Development of homologous radioimmunoassays for equine growth hormone and equine prolactin and their application to the detection of circulating levels of hormone in horse plasma

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Summary — Highly purified and well-characterised preparations of equine prolactin and growth hormone from equine pituitary glands were employed to set up highly sensitive and specific homologous radioimmunoassays (RIA) for the measurement of hormone in horse plasma. The limit of sensitivity of the GH RIA was 1.2 ng/ml with mean intra- and inter-assay coefficients of variation (CV) of 6.6 and 10%, respectively. The sensitivity of the equine prolactin (ePRL) RIA was 0.5 ng/ml with mean intra and inter-assay CV of 9.1 and 15.6%, respectively. Dose–response curves of a crude pituitary gland extract and plasma samples collected from a mare and foal were parallel to the standards and the PRL RIA was clinically validated by administration of thyrotropin-releasing hormone (TRH). Plasma samples taken at 15 min intervals over 24 h from lactating mares gave 24 h mean GH values in the range 5.5 to 7.95 ng/ml. Large intermittent elevations of GH activity were detected. The mean 24 h PRL concentrations were between 3.2–10.4 ng/ml in the lactating animals, with higher concentrations earlier in lactation. Long episodic bursts of PRL were detected.

equine / purification / characterisation / radioimmunoassay / equine prolactin / equine growth hormone

Résumé — Mise au point de tests radio-immunologiques homologues pour la détection de l'hormone de croissance et de la prolactine équines et leur application pour la détection des concentrations d'hormones circulantes dans le plasma de cheval. La prolactine (PRL) et l'hormone de croissance (GH) ont été purifiées à partir d'hypophyses de cheval. Ces préparations d'hormones ont permis de préparer des anticorps spécifiques et de mettre au point des tests de RIA homologues. La limite de détection du RIA de l'hormone de croissance est de 1.2 ng/ml, avec des moyennes de coefficients de variations (CV) inter- et intra-tests de 6,6% et 10% respectivement. La sensibilité du RIA et

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de la PRL est de 0,5 ng/ml avec une moyenne de CV inter- et intra-tests de respectivement 9,1% et 15,6%. Les courbes d’effet-dose d’extrait brut d’hypophyse et d’échantillons de plasma prélevés sur une jument et un poulain sont parallèles aux courbes standard et le RIA de la PRL a été validé cliniquement par injection de la «thyrotropin releasing hormone» (TRH). Des niveaux de GH à 24 h de 5,5 à 7,5 ng/ml de GH ont été détectés dans des échantillons de plasma de juments allaitantes prélevés à 15 min d’intervalle sur une période de 24 h. Des variations élevées, courtes et intermittentes de GH ont été détectées. Les concentrations moyennes à 24 h de PRL se situent entre 3,2 et 10,4 ng/ml pendant la lactation, avec des niveaux plus élevés dans les phases précoces de lactation avec de longs pics épisodiques.

équin / purification / particularité / dosage radio-immunologique / prolactine / hormone de croissance

INTRODUCTION

Prolactin (PRL) and growth hormone (GH) are protein hormones produced in the anterior pituitary gland, which are involved either directly or indirectly in growth and lactation in all mammalian species. Although equine GH (eGH) has been purified (Saxena and Henneman, 1966; Hartree et al, 1968; Conde et al, 1973) and antisera have been raised in rabbits (Poskus et al, 1976) inadequate supplies of the hormone have precluded the development of a homologous radioimmunoassay (RIA) for its measurement. A role for eGH in the anabolic process of foal growth is presumed and a galactopoietic role similar to that in ruminants has been suggested (Worthy et al, 1986). The purification of equine PRL (ePRL) has been previously described (Chen et al, 1979; Li and Chung, 1983) and there are several RIAs available for its measurement (Roser et al, 1984; Johnson, 1986; Thompson et al, 1986a; Worthy et al, 1986). Prolactin is likely to control the growth and function of the mammary gland during a normal gestation and lactation (Worthy et al, 1986) and may also be involved in ovarian function in the mare (Roser et al, 1987). It has also been suggested that PRL may be involved in infertility post partum (Worthy et al, 1986).

