Amphibian hormonal induction

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Introduction

This document describes techniques and protocols for the use of hormones. It then discusses the use of pituitaries, and provides some examples of species or species groups.

The induction of ovulation of amphibians using hormones has been used since the early 20th century. Some amphibians have never reproduced in captivity without hormones and using hormonal induction many species can be reproduced at will.

Conservation breeding programs for the flagship species, the Wyoming toad (Bufo baxteri), Boreal toad (Bufo boreas boreas), Puerto Rican crested toad (Peltophryne lemur), and Chinese giant salamander (Andrias davidianus) are either completely or largely dependent on hormonal induction. Commercial production of African clawed toads (Xenopus spp.) and also numerous other species rely on hormonal induction.

One of the great advantages of hormonal induction is the production of oocytes without fertilisation by males. These can then be fertilised by the sperm of many males to increase genetic variation in offspring, or used for Intracytoplasmic sperm injection (ICSI), nuclear transfer or other biological studies.

Several classes of compounds are used to induce amphibians - hormones as purified compounds, compounds such as pimozide, and pituitary extracts. The primary compound for the artificial ovulation of amphibians is Luteinising Hormone Releasing Hormone analogue (LHRHa). There are also two steroid hormones of use for amphibian induction, human Chorionic Gonadotrophin (hCG) and Progesterone. Other compounds of value to ovulation are dopamine blockers such as Pimozide and compounds such as oxytocins and vasotacin.

Certain types of stress can be detrimental to the maturation of ovaries and ovulation. However, stress is an essential component to the function and behaviour of amphibians and contributes to successful ovulation and spawning under natural conditions. Dopamine blockers such as Pimozide may reduce the negative effects of stress on hormonal induction in amphibians.

Pituitaries are very successful in ovulating some amphibians where purified hormones do not succeed. The disadvantage of pituitaries is that they may not be commercially available in all countries, and that they have the potential to transmit disease. Therefore, pituitaries should not be used in amphibians that are for release.

Hormones can help ovulate amphibians that are not in good reproductive condition. Nevertheless, conditioning through good husbandry, nutrition and other forms of environmental entrainment are recommended to achieve the best results from hormonal induction.

Hormones can be injected as aqueous solutions preferably as sterile SAR or Phosphate buffered saline, or applied as topic ointments in dimethyl sulfoxide (DMSO) or other carriers, or as implants. Fine needles such as 27-30 gauge and 0.5-1.0cc syringes are optimal to inject the most common volume of 100µl (0.5ml insulin syringes are particularly
useful). Some compounds and particularly progesterone require solvents such as dimethyl sulfoxide (DMSO) or alcohol to dissolve before adding to aqueous solutions.

Recommended reading for the use of hormones in the ovulation of amphibians:

Goncharov et al.1989; Browne et al. 2006 a,b.

**Which hormone to use and when**

Both theoretical and experimental evidence by 1989 showed that LHRHa was a more universal hormone for induction of oocytes from females than hCG. However, some large conservation breeding programs including have until recently used hCG (Browne et al 2006 a, b).

The tradition of using hCG with toads evolved from the use of toads for human pregnancy testing However, for the induction of sperm from male toads hCG is often preferred to LHRHa. The use of standard mammalian gonadotropins such as hCG have proved effective only in a limited number of species, and can also result in an immune response that involves the use of increasingly higher doses for repeated spawning or spermiation.

LHRHa is capable of stimulating the entire complex of reproductive processes and producing healthy offspring. Evidence shows that LHRHa is a universal stimulant to female reproduction, providing it is administered at a time when the individual has undergone gamete maturation and is ready to respond to stimulus However, there has been considerable variance in sensitivity between species, some being acutely sensitive and showing a response after a single injection of LHRHa (2mg/kg; 2µg/g) and others responding only after repeated (up to five times) application of large doses (up to 8mg/kg; 8µg/g).

Pituitary gland extracts have been widely used in the induction of frogs in the USA. The use of pituitary gland of the same or a closely related species have several shortcomings; 1) to obtain the pituitary the donors must be sacrificed, 2) it could difficult, or impossible, especially with rare species to obtain pituitary of the same or a related species, 3) even an effective preparation an incorrect level may produce negative results.

