Ordinance of the FVO 455.163
on laboratory animal husbandry, the production of genetically modified animals and the methods of animal experimentation (Animal Experimentation Ordinance)
of 12 April 2010 (as at 1 May 2010)

On the basis of Article 124 Paragraph 2, 136 Paragraph 2, Article 142 Paragraph 4 and Article 209 Paragraph 1 of the Laboratory Animal Ordinance of 23 April 2008: (TSchV), the Federal Veterinary Office (FVO) decrees the following:

Section 1: Scope

Article 1

This ordinance contains regulations on:

a. the husbandry of laboratory animals;
b. the production, breeding and husbandry of genetically modified animals and mutants that have a clinical pathological phenotype;
c. the registration and documentation of constraints and reporting procedures;
d. the definition of the degrees of severity;
e. intercantonal animal experiments;
f. applications and reports concerning laboratory animal facilities and animal experiments.

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1 SR 455.1
Section 2: Husbandry of laboratory animals

Article 2 Monitoring of laboratory animals (Article 121 TSchV)

1 Technical installations such as ventilation and automatic drinking systems shall be fitted with an alarm device if the failure or a malfunction of these systems can cause harm to the animals.

2 The condition of the animals' housing environment, especially bedding, feed and water, and also the well-being of the animals shall be checked daily.

3 The well-being of small rodents must be checked when they are transferred to clean cages. The animals must also be visually inspected at least three times a week. During the weekend, the condition of the housing environment and the well-being of the animals need not be checked if it can be shown that the animals are not adversely affected by this procedure.

4 The animals listed in Annex 3 Table 1 TSchV are considered small rodents.

5 If an animal shows signs of any constraints, these should be indicated on the enclosure or cage.

6 The frequency of checks as specified in Paragraphs 2 and 3 shall be increased according to the constraints observed.

7 A record shall be kept of the checks.

Article 3 Individual housing of incompatible animals (Article 119 Para. 2 TSchV)

A record shall be kept of the beginning and end of individual housing for incompatible animals and of special events arising during such housing.

Article 4 Run for dogs (Article 71 Para. 2 TSchV)

The run for dogs may be in an outdoor enclosure.

Article 5 Marking of small rodents (Article 120 TSchV)

1 Invasive methods such as tattoos, microchips, ear notches or amputation of toe tips may be used for marking small rodents intended for breeding.

2 For marking small rodents not intended for breeding, the use of invasive methods must be justified in the context of the specific experiment.

3 Marking with ear tags is not permitted.

4 If marking is indispensable for genotyping, the marking and biopsy must be combined
Article 6  Measures and procedures in animal rooms (Article 135 Para. 9 TSchV)

The following measures and procedures may be performed in rooms where animals are kept:

a. Marking procedures;
b. Administration of substances or food, such as brief injections or gavage
c. Taking of samples such as blood, hair, urine and saliva.

Article 7  Documentation (Article 114 TSchV)

1 The allocation and instruction of personnel taking care of laboratory animals must be recorded in a reproducible way.

2 In the animal rooms it must be clearly indicated for each animal who is responsible for compliance with animal welfare regulations.

3 Personnel must have access at all times to the documentation on constraints and on the criteria for euthanasia.

Article 8  Qualification of animal caretakers (Article 116 TSchV)

At least one third of personnel in animal caretaker positions must hold a qualification as animal attendant as specified in Article 195 TSchV.

Section 3: Production, breeding and housing of genetically modified laboratory animals and mutants that have a clinical pathological phenotype

Article 9  Approved methods for producing genetically modified animals (Article 142 Para. 4 TSchV)

1 The approved methods for producing genetically modified animals are listed in Annex 1.

2 A method may be approved if it is in widespread practical use and best ensures the well-being of the animals compared with other methods. Both the implementation of the procedures and measures and the success rate and number of surplus animals must be taken into account.

3 The approved methods shall be performed according to a standard protocol in a manner that best ensures the well-being of the animals.