MATERIALS AND METHODS

Purchased supplies

Sodium \textsuperscript{125}I-iodide and \textsuperscript{35}S were purchased from Amersham PLC UK; Iodogen, Pierce Laboratory Rockford, IL, USA; ampholine PAG plates, diethylaminoethyl (DEAE) sephacel, phenyl sepharose CL-4B, sephadex G-100, and molecular weight markers were purchased from Pharmacia, Milton Keynes UK; tissue culture media, Gibco Biocult, Paisley, Scotland; JB-4 resin kit, Polysciences Inc, Warrington, PA, USA; thyrotropin-releasing hormone, bovine serum albumin, adjuvants and reagents for PAGE, Sigma Chemical Co, Poole, Dorset UK; lithium heparin blood sampling tubes (100 x 16 mm), intravenous cannulae (Branula G-14 and vygon mosquitos 123 G-13, 70 mm), Biovet, Mullingar, Rep of Ireland; donkey anti-guinea-pig gamma globulin, Guildhay Antisera Ltd, Guildford, Surrey, UK.

Hormones and reagents

Ovine PRL (NIAMDD-oPRL-14, 31 iu/mg), bovine PRL (NIAMDD-bPRL-6, 30 iu/mg), ovine GH (NIH-oGH-S-11, 0.56 iu/mg), bovine GH (NIH bGH-B-18, 0.81 iu/mg), rat PRL (NIDDK-rPRL-B-6,2 iu/mg), rat GH (NIDDK-rGH-B-12, 1.8 iu/mg) were obtained from the NIAMDD, Baltimore, MA, USA; porcine PRL (USDA pPRL-B-1, 34 iu/mg) and porcine GH (pGH-B-1, 1.5 iu/mg), were obtained from D Bolt at the USDA, Beltsville, MA, USA; human GH (MRC 1st International Standard for immunoassay B 66/217). The equine gonadotrophin fraction was obtained from M Kelly, Faculty of Agriculture, University College Dublin.
and was a 150 mM ammonium sulphate, pH 4 extract from equine pituitary glands. The equine pituitary extracts were alkaline extracts (100 mM ammonium acetate, pH 8.5) dialyzed and lyophilized from whole equine pituitary glands.

**Animals**

Pigeons of mixed breed and sex (adults, aged over 2 years and weighing between 298 and 537 g) were obtained from the Department of Psychology, Thornfield, University College Dublin. They were housed under uniform temperature conditions (25–26°C) in a room artificially illuminated during the normal daylight hours. They were kept for at least 1 week before being used in assays and were caged in pairs and fed pigeon chow and water ad libitum. Albino Schofield mice were obtained from Trinity College Dublin and were maintained as a randomly bred colony for at least 9 generations before experimental use. Snell dwarf mice were obtained from AT Holder and were maintained as a breeding colony at the animal facility at the NIRD Shinfield, Reading.

**Tissue sources**

Pituitary glands were collected from horses of mixed breed (Irish horse abattoir, Straffan, Co Kildare) and were frozen immediately in liquid nitrogen and stored frozen at -20°C until use. Livers and mammary glands were collected from rabbits (Western Laboratories Ltd) and mares (Irish horse abattoir) and were kept at -20°C for up to 3 months before preparation of microsomes for radioreceptor assays (RRAs).

**Extraction of GH and PRL from equine pituitary glands**

**Growth hormone**

Extraction of the hormone from horse pituitary glands was by the general extraction procedure for pituitary hormones (Licht et al, 1977). Pituitary glands were homogenized in 5 volumes 25 mM ammonium bicarbonate, pH 9.0 and extracted for 2 h at 4°C. Following centrifugation (10 000 g for 15 min) the residue was re-extracted as above and the pellet discarded. The pooled supernatants were brought to 0.15 M ammonium sulphate, adjusted to pH 4.0 and stirred for 1 h at 4°C. After centrifugation as described above the supernatant was discarded and the residue containing the GH (Farmer et al, 1975), PRL and adrenocortico-trophic hormone was resuspended in 50 mM ammonium acetate (155 ml) pH 9.5.

**Prolactin**

Equine PRL was prepared from horse pituitary glands by the method of Li and Chung (1983) (with modifications). Acid-acetone powder (9.5 g) was extracted overnight in 0.1 M ammonium acetate, pH 9.5. The insoluble material was removed by centrifugation at 9 500 g for 20 min and the supernatant was adjusted to pH 5.7. The isoelectric precipitate was recovered by centrifugation and dissolved in 25 mM ammonium acetate, pH 9.5 (150 ml). Aliquots (50 ml) were stored frozen at -20°C.