Sensitivity to hormones is a function of several factors including species specificity, reproductive condition (physiological state of gonads and neuroendocrine system). For effective induction, doses, varying response between individuals, and the number and intervals between injections may vary. The doses may need to be varied during the course of the experiment according to the behaviour of the mating pair.
Luteinizing Hormone Releasing Hormone analogue (LHRHa) Sigma L4513

Of the different types of LHRHa available Sigma L4513 (D-Ala6, des-Gly10, ethylamide LHRHa) is the only one that has been proven to be widely effective with amphibians. LHRHa vials from Sigma contain 1mg of powder that should be dissolved in 2ml of phosphate buffered saline or SAR; resulting in a concentration of 0.5mg/ml (500 µg/ml). This stock solution can then be stored indefinitely in Eppendorf tubes at -80°C in 100 µl aliquots, which will give one dose of 50µg for example at 2 µg/g for a 25g male toad.

A milligram (mg) is one thousandth of a gram (1 x 10^-3g), a microgram (µg) is one millionth of a gram (1 x 10^-6g), and a picogram (pg) is one billionth of a gram (1 x 10^-9g).

Human Chorionic Gonadotropin (hCG) Sigma C1063, Intervet Inc. Choluron®

Human Chorionic Gonadotropin (hCG) comes as a powder and must be dissolved into an aqueous solution before administration by injection. The most convenient units are vials at 2500 or 5000 International Units (IU). Store hCG in the fridge at 4°C for short periods or frozen for months at -80ºC.

Table 2. Standard vial amount from *Sigma hCG; #Intervet Inc.Choluron®. IU = international units. The volumes are products of 100µl plus 20, 40, or 80µl for vial loss.

<table>
<thead>
<tr>
<th>Vial (IU)</th>
<th>Water (µL)</th>
<th>No. 250 IU doses</th>
<th>IU/100µL</th>
</tr>
</thead>
<tbody>
<tr>
<td>*2500</td>
<td>1020</td>
<td>10</td>
<td>250</td>
</tr>
<tr>
<td>*5000</td>
<td>2040</td>
<td>20</td>
<td>250</td>
</tr>
<tr>
<td>#10000</td>
<td>4080</td>
<td>40</td>
<td>250</td>
</tr>
</tbody>
</table>

Priming

Priming is useful in preparing females for the final induction of oocytes especially where the state of ‘Reproductive conditioning’ is uncertain. Typical priming doses are 10% of the doses used to finally induce ovulation. For extended priming administer LHRHa ten days and three days before the final ovulatory dose. Priming administered about 24hrs before administration to bring oocytes to stage 6 may also be advantageous even with well conditioned females. However, females in optimal reproductive condition may be very sensitive to hormones. In these cases priming should not be used as even very low doses of hormones may produce ovulation.

When the fertilisation of oocytes is critical, males may be induced about 12hrs after the administration of priming doses and placed with females in spawning tanks.

Typical induction protocol for females

From an overview of the traditional and recent studies the optimal dosage of hormones for ovulation is 2-8 µg/g. So for a 50 g toad is 60µg LHRHa, or 20µg LHRHa with 500IU hCG could be used (IU, International Units). Progesterone used with LHRHa or hCG offers considerable potential for improving traditional methods (see Progesterone this doc). Females typically ovulate about 12-24hrs after an ovulatory dose of hormones. However, this varies with factors including species, female age, reproductive condition, and temperature.
Table 1. The different doses of hormones and the amount per gram of amphibian weight in a study of the Fowler toad (*Bufo fowleri*; Browne et al. 2006b).

<table>
<thead>
<tr>
<th>Compound</th>
<th>Administered dose</th>
<th>Amount per gram of toad weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>LHRHa</td>
<td>4 μg</td>
<td>0.12 μg</td>
</tr>
<tr>
<td>LHRHa</td>
<td>20 μg</td>
<td>0.61 μg</td>
</tr>
<tr>
<td>LHRHa</td>
<td>60 μg</td>
<td>1.8 μg</td>
</tr>
<tr>
<td>hCG</td>
<td>500 IU hCG</td>
<td>10 IU</td>
</tr>
</tbody>
</table>

hCG alone: A recommended ovulatory dose for females is 10IU hCG per gram. It simplifies the process to standardize the volume for storage to 100μl (Table 1). For medium sizes toads this approximates to 500IU for a 50g female (200 μl). Males are smaller and only need 300 IU (Browne et al. 2006a,b).