4 The laboratory animal facility shall keep a record of the success rate in the use of the approved methods for the attention of the cantonal authorities. If the success rate is unsatisfactory, the laboratory animal facility must take action to improve the performance.
Article 10  Genotyping (Article 120 Para. 1 and 123 TSchV)

1 The following methods and combinations thereof are approved for genotyping in production and breeding of genetically modified animals:
   a. non-invasive methods such as the investigation of faeces or saliva;
   b. methods combined with the marking of animals;
   c. blood sampling.

2 Tail biopsies are only permitted in particular cases justified by the experiment in question. Not more than 5 mm of the tail may be removed.

3 In small rodents the following methods for combined genotyping and marking are permitted:
   a. Amputation of the distal phalanx of a toe in the first 12 days after birth; not more than two toe tips per animal may be amputated;
   b. Identification by means of ear perforation or notching after weaning.

Article 11  Phenotyping (Article 124 TSchV)

In the production and breeding of animal lines or strains, the killing of animals for anatomical or pathological purposes and investigations such as behavioural studies with mild constraint or blood sampling are permitted provided they serve to characterise the animal lines and strains. The studies shall be performed in a manner that best ensures the well-being of the animals.

Section 4: Record and documentation of constraints and reporting procedures

Article 12  Basic principles of recording constraints in small rodents (Article 124 TSchV)

1 The recording of constraints in small rodents must be documented. The following data must be entered:
   a. Results of inspections according to Annex 4;
   b. Time and date of inspections and person carrying out inspections.

2 The frequency of inspections and the traits to be observed shall be constantly adjusted based on new findings from monitoring or from animal experiments.

3 Measures for reducing constraints and criteria for euthanasia must be implemented immediately. The implementation must be documented.
Article 13  Recording constraints in small rodents (Article 124 TSchV)

1 The head of the laboratory animal facility is responsible for the recording of constraints. In particular, he or she makes sure that
   a. the persons involved in the monitoring of genetically modified lines or lines that have a clinical pathological phenotype
      1. have sufficient time at their disposal to carry out and document the inspection in a manner that best ensures the welfare of the animals,
      2. maintain state-of-the-art knowledge in the field of recording constraints,
      3. are immediately informed about new findings on clinical signs of constraint in the lines to be assessed;
   b. the basic principles stated in Article 12 are adhered to.

2 The list of traits to be checked according to Annex 4 shall be supplemented for each line with traits that can be expected or not excluded on the basis of the genetic modification.

3 Constraint records and reproduction and mortality data shall be constantly evaluated and compared with existing data on animals with the same genetic background.

Article 14  Recording constraints in new or insufficiently characterized lines of small rodents (Article 124 TSchV)

1 The new or insufficiently characterized lines of genetically modified small rodents shall be inspected for traits according to Annex 4 while changing cages and observed at least once in between cage changes.

2 Newborn animals shall be inspected for traits according to Annex 4 within the first five days and thereafter checked and observed at intervals as defined in Paragraph 1 until they are weaned.

3 During the first three generations, all animals shall be checked and observed as stipulated in Paragraphs 1 and 2.

4 If a total of 100 animals from at least three generations have been checked and no constraints have been detected, the line is deemed to be free of clinical pathological phenotype.

Article 15  Recording of constraints in small rodent lines likely to have a clinical pathological phenotype (Article 124 TSchV)

1 A small rodent line is deemed likely to have a clinical pathological phenotype if
   a. evidence of genetically related constraints are found in several animals or
   b. analysis of the data shows an increased mortality or reproduction problems.

2 In lines likely to have a clinical pathological phenotype Article 14 is applicable for the recording of constraints.
**Article 16** Recording of constraints in small rodent lines that have a clinical pathological phenotype. (Article 124 TSchV)

In lines that have a clinical pathological phenotype, the scope of the inspections and the list of traits to be studied and documented shall be stipulated in the decision according to Article 127 TSchV.

**Article 17** Provisional reporting of constraints in small rodent lines (Article 126 and 145 Para. 1a TSchV)

1 If similar constraints are found in several animals of a new or insufficiently characterized line or of a line likely to have a clinical pathological phenotype the head of the laboratory animal facility must report the constraints observed to the cantonal authorities (provisional report).