**Electrophoretic characterisation**

SDS polyacrylamide gel electrophoresis was carried out as described previously (Laemmli, 1970). Discontinuous polyacrylamide gel electrophoresis was carried out as described by Davis (1964) Isoelectrofocusing was carried out on ready-made acrylamide gels (5% acrylamide, 3% bis-acrylamide) in the pH range 3.5–9.5. Hormone samples (20 μg) in 1% glycine were focused at 10°C for 1.5 h at 30 W, 1 500 V and 50 mA (LKB 2103 power supply settings). The pH gradient was determined using a surface pH electrode.

**Bioactivity**

Standard ovine PRL (oPRL) and bovine GH (bGH) RRAs were set up to characterise the binding activity of ePRL and eGH to lactogenic and somatogenic receptors. Microsomal membranes from lactating rabbit mammary gland, pregnant mare mammary gland and pregnant rabbit liver (Shiu et al, 1973; Tsushima and Friesen, 1973) and plasma membranes from rabbit and mare liver were used (Parke and Forsyth, 1975). Mouse mammary gland bioassay was carried out as described by Hayden et al, 1991, and the local or micro method of pigeon crop sac bioassay was
used (Nicoll, 1967). The Snell dwarf mouse bioassay was carried out as described by Holder et al. (1980).

Immunoactivity

The crossreactivity of eGH was examined in an homologous pGH RIA (Buttle, 1987a). The crossreactivity of ePRL was examined in an homologous ePRL RIA (Worthy et al., 1986) and in an homologous pPRL RIA (Buttle, 1987b).

Assay development and validation

Radioiodinations

Equine GH and ePRL (5 μg) were radiolabelled with Na $^{125}$I (0.5 mCi) by the iodogen method (Salacinski et al., 1981). Labelled hormone was separated from free iodine by gel filtration chromatography on Sephadex G-100 previously coated with a solution of 0.5% BSA in RIA buffer. The specific activity, estimated by self-displacement analysis (Abdul-Ahad, 1984) was 36 ± 7.9 (sd) μCi/μg (n = 6) for ePRL and 41.5 ± 10 (sd) μCi/μg (n = 5) for eGH.

Growth hormone

Antiseras were raised in 2, 4-month-old Duncan Hartley strain guinea pigs with 3, monthly, subcutaneous injections of 750–950 μg of eGH each, emulsified in Freunds incomplete adjuvant. A final boost of 215 μg was given and the animals were bled by cardiac puncture 2 weeks later. Titre studies indicated that the dilution of antibody that bound 60% of the $^{125}$I-eGH in the absence of competitor was 1:10 000 (final). One set of antisera was of higher affinity and this was employed in the RIA. The antiserum was used at a final dilution of 1:10 000. All dilutions were made up in RIA buffer (50 mM sodium phosphate, pH 7.5, containing 0.5% BSA and 150 mM NaN$_3$). The assay tubes contained dilutions of standard eGH (0.98–250 ng/ml in 100 μl), or of plasma sample (1:2 and 1:4), antiserum (100 μl) and $^{125}$I-eGH (20 000 cpm in 50 μl). The antibody was allowed to preincubate with the standard for 48 h before addition of the tracer and incubation overnight at 4°C. This was followed by the addition of normal guinea-pig serum (50 μl, diluted 1:700) and donkey anti-guinea-pig gamma globulin (50 μl, diluted 1:56). After incubation for a further 24 h, separation of the precipitated (bound) and free radioactive activity was achieved by adding 1 ml of a cold 4% solution of polyethylene glycol in phosphate-buffered saline and centrifugation at 2 500 g for 30 min at 4°C. The supernatant fraction was drained and the antibody bound hormone in the precipitate was counted on an automatic 1260 multigamma 11 counter (74% efficiency).

Administration of hGRF(1–29)

Three Dutch warm-blooded mares between 8 and 18 years of age and weighing between 463 and 546 kg were used. The animals were used in a 2-factor cross over design. One horse was injected with 670 μg hGRF(1–29) (Kabivitrum r, Stockholm, Sweden) dissolved in 5 ml saline intravenously and the other 2 were given the equivalent volume of saline. The next day each horse was given the opposite treatment. Blood samples were collected from the left jugular vein via an indwelling catheter at various times before and after administration of saline or hGRF(1–29).

Effect of fasting on GH concentrations

Six Dutch warm-blooded horses (3 mares and 3 geldings) between 9 and 12 years of age and weighing between 580–634 kg were used. Blood samples (5 ml) were collected from the jugular vein by venepuncture at 10.00, 10.30, 11.00, 11.30 and 12.00 h prior to fasting and at the same time points after 16 h fasting.