For smaller amphibians adjust the dose proportionately to body weight within a volume of 50-100μl. For instance for small 5g female dilute the 500IU/100ml dose into 900ml and administer 100μl.

Injections should be intraperitoneal or through the dorsal lymph sac (see Handling and injecting amphibians). Once females receive the final ovulating dose spawned oocytes may be fertilised by induced males or females can spawn into Simplified Amphibian Ringers for *in vitro* fertilisation (*In vitro fertilisation*). If the oocytes are to directly fertilised induced males should be introduced to the females (see below). The pairs should be placed in an appropriate depth of water. Be careful to have a solid place for females to rest otherwise they can drown when amplexing.

**Typical induction protocol for males**

With toads inject 35g males with 300 IU of hCG preferably also with 20μg LHRHa. Males produce sperm within about three hours and sperm production lasts about 8-12 hrs. Because the female begins laying eggs around 10-24 hours after her injection, a synchronized injection scheme is important to optimize fertilization (Browne et al. 2006a,b).

**Progesterone P 8783**

Progesterone P 8783 comes in a 1g bottle of powder. Progesterone has a poor solubility in aqueous solution. At 25°C, the maximum solubility is about 7μg/ml.

In post spawning Fowler toads (*B. fowleri*) the use LHRHa and hCG in various combinations with progesterone showed that only treatments with progesterone produced ovulation. Overall, results suggest that progesterone 100 μl at 7μg/ml with a dose between 20μg and 60μg of LHRHa may be optimal for the induction of ovulation in toads with poorly developed stage 6 oocytes.


**Pimozide Sigma P 1793**

Pimozide blocks estradiol produced by the gonads preventing the release of luteinising hormone from the pituitary. Hence, pimozide circumvents the need for environmental entrainment to mature and ovulate amphibians.
Pimozide (Sigma P 1793) comes in 500 mg bottle of powder. Pimozide can be dissolved in dimethyl sulphoxide (Me₂SO; Sigma D 2438) at 18mg/ml.

The use of pimozide with amphibians is at an early experimental stage. Browne et al. 2006 with the Fowler toad *B. fowleri* showed no effect of pimozide on improving ovulation. However, pimozide had proved very successful with some fish species. Optimal doses for carp *Cyprio carpio* are 10 µg/kg LHRHa with 5 mg/kg pimozide. Indian catfish *Heteropneustes fossilis* can be induced to spawn by LHRHa alone in a dosage range of 0.15-0.2 mg/kg body weight or in combination with pimozide at a very low dosage of 0.05 mg/kg.

**Examples**

**African Clawed Toads Xenopus spp.**

Typically *X. laevis* are induced by a priming dose of 10IU hCG 12-72 hrs before the ovulatory dose of 100 - 200 IU hCG. A priming dose of 20 IU can ovulate. With older females 100 IU produces ovulation after 6-8 hours, whereas 200 IU produces ovulation after 4 hours. For fertilisation through amplexus in the evening only the ovulatory dose of 200 IU hCG for females is given and 100 IU hCG for males. High temperatures of 30ºC will prevent mating and low temperatures of 16ºC cause developmental abnormalities.

**Pituitaries (with the Northern Leopard frog Rana pipiens)**

Hypophysation in its strictest sense is the process of inducing breeding through injection of pituitary extract from pituitary glands of amphibians of the same species. Anuran pituitaries as a dried powder can be purchased from supply houses or can be sampled directly from frogs or toads. The advantage of commercial pituitaries is that they may be obtained aseasonally, and that their purchase saves the trouble of harvesting them. The disadvantage of commercial pituitaries is that interspecific differences in their efficacy in not well known, and that pituitaries sampled from a local population have less chance of spreading disease. *Rana pipiens* pituitaries and ovulation kits can be obtained commercially.

The earlier before the normal spawning time the greater number of pituitaries and amount of progesterone are needed to induce ovulation (Table 2).

Table 2. The doses of progesterone when combined with pituitaries use to ovulate *Rana pipiens*. NB: Doses shown are calculated as female pituitaries; two male pituitaries are equivalent to one female pituitary. Both the pituitary and progesterone should be injected into the coelomic cavity about 48 hours before scheduled fertilization.

