Extraneous agents.*
    - On the pigs used for the safety test carry out tests for antibodies. The vaccine complies with the test if it does not stimulate the formation of antibodies, other than those against the vaccine virus, against viruses pathogenic for pigs or against viruses that could interfere with the diagnosis of infectious diseases of pigs (including the viruses of the pestivirus group).

- Inactivated viral vaccines for chicken
  - *Extraneous agents.*
    - Use the chickens from the safety test. 21 days after injection of the double dose of vaccine, inject 1 dose by the same route into each chicken. Collect serum samples from each chicken 2 weeks later and carry out tests for antibodies against the following agents: avian encephalomyelitis virus, avian leucosis viruses, egg-drop syndrome virus, avian infectious bursal disease virus, avian infectious laryngotracheitis virus, influenza A virus, Marek’s disease virus, Newcastle disease virus.

Specific Monographs for Veterinary Vaccines

- Live viral vaccines for Mammals
  - *Extraneous agents.*
    - Neutralise the vaccine virus with a suitable monospecific antiserum against the vaccine virus and inoculate into cell cultures known for their susceptibility to viruses pathogenic for the target species. Carry out at least one passage and maintain the cultures for 14 days. The vaccine complies with the test if no cytopathic effect develops and there is no sign of the presence of haemadsorbing agents. (Carry out a specific test for pestiviruses.)

- Avian Live viral vaccines
  - *Extraneous agents.*
    - The vaccine complies with the tests for extraneous agents in batches of finished product (2.6.25).
Limitations of the approach in handling extraneous agents

- No harmonized approach, requirements scattered over several texts
- Testing requirements are different from species to species
  - Very precise for avian vaccines
  - List of agents to be tested for
  - General approach for vaccines for mammals
  - No policy for vaccines for fish
- Molecular methods are neglected
- Focus on laboratory testing only
- Consequences of good manufacturing are not taken in account

Limitations of the approach to handle extraneous agents

- Tests in tissue culture, embryonated eggs (or animals)
  - Detailed description of the test to be performed
    - Number and size of test replicates
    - Volume of inoculate
    - Incubation time and temperature
    - Test evaluation: macroscopic and microscopic examination, staining of cultures, cytopathic effects, hemadsorption, immunofluorescence ...
  - No positive controls
  - Suitability of the test justified by historic experience
Harmonization

- All requirements in one chapter

5.2.5 Management of extraneous agents in immunological veterinary medicinal products

- Applies to all substances: seed material, material used during production, etc
- Risk management
- Control measures
  - Starting material
  - Control measures during production
  - Methods for detection of extraneous agents
    - 2.6.1 Sterility
    - 2.6.21 Nucleic acid amplification techniques
    - 2.6.7 Mycoplasmas
    - 2.6.37 Principles for the detection of extraneous viruses in immunological veterinary medicinal products using culture methods

- ANNEX I: LIST OF EXTRANEOUS AGENTS TO BE CONSIDERED FOR THE RISK ASSESSMENT
Extraneous agents testing in the European Pharmacopoeia (Ph. Eur.) for IVMPs

The overall new approach
EDQM training on the Management of Extraneous Agents in IVMPs

April 1-2, 2020 EDQM, Strasbourg

Catherine LANG, EDQM, Group 15V Secretary
STRUCTURE

- Revision of the extraneous agents testing approach
  - Where are we now and what are the next steps?
  - Aim and drivers
  - Impact on the Ph. Eur. texts
  - New approach brings opportunities and benefits

Training replaced by a GoToWebinar on 1st April 2020

NEW: Training live-broadcast on the Management of Extraneous Agents in IVMPs

REGISTRATION FEES
No registration fees.

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Where are we now and next steps

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<tr>
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<td>EDQM training for all interested parties on 1st April 2020</td>
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<td>2 Hearings and consultation in Phpa 30.2 deadline for comments 30/06/2018</td>
<td>Ph. Eur. COM adopted</td>
<td>Ph. Eur. Supplement 10.2 Publication: January 2020 Implementation: 1 July 2020</td>
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<tr>
<td>EDQM Explanatory document</td>
<td>EDQM Technical guide</td>
<td>EDQM Concrete examples</td>
<td>EMA Q&amp;A document</td>
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EA testing adopted 06/2019; publication 01/2020; implementation 07/2020

Revision of the extraneous agents testing approach:

- **43 Ph. Eur. texts** involved in the revision
- Requirements collated and harmonised, compiled in chapter 5.2.5 “Management of extraneous agents in immunological veterinary products”
- Risk assessment + list of extraneous agents (avian revised list + EMA IWP list as is)
- Open to any fit-for-purpose method

**OTHER CHANGES (out of scope of the training)**

- Deletion of the test for Specified Extraneous Agents in some individual monographs (batch test using 2 animals (pigs, cattle, rabbits))
- To continue to allow the use of antibiotics during vaccine production, but request justification for such use (general monograph 0062 – Vaccines for veterinary use).
- To allow identification of live vaccines by any suitable method instead of using an immunostaining/neutralisation test in cultures - cells or SPF eggs - with a monospecific antiserum or monoclonal antibodies.
Impact on the Ph. Eur. texts: EA testing

Deleted texts

• 2.6.24. AVIAN VIRAL VACCINES: TESTS FOR EXTRANEOUS AGENTS IN SEED LOTS
• 2.6.25. AVIAN LIVE VIRUS VACCINES: TESTS FOR EXTRANEOUS AGENTS IN BATCHES OF FINISHED PRODUCT

These chapters, which contain detailed protocols for testing extraneous agents, will be suppressed from the Ph. Eur. as of 1st of July 2020.

⇒ current methods still acceptable provided they fulfill the new requirements (see new chapter 2.6.37)

Still accessible in Ph. Eur. archives

https://pheur.edqm.eu/app/arch/search/

Impact on the Ph. Eur. texts - Revised texts

General chapter 5.2.5 (Core chapter)

"Substances of animal origin for the production of IVMPs" now "Management of extraneous agents in immunological veterinary products"

- Risk management: already described but only for substances of animal origin (e.g. Trypsin)
- Compilation of all existing requirements, collated and harmonised from other chapters and monographs
- The scope modified to cover the whole production process

- Includes ANNEX I: List of extraneous agents to be considered for the risk assessment:
  consolidates the list of avian agents already published in the Ph. Eur., includes the list of mammalian and fish agents which are currently outlined in the EMA guideline on requirements for the production and control of IVMPs (EMA/CVMP/IWP/206555/2010-Rev.1).
**Chapter 5.2.5 Annex I**

- **Non-exhaustive**: any new agent should be tested even if not on the list yet
- **Regularly reviewed** + Requests for revision supported by data

**Chapter 5.2.5 Annex II**

- Gives an example of a testing strategy decision tree for substrates which is irrespective of the target and sources species

For more information, please consult the revised version of the technical guide (available around 8th April) and in the explanatory document "What has changed and Why"

More examples to follow (EDQM training)
Impact on the Ph. Eur. texts - Revised texts

General chapter 5.2.5 new section 4-3 "Methods of detection of extraneous agents"

- Allows the use of new technologies, particularly in vitro methods such as molecular techniques
- States general prerequisites for suitable methods including, e.g. specification for positive and negative controls.
- Focuses on specific information regarding testing
- References new general chapter 2.6.37. “Principles for the detection of extraneous viruses in IVMPs using culture methods.”

General principles and examples of parameters to be taken into account to use fit-for-purpose methods (manufacturers have to check that the method is able to detect what they are looking for)

Impact on the Ph. Eur. texts - Revised texts

0062 - VACCINES FOR VETERINARY USE

5.2.4. CELL CULTURES FOR THE PRODUCTION OF VACCINES FOR VETERINARY USE

0030 - IMMUNOSERA FOR VETERINARY USE

Rearrangements in view of the new chapter 2.6.37 and the revised chapter 5.2.5.

+ Thirty-eight individual monographs

To know more: see EDQM press release
New Approach brings opportunities & benefits

• New approach based on risk assessment allows reduction in testing during manufacture and deletion of unnecessary tests for EAs on final product
• Comprehensive requirements for EA testing are centralised, Ph. Eur. texts now cover all species, this brings more clarity (no duplication, no discrepancies)
• Flexibility to choose methods for specific EA testing – fit-for-purpose sensitive techniques reflecting progress in science
• Methods no longer described in detail, building in flexibility of approach and allowing tailoring to individual product needs
• Use of state-of-the-art methods results in reduction of in vivo testing (decreased reliance on less robust methods)
• Coordinated Ph. Eur./EU approach, important also in the context of VICH
• Reduction in costs per batch

Huge thanks to everyone involved

• Experts of Group 15V and its Chairs, Céline Lorteau, Lukas Bruckner, Prof. Person
• EDQM team in charge of the secretary to Group 15V