Prolactin

Antiseras to ePRL were raised in 2, 3-month-old New Zealand White rabbits using 3 injections of approximately 350 μg ePRL at 2–3 week intervals, in Freund's complete adjuvant for the first injection and incomplete adjuvant subsequently. The antiserum bound between 40 and 60% of $^{125}$I-ePRL at a final dilution of 1:5 000 and further immunizations failed to boost greater responses. Both antisera revealed similar specificities and affinities and they were pooled.

All assay reagents were made up in RIA buffer. The assay tubes contained dilutions of standard ePRL (0.48–250 ng/ml in 100 μl) or plasma samples (1:2 and 1:4), antiserum (100 μl) and $^{125}$I-ePRL (20 000 cpm in 50 μl). The sensi-
tivity of the assay was increased by preincubating the standard with antibody for 72 h before addition of tracer. Incubation was at 4°C and was followed by the addition of normal rabbit serum (50 µl, diluted 1:700) and sheep anti-rabbit gamma globulin (50 µl, diluted 1:28). The assay was completed as described above.

Administration of TRH

To determine if the PRL RIA detected changes in PRL concentrations in plasma in response to TRH, blood samples were taken from each of 3 animals (a non-lactating mare, a lactating mare and a stallion) via indwelling jugular catheters implanted 24 h before TRH injection (2.34 mg in 1 ml of sterile saline, administered intravenously). A lactating mare received an injection of saline only. Blood was collected at 15 min intervals from -60 to +165 min relative to injection.

Twenty-four-hour profiles of growth hormone and prolactin

Blood samples were taken at 15 min intervals for at least 24 h into heparinized tubes from an indwelling jugular catheter implanted 24 h before sampling. Two Connemara ponies (#101, age unknown, and #105, age 14 years) and 3 cross-breeds (throughbred x Irish draught, #102, #103 and #104) ranging in age from 9 to 20 years. Mares #101, #102 and #103 were lactating. Sampling times were as follows: (1) #101, week 1 of lactation from 6-7 March (starting at 06.45 h on d1 to 07.15 h on d2), week 3, from 22-23 March (06.30 h on d1 to 07.00 h on d2) and week 6, from 16-17 April (06.45 h on d1 to 07.15 h on d2); (2) #102, week 1 of lactation, 23-24 May (10.15 h on d1 to 10.30 h on d2) and week 8, 14-15 July (10.15 h on d1 to 10.30 h on d2); and (3) #103, week 2 of lactation (sampled in May as described). Horse #104 was a non-lactating mare and #105 was a stallion (sampled in mid-July as described). Plasma was collected by centrifugation at 1500 g for 20 min and stored at -20°C until assay. Animals were subjected to the natural photoperiod, temperatures and feeding regimes throughout the period of study.

Statistical analysis

RRA and RIA dose–response curves were analyzed using the computerized, non-linear, least-squares curve fitting routine (Allfit) of De Lean et al (1978). Potency estimates of the hormones were obtained using the ED50 values (50% maximally effective dose). In the mouse mammary gland bioassay, the relationship between the lactogenic hormone concentration of the medium and the secretory grades of the explants were evaluated by linear regression analysis and by the Bonferroni t-test (Gill, 1986) to determine if hormonally stimulated effects were significant. Validity tests were carried out on the results of the pigeon crop sac bioassay using the methods for the 6-point parallel line assay of Wardlaw (1985). Potency estimates and 95% confidence limits were made using the method of Finney (1978) employing slope and variance in the final calculations. Results of the assay of plasma samples were calculated using the RIA program of the 1260 multit gamma 2 counter based upon cpm versus log concentration. A paired t-test and a sign test were used to determine the significance of the timing or the GH concentration of the GH peak in the GRF challenge test. The Hotelling's T² statistic was used to determine if GH concentrations differed with feeding status (Morrison, 1967; Rosner, 1990).

RESULTS

Isolation of equine GH and equine PRL

Growth hormone

Table I summarises the steps involved in the purification of eGH. The initial pH 9 extract had quite a low specific activity relative to bGH in the RRA. This was followed by an ammonium sulphate-assisted acid precipitation of the GH resulting in a 5-fold increase in the purity of eGH and 50% recovery of GH-like activity. The specific activity of this fraction relative to bGH was 0.27. Growth hormone eluted from a sephadex G-100 column with a KaV of 0.416 and a Vₑ/Vₒ ratio of 2.25. Gel filtration resulted in a 2.3-fold purification over the previous step and a specific activity of 0.57. The elution profile of eGH from phenyl sepharose was rather broad, eluting
between 90 mM and 38 mM ammonium acetate (fig 1A). Active fractions were pooled and corresponded to 42% of the GH applied, a 1.2-fold increase in purity over the previous step and a specific activity of 0.66. Ion exchange chromatography was highly efficient in separating the PRL from the GH. Growth hormone was eluted in 2 broad bands, one was unadsorbed by the gel and was free of equine PRL (fig 1B) and the other was eluted between 0.06 M and 1.45 M sodium chloride. Only the breakthrough fractions were pooled and contained 88% of the GH activity applied with a specific activity of 1 and an overall yield of 0.5 mg GH per g of fresh pituitary glands.