2 The provisional report must contain the following information:
   a. a precise description of the observed constraints in the summary of the constraints record;
   b. basic scientific data as specified in Annex 2;
   c. planned additional observations;
   d. planned or initiated measures to reduce constraints and their expected impact.

3 The provisional report must be submitted within two weeks of the constraints being observed.

4 If constraints are confirmed by additional recordings, the head of the laboratory animal facility must submit a definitive report as stipulated in Article 18. If the initial constraints are not confirmed, he or she must likewise report this to the authorities.

**Article 18** Definitive report of constraints in small rodent lines (Article 126 and 145 Para. 1a TSchV)

1 The definitive report of constraints in small rodent lines must be submitted at the latest when 100 animals have been checked as stipulated in Article 14.

2 The definitive report must contain the following information:
   a. basic scientific data as specified in Annex 2;
   b. specific observation plan and results of constraints record including degree of severity;
   c. measures to be taken to reduce constraints and their impact;
   d. weighing the observed constraints on the animals against the potential benefit for research, therapy or diagnostics and the likelihood of this benefit being realised;
   e. intended scope of breeding and the number of animals to be used in animal experiments.
**Article 19**

Recording of constraints in new or insufficiently characterised fish lines (Article 124 TSchV)

1 In the case of new or insufficiently characterised lines of genetically modified fish, the recording of constraints includes
   a. observation of swimming behaviour and, if possible, swarm behaviour;
   b. recording of reproduction performance;
   c. check on general health;
   d. test for clinical symptoms;
   e. test for morphological changes.

2 The reproduction data shall be constantly evaluated and compared with existing data on animals with the same genetic background.

**Article 20**

Recording of constraints in fish lines that probably have a clinical pathological phenotype (Article 124 TSchV)

1 A fish line is regarded as probably having a clinical pathological phenotype if
   a. evidence of genetically related constraints is found in several animals or
   b. analysis of the data shows increased mortality or reproduction problems.

2 In fish lines that probably have a clinical pathological phenotype, Article 19 is applicable for the recording of constraints.

**Article 21**

Recording of constraints in fish lines that have a clinical pathological phenotype (Article 124 TSchV)

In fish lines that have a clinical pathological phenotype, the scope of the inspections and the list of traits to be studied and documented shall be stipulated in the decision according to Article 127 TSchV.

**Article 22**

Reporting procedure for genetically modified fish lines that have a clinical pathological phenotype (Article 126 and 145 Para. 1a TSchV)

The report of a genetically modified fish line that has a clinical pathological phenotype comprises the information defined in Article 126 Paragraph 2 TSchV including the weighing of constraints against benefits as specified in Article 18 Paragraph 2 d of this ordinance.
Article 23

Data sheet for genetically modified lines and mutants that have a clinical pathological phenotype (Article 124 TSchV)

1 In the case of genetically modified lines and mutants that have a clinical pathological phenotype, the most important information shall be entered in a summarising document (data sheet). The data sheet contains the following information:

   a. basic scientific data as stipulated in Annex 2;
   b. summary of constraints record as stipulated in Annex 3;
   c. where applicable, the decision on lines that have a clinical pathological phenotype (Article 127 TSchV).

2 The data sheet shall be supplied to the cantonal authorities at the latest when an application for animal experiments using the line or mutant in question is submitted for approval or when a report is submitted concerning this line or these mutants.

3 It serves as a communication as stipulated in Article 13 of the containment ordinance of 25 August 1999 when genetically modified animals are moved from one enclosed system to another. When a new line that is not yet sufficiently characterised or a line that probably has a clinical pathological phenotype is passed on, all data available up to this time shall be provided with the animals.