<table>
<thead>
<tr>
<th>Month</th>
<th>Pituitaries Alone</th>
<th>OR Pituitaries</th>
<th>+ Progesterone (mg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sept-Oct</td>
<td>10-12</td>
<td>2</td>
<td>5.0</td>
</tr>
<tr>
<td>Nov-Dec</td>
<td>6-8</td>
<td>2</td>
<td>2.5</td>
</tr>
<tr>
<td>Jan-Feb</td>
<td>4-5</td>
<td>1</td>
<td>2.5</td>
</tr>
<tr>
<td>Mar-Apr</td>
<td>1</td>
<td>1</td>
<td>-</td>
</tr>
</tbody>
</table>
The following description is based on *R. pipiens*, whose ovulation has been most routinely conducted under laboratory conditions. However, pituitaries are readily collected following the technique described by Rugh (1965) and used immediately or stored frozen for days.

**Breeding with amplexus**

If breeding is to be accomplished by amplexus, both male and female should be treated. Follow the above table for pituitary dosage for females. Males will respond to half the pituitary dose; progesterone is not needed. The injected animals should be placed in an aquarium with 200-250 mm (8-10 in.) of dechlorinated water at 20-22°C. The animals should be placed above plastic screening with a coarse mesh 9.5 mm approximately 50 mm (2 in.) above the floor of the aquarium to protect the egg masses. Depending on the season, fertile eggs should be available in 1848h if the aquarium is placed in an undisturbed area. Note that the precise time of fertilization is not known when this technique is used.

**Hormones inducing salamanders**

Spanish newt, *Pleurodeles waltl* embryos were obtained either from Xenopus Ltd (UK) or from breeding pairs maintained in the laboratory. In either case, females were induced to spawn by subcutaneous injections of 1 mg/ml of LHRHa (Sigma L4513) at 0.1mg/kg body weight (0.1µg/g). Up to 400 fertile eggs could be obtained from a single female during the breeding season (November-March) (Fekete and Brockes 1988).

**Hormones inducing Puerto Rican crested toad (*Peltophryne lemur*)**

LHRHa is used for females and hCG for males. LHRHa is diluted to make a 100 microgram/ml solution and frozen in 1 ml syringes prior to use. The shelf life of the dry product is 2 years. Toads are injected using a dose of 0.1 µg/g subcutaneously (intraperitoneal injection can also be used).

Although there was no difference in peak sperm concentrations between LHRH and hCG, we still recommend using hCG because of the timing of the spermiation and an enhanced behavioural response. The timing of the peak appears to be sooner with hCG (3 to 6 hours) compared to LHRH (24 hours). The hCG dose that we recommend for males is 10 IU/g subcutaneously (Lentini 2007).

**USSR program for breeding amphibians**

In 1983 the USSR established a working group to provide technologies for the conservation of amphibians (Goncharov et al 1989).

1. to develop methods for maintaining amphibian species of interest;
2. to develop methods for obtaining healthy offspring in captivity;
3. to develop the most economical methods for egg incubation and rearing, that is, the maximum yields for the minimum expense;
4. to carry out practical measures for breeding and rearing of sufficient numbers of animals to ensure captive conservation of the chosen species;
5. to carry out practical measures for the establishment of new populations in nature based on animals bred and reared in captivity.

Goncharov et al. (1989) found there was considerable variance in sensitivity between species, some being acutely sensitive and showing a response after a single injection of LHRHa (0.2 mg/kg) and others responding only after repeated (up to five times) application of large doses (up to 0.8mg/kg).
In this program 37 species in 17 genera were successfully induced with LHRHa:

*Onychodactylus fisheri*
*Salamandrella keyserlingii*
*Mertensiella caucasica*
*Triturus alpestris, T. cristatus, T. helveticus, T. karelini, T. montandoni, T. vittatus, T. vulgaris*
*Bufo bufo, B. calamita, B. danatensis, B. marinus, B. melanostictus, B. viridus*
*Bombina bombina, Bo. maxima, Bo.orientalis, Bo.variegata*
*Hyla arborea, H. japonica, H. savignyi*
*Litoria caerulea*
*Osteopilus septentrionalis*
*Ceratophrys cranwelli, C. ornate*
*Discophus antongilli*
*Kaloula pulchra*
*Microhyla ornata, M. pulchra*
*Pelobates fuscus, P. syriacus*
*Pelodytes caucasicus*
*Xenopus borealus, X. laevis*
*Rana adspersa, R. macrocnemus, R. nicobariensis, R. pipiens, R. temporaria*

**References:**