**Prolactin**

Table II summarises the purification of ePRL. Substantial losses of up to 65% of oPRL-like RRA activity were detected after isoelectric precipitation of ePRL. This was prevented in later runs by standing the extract for up to 48 h at 4°C and reharvesting the precipitate. On a sephadex-G 100 column ePRL was eluted as a single peak with an elution volume 2.1 times that of the void volume \( V_e / V_0 = 2.1 \) corresponding to a KaV of 0.4 (fig 1B) resulting in a 2.5-fold increase in purity and over 60% recovery relative to the preceding step. Equine PRL eluted from the ion exchanger at 0.1 M sodium chloride and yielded almost 90% of the applied hormone. An increase of 4-fold in purity over the product of the previous step was achieved, giving an overall purity of 10-fold. The low potency of the final preparation of ePRL relative to oPRL in all of the lactogenic RRAs (< 2 iu/mg) precluded their use as quantitative assays. The rabbit mammary gland RRA was therefore used.

**Table I. Summary of purification of eGH from a 50 g wet weight of equine pituitary glands.**

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Total bGH equiv (mg)+</th>
<th>Protein (mg)</th>
<th>Specific activity (mg/mg)+</th>
<th>Purification (fold)</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH 9 ammonium bicarbonate extract</td>
<td>259.8</td>
<td>5929</td>
<td>0.043</td>
<td>5.58 (5.6)</td>
</tr>
<tr>
<td>pH 4 ammonium sulphate precipitate</td>
<td>130.7 (50.3)</td>
<td>546</td>
<td>0.240</td>
<td>13.27 (2.4)</td>
</tr>
<tr>
<td>Post-gel filtration</td>
<td>75.4 (57.7)</td>
<td>132.3</td>
<td>0.571</td>
<td>15.44 (1.2)</td>
</tr>
<tr>
<td>Post-hydrophobic interaction chromatography</td>
<td>31.9 (42.3)</td>
<td>48</td>
<td>0.66</td>
<td>10.58 (1.7)</td>
</tr>
<tr>
<td>Post-anion exchange chromatography</td>
<td>28.0 (87.9)</td>
<td>25.5</td>
<td>1.1</td>
<td>25.58 (1.7)</td>
</tr>
</tbody>
</table>

Values in parentheses are relative to the preceding step; +: by radioreceptor assay.

**Table II. Summary of purification of ePRL from a 500 g wet-weight batch of equine pituitary glands.**

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Total ovine PRL equiv (mg)+</th>
<th>Protein (mg)</th>
<th>Specific activity (mg/mg)+</th>
<th>Purification (fold)</th>
</tr>
</thead>
<tbody>
<tr>
<td>pH 5 isoelectric precipitate</td>
<td>0.53</td>
<td>95</td>
<td>0.006</td>
<td>1.85 (2.20)</td>
</tr>
<tr>
<td>Post-gel filtration</td>
<td>0.32 (60.4)</td>
<td>42</td>
<td>0.008</td>
<td>2.54 (1.37)</td>
</tr>
<tr>
<td>Post-ion exchange</td>
<td>0.28 (87.5)</td>
<td>9</td>
<td>0.031</td>
<td>10.37 (4.08)</td>
</tr>
</tbody>
</table>