Section 5: Definition of the degree of severity of constraints

Article 24

Categories of constraint resulting from experimental procedures or measures (Article 136 Abs. 2 TSchV)

The following four categories are used for constraints on animals resulting from procedures or measures in the context of animal experiments:

   a. Severity grade 0 – no constraint: Procedures and actions performed on animals for experimental purposes that do not inflict pain, suffering or harm on the animals, engender fear or impair their general well-being;
   b. Severity grade 1 – mild constraint: Procedures and actions performed on animals for experimental purposes that cause short-term mild pain or harm or a mild impairment of general well-being;
   c. Severity grade 2 – moderate constraint: Procedures and actions performed on animals for experimental purposes that cause short-term moderate or medium to long-term mild pain, suffering or harm, short-term moderate fear or short to medium-term severe impairment of general well-being;
   d. Severity grade 3 – severe constraint: Procedures and actions performed on animals for experimental purposes that cause medium to long-term moderate pain or severe pain, medium to long-term moderate harm or severe harm, long-term severe fear or a severe impairment of general well-being.
Article 25  Categories of constraint due to genetic modifications  
(Article 136 Abs. 2 TSchV)

The following four categories are used for constraints on animals resulting from genetic modifications:

a. Severity grade 0 – no constraint: Genetic modifications that do not inflict pain, suffering or harm on the animals, engender fear or impair their general well-being;

b. Severity grade 1 – mild constraint: Genetic modifications that cause mild pain or harm or a mild impairment of general well-being;

c. Severity grade 2 – moderate constraint: Genetic modifications that cause moderate pain, suffering or harm, fear or impairment of general well-being;

d. Severity grade 3 – severe constraint: Genetic modifications that cause severe pain, long-term suffering, severe harm, severe fear or a severe impairment of general well-being.

Article 26  Constraints to be considered for assessing the acceptability of an experiment  (Article 136 Abs. 2 TSchV)

To assess the acceptability of an experiment, consideration shall be given to the constraints defined in Articles 24 and 25 and also to further constraints imposed on the animals through debasement, through radical intervention in their appearance or their capabilities or through excessive instrumentalisation.

Section 6: Intercantonal animal experiments

Article 27  (Article 139 Abs. 2 TSchV)

1 In the reports on intercantonal animal experiments, the numbers of animals shall be separated by canton.

2 If animals change location during the experiment, they shall only be registered in the canton where the experiment mainly took place.

Section 7: Applications and reports concerning laboratory animal facilities and animal experiments

Article 28  Content of applications for approval of a laboratory animal facility  
(Article 122 TSchV)

Applications for the approval of laboratory animal facilities must contain the following information:

a. A statement of the purpose of the laboratory animal facility;

b. Animal species housed in the facility and capacity of the facility for each species;
c. Number and size of rooms, such as animal stalls, laboratory, rooms for procedures and support rooms, as well as air conditioning and lighting;
d. Housing installations, hygiene standards and access regulation, as well as cleaning standards;
e. Monitoring standards for feeding, the cages and the animal stalls;
f. Details on the origin, genetic modification, marking and husbandry of the animals;
g. Details on the method of production, breeding and husbandry of genetically modified animals or lines that have a clinical pathological phenotype;
h. Health monitoring;
i. Disposal of animal cadavers;
j. Emergency concept;
k. Name of the head of the facility and his or her deputy;
l. Number and qualification of personnel;
m. Description of animal inventory control, including the documentation of recording of constraints where applicable.

Article 29 Content of reports on laboratory animal facilities
(Article 145 Para. 1 b TSchV)

1 Reports on laboratory animal facilities must contain the following information:
   a. Number of animals born in the facility, counted at the time of weaning;
   b. Number of animals imported from abroad.

2 The numbers of animals shall be itemised by animal species.

3 Lines that have a clinical pathological phenotype shall be reported individually. Their designation must match that in the data sheet.