* Values in parentheses are relative to the preceding step; +: by radioreceptor assay.
Fig 1. Purification of eGH (A) and ePRL (B). (A) Elution of crude eGH from a column of Sephadex G-100 (5 x 63 cm) with 25 mM ammonium acetate pH 8.5 (top). Fractions of 15 ml were collected at a flow rate of 1 ml/min. Fractions containing eGH activity were pooled and dialyzed overnight in 100 mM ammonium acetate, pH 8.5 and applied to a column of phenyl Sepharose CL-4B (1.5 x 8 cm) in 100 mM ammonium acetate, pH 8.5 (middle). Elution was achieved using a linear gradient of acetonitrile (0–30% V/V) and ammonium acetate from 100 to 10 mM. Active fractions were pooled and applied to a column of DEAE-Sephacel (1 x 4 cm) equilibrated with 25 mM ammonium acetate pH 8.5 containing 20%(V/V) acetonitrile (bottom). After sample application at A non-adsorbed proteins were eluted with starting buffer at B and adsorbed material was eluted in a gradient from 25 to 200 mM NaCl at C. Fractions of 11 ml were collected at a flow rate of 0.25 ml/min. Equine GH was recovered in the breakthrough fractions and was free of ePRL (which was eluted at 100 mM NaCl in a gradient from 25–200 mM NaCl) (B) Elution of crude ePRL from a column of Sephadex G-100 (5 x 76 cm) with 25 mM ammonium acetate, pH 9.5 (top). Fractions of 20 ml were collected at a flow rate of 1 ml/min. Fractions containing PRL were pooled and applied to a column of DEAE-Sephacel (2.5 x 8.5 cm) (bottom). The column was equilibrated with starting buffer at A (25 mM ammonium acetate, pH 8.5, 15% acetonitrile) and fractions of 11 ml were collected at a flow rate of 0.25 ml/min. Non-adsorbed protein was eluted with starting buffer at B, followed by step-elution of ePRL with 50 mM NaCl at C and 100 mM NaCl at D. Fractions were monitored for protein by absorbance at 280 nm ( ), for PRL activity by the rabbit mammary gland RRA (O) and for GH activity in a rabbit liver RRA (●).
to locate, but not quantify, ePRL. The overall yield of ePRL was 10 mg from 500 g of pituitary glands.

**Electrophoretic characterisation**

On a non-denaturing gel eGH was resolved into 3 major bands with relative mobilities (Rf) of 0.38, 0.47 and 0.58 and 1 minor band with an Rf of 0.65. Equine PRL was resolved into 3 major bands with Rfs of 0.65, 0.73 and 0.81 and 2 minor bands with Rfs of 0.58 and 0.61. Ovine PRL, which was run in parallel, was resolved into 2 major bands with Rfs of 0.65 and 0.73. (fig 2A). On a 12.5% SDS gel, eGH was resolved into 4 protein bands with molecular weights of 22 500 Da for the major band and 19 000, 16 500 and 15 200 Da for the 3 minor components (which appeared only after several months of storage). Equine PRL migrated as 2 protein bands with a major band of 26 000 Da and a minor band of 29 000 Da. Similarly oPRL was resolved into 2 protein bands of 25 000 and 29 000 Da (fig 2B). Using broad range isoelectrofocusing, ePRL was resolved into 2 bands with isoelectric points of 5.8 and 6.2 and the 2 bands focused from eGH were 8.1 and 8.2 (data not shown).

**Bioactivity**

Potency estimates of eGH relative to bGH in the GH RRAs (using receptor sources from rabbit liver and bGH as standard and label) were high, ie 3.6 times that of bGH when pregnant rabbit liver microsomalms were used (fig 3, bottom left). The eGH displacement curve was parallel to the standard bGH. When mare liver plasma membranes were used as the receptor source and bGH as

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**Fig 2.** Analytical non-denaturing PAGE (A) of purified eGH, purified ePRL and the standard oPRL (NIAMDD oPRL-14). Alkaline discontinuous gel electrophoresis at pH 8.9 in a 7.0% acrylamide resolving gel was carried out on 20 µg of each hormone at a constant current of 20 mA and 180 V for 2 h on a Biorad mini-gel system. Analytical sodium dodecyl sulphate PAGE (B) of purified eGH, purified ePRL and standard oPRL (NIAMDD oPRL-14, lane 4). Electrophoresis was carried out at pH 8.8 in a 12.5% acrylamide resolving gel. Hormone samples (20 µg) were prepared for electrophoresis by dissolving each in 50 µl NaOH (0.01 M) followed by addition of 0.05 M Tris pH 6.7 (50 µl). The molecular weight markers were: alpha lactalbumin (Mr 14 000); trypsin inhibitor (Mr 20 100); carbonic anhydrase (Mr 30 000); ovalbumin (Mr 43 000); albumin (Mr 67 000); and phosphorylase B (Mr 94 000).
the standard and radioligand, potency estimates of eGH were low (0.045) relative to bGH corresponding to an activity of 0.04 iu/mg (fig 3, bottom right). Total binding of $^{125}$I bGH to receptors ranged from 15.4 to 30%, with non-specific binding in the range 4.8 to 11%. Potency estimates of ePRL relative to oPRL, in all of the PRL RRAs (using rabbit or mare mammary gland or liver as receptor sources and oPRL as the standard and tracer) were low (0.01–0.04) corresponding to activities in the range 0.37 to 1.2 iu/mg (fig 3, top left and right). All of the ePRL displacement curves were parallel to that of the standard, except that which used lactating rabbit mammary gland as the

Fig 3. Characterisation of (bottom left and right) the binding of $^{125}$I-labelled bGH and unlabelled eGH to crude membranes prepared from rabbit and mare liver and the binding of $^{125}$I-labelled oPRL and radioinert ePRL to crude membranes prepared from rabbit and mare (top left and right) mammary tissue. Stock solutions of hormones (2 µg/ml) were serially diluted in assay buffer (25 mM Tris, 10 mM CaCl₂, 0.1% BSA, pH 7.8) to give hormone concentrations in the range 0.098 ng to 5 µg/ml. Into plastic test tubes were added 0.1 ml each of standard solution or sample, tracer and membrane preparation. Total binding was estimated by substituting assay buffer instead of hormone and non-specific binding estimated by using an excess (20 µg) of standard. The tubes were incubated for 18 h at 4°C and the reaction was stopped by adding 2 ml of ice-cold assay buffer followed by centrifugation at 3 000 g for 20 min. The supernatants were aspirated and the pellets assayed for radioactivity.
receptor source. Total binding of $^{125}$I labelled oPRL ranged from 17 to 20%, with non-specific binding in the range 4.3 to 9%. In the Snell dwarf mouse bioassay, dwarf mice responded to the hormones by increased growth rates and uptake of $^{35}$SO$_4$ into cartilage. Both porcine and eGH demonstrate similar dose–response curves, with linearisation towards higher doses (40 to 160 µg) (fig 4). In the mouse mammary gland bioassay increasing doses of both equine and oPRL significantly increased the secretory grades of the cultured explants over control grades. However, the response with ePRL was significantly lower at comparable doses of lactogenic hormone. Comparison of the ED$_{50}$ values obtained from linear regression of dose–response curves gave a potency estimate of 0.4 relative to oPRL corresponding to 12.4 iu/mg. In the pigeon

crop sac bioassay ePRL gave a potency estimate of 29.23 iu/mg (15.8–53.6 iu/mg).

**Imnoactivity**

Equine GH cross-reacted fully in the pGH RIA of Buttle (1987a) and parallel dose–response curves were obtained (fig 5A). The potency of ePRL relative to pPRL in a homologous pPRL RIA was 0.96 (32.6 iu/mg) (fig 5B). Equine PRL cross-reacted fully in the homologous ePRL RIA of Worthy et al (1986).

**Radioimmunoassay of eGH**

A dose–response curve for eGH in the eGH RIA is presented in fig 6A. The limit of sensitivity of the assay, defined as the concentration of hormone which gave a $B/B_0$ ratio of 90% was 1.2 ng/ml (where $B$ is specific binding and $B_0$ is total binding). Ovine, bovine and human GH at concentrations of up to 800 ng per tube showed no significant displacement of $^{125}$I eGH, only pGH showed inhibition with a cross-reactivity of 0.24% (fig 6A) however the slope of the dose–response curve was non-parallel to the eGH standard. Of the PRLs tested, ovine, bovine, rat, porcine and equine, at concentrations up to 400 ng per tube, none showed any significant displacement; only ePRL showed inhibition with a slight cross-reactivity of 0.9%. The equine gonadotropin fraction did not cross-react in the assay. Dose–response curves of a crude pituitary gland extract and plasma samples collected from a mare and foal are also shown and dose–response curve analysis revealed that all curves were parallel. The quantitative recovery of eGH standard, serial dilutions from 2.5 to 20 ng/ml, added to equine plasma was 86.2 ± 3.3 (se)% with a mean intra- and interassay CV of 6.0 and 9.9%, respectively ($n = 6$) for repeated determinations on plasma pools.