4 In the case of lines free of clinical pathological phenotype, the following may be summarised per species:
   a. genetically modified lines;
   b. non-genetically modified lines.
**Article 30**  
Content of applications for the approval of animal experiments  
(Article 139 Para. 1 TSchV)

Applications for the approval of animal experiments must contain the following information:

a. Species, number, gender and origin of animals that are to be used;

b. Information on the use of genetically modified animals or mutants that have a clinical pathological phenotype, including the data sheet stipulated in Article 23 of this ordinance and, where applicable, the decision stipulated in Article 127 TSchV;

c. Address of the laboratory animal facility;

d. Rooms, infrastructure and location of experiment to be conducted;

e. Listing of resource manager, study director, deputies and persons conducting the experiments;

f. Number and qualification of personnel;

g. Objective of experiment;

h. Methodology, including timetable for different parts of experiment and times for interim analysis and, where necessary, interim reports;

i. Total duration of experiment;

j. Likely impact of constraints on the condition of the animals, including degree of severity of constraints for each part of the experiment or group;

k. Monitoring of animals and criteria for euthanasia;

l. Analysis of results;

m. Justification for the experiment, the methodology, the constraint on the animals and the number of animals;

n. Weighing of constraints on animals against benefits.

**Article 31**  
Content of reports on animal experiments  
(Article 145 Para. 2 TSchV)

1 Reports on animal experiments must contain the following information:

a. Species of the animals used and, where applicable, details of the genetically modified lines or lines with a clinical pathological phenotype to which they belong;

b. Number of animals used per calendar year;

c. Origin of animals;

d. Constraint on animals;

e. Use of animals after the experiment;

f. Results and assessment of the experiment.

2 The reports shall be written for animal experiments performed and not performed.
Section 8: Commencement

Article 32

The commencement date of this ordinance is 1 May 2010.

12 April 2010  Federal Veterinary Office

Hans Wyss
Recognised methods for producing genetically modified animals

a. Crossing of genetically modified lines;
b. Pronuclear injection in mouse, rat, rabbit and guinea pig;
c. Injection and aggregation of embryonic stem cells in mouse and rat;
d. Use of viral vectors in mouse and rat;
e. Intracytoplasmic sperm injection in the mouse;
f. Injection into the cytoplasm or the yolk sac of early embryonic stages (1 to 16-cell stage) in zebrafish.
**Basic scientific data**

The following basic scientific data shall be provided on a breeding line:

a. Animal species;

b. Name of line;

c. Type of genetic modification, database reference, literature, purpose of line;

d. Producer, method of production, year of production, generation class, status of breed (discontinued, cryo-preserved);

e. Genotype, genetic background, hygiene status.
Annex 3
(Article 23 Para. 1 b)

Summary of constraints record

The following information shall be provided on the procedure and results of recording the constraints:

a. Scope of constraints record, results from mortality and reproduction data;
b. Status of constraints record (under investigation, reported, completed);
c. Description of the phenotype, assessment of the constraint, degree of severity, expression of transgene (dominant/recessive, conditional, inducible);
d. Specific needs constraint-reducing measures.
**Annex 4**
(Article 12 Para. 1 a, 13 Para. 2 and 14 Para. 1 and 2)

**Constraint record in genetically modified small rodent lines and mutants that have a clinical pathological phenotype**

Table 1: Record of constraints in genetically modified small rodent lines and mutants that have a clinical pathological phenotype

<table>
<thead>
<tr>
<th>Constraint</th>
<th>Nest inspections (Article 14 Para. 2 and 15 Para. 2)</th>
<th>Inspections during cage changing (Article 14 Para. 1 and 15 Para. 2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of young, colour, size differences</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>Food intake (milk spot)</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>General condition (size, tonus, nutritional status etc.)</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>External visible malformations</td>
<td>x</td>
<td></td>
</tr>
<tr>
<td>Dead animals, cannibalism, if necessary <em>post mortem</em> tests</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>Other noticeable problems, such as bite wounds</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>Clinical symptoms (tremor, convulsions, lameness etc.)</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>Other morphological traits according type of genetic modification</td>
<td>x</td>
<td>x</td>
</tr>
<tr>
<td>Behaviour during cage changing (apathy, nervousness etc.)</td>
<td></td>
<td>x</td>
</tr>
<tr>
<td>Nest building, condition of nest</td>
<td></td>
<td>x</td>
</tr>
<tr>
<td>Fur, eyes, body orifices</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>