![Fig 4.](image-url)
Radioimmunoassay for equine GH and equine PRL

Radioimmunoassay of ePRL

A dose–response curve for ePRL in the RIA is presented in figure 6B. The limit of sensitivity of the assay was 0.5 ng/ml. Of the PRLs tested (ovine, bovine and rat) at concentrations of up to 800 ng per tube, none showed any significant displacement. Only pPRL showed inhibition with a slight cross-reactivity of 0.4%. Ovine, bovine or pGH at concentrations of up to 800 ng per tube showed no significant displacement of $^{125}$I ePRL. Only eGH showed slight inhibition with a cross-reactivity of 0.1%. The effect of different dilutions of crude alkaline extract of equine pituitary gland and of plasma samples collected from a mare and a foal are
Fig 6. Dose–response curves of different hormone preparations in the homologous eGH RIA (A) and in the homologous ePRL RIA (B). In A, the preparations used were porcine GH (USDA-B1), bovine GH (NIH-B18) and purified equine PRL. The results for ovine PRL (NIH-P-S12), bovine PRL, (NIAMDD-bPRL-6), rat PRL (NIDDK B-6), porcine PRL (USDA B-1), ovine GH (NIH S-11), and human GH (MRC) are contained within the broken lines. In B, the preparations used were porcine PRL (USDA B-1), ovine PRL (NIH-P-S12), bovine PRL (NIAMDD bPRL-6) and purified equine GH. The results for ovine GH (NIH-S-11), bovine GH (NIH-b18), porcine GH (USDA-B-1) and rat PRL (NIDDK-rPRL-B-6) were contained within the broken lines. PE (an alkaline extract of equine pituitary gland), GnT (an equine gonadotrophin fraction) and P1 and P2 (dilutions of plasma samples from a mare and a foal respectively).
presented and dose–response curve analysis revealed that all curves were parallel. Recovery of ePRL standard (serial dilutions from 2.5 to 20 ng/ml) added to equine plasma was 111 ± 5.36 (se)% with a mean intra- and interassay CV of 9.1 and 15.6%, respectively (n = 6).

**Effect of administration of TRH on plasma PRL and GH**

Two out of the 3 TRH injected animals responded with a rapid surge of PRL within 15 min. The vehicle injected lactating mare, as expected, did not exhibit any increase in PRL (fig 7A). PRL concentrations in the non-lactating mare were highest 45 min post-injection reaching 10.9 ng/ml and declined thereafter to 4.9 ng/ml 150 min post-injection. PRL secretion in the stallion was elevated to an even greater extent (22.8 ng/ml) with the time course of the effect approximating to that of the non-lactating mare. The TRH injected lactating mare failed to respond as rapidly as the others, but exhibited a rise in PRL from 2 to 7 ng/ml between 120 and 150 min post-injection. This mare had also experienced a natural pulse of PRL just before TRH injection. In all of the TRH injected animals there was no immediate rise in GH. In the non-lactating mare concentrations ranged from 4 to 8 ng/ml, in the stallion from 2.5 to 8 ng/ml and in the lactating mare from 1 to 3.5 ng/ml. The vehicle-injected lactating mare had concentrations in the range 2 to 10 ng/ml and exhibited a large pulse of GH 30 to 120 min post-vehicle injection (fig 7B).

**Administration of hGRF(1-29) and the effect of fasting on plasma GH**

Overall mean GH concentrations did not differ significantly in the saline or GRF-treated animals over the sampling period with concentrations of 3.83 ± 0.31 (se) ng/ml (n = 3) for the saline group and 5.32 ± 1.18 (se) ng/ml (n = 3) for the hGRF(1-29)-treated group. Both groups exhibited a pulse of GH after infusion at 12.00 h and although the GH pulse in the GRF-treated group appeared earlier (5 min post-infusion, cf, 15 min for the saline treated) the timing was not significant (fig 8). Similarly fed and fasted horses had concentrations of GH which did not differ significantly (3.81 ± 0.96 (se) ng/ml (n = 6) (fed) and 4.5 ± 0.79 (se) ng/ml (n = 6) (fasted) over the time period examined.

**GH and PRL secretory profiles**

The profiles are characterized by large intermittent elevations in GH concentrations separated by trough periods during which GH returns to basal levels (fig 9A). In mare #101, concentrations of GH ranged from 1.8 to 29.08 ng/ml with up to 7-fold changes detected in a time period of 1 h. Mean 24-h GH were 5.3, 6.4 and 5.5 ng/ml in weeks 1, 3 and 6 of lactation. In mare #102, GH concentrations ranged from 1.7 to 89.6 ng/ml with up to 23-fold changes in GH detected in a time period of 1 h. Mean 24-h GH concentrations were 6.26 and 7.14 ng/ml in weeks 1 and 8 respectively. In the non-lactating mare #104 the overall 24-h mean GH was 5.9 ng/ml and in the stallion #105, 7.3 ng/ml with no distinct elevations in hormone concentration (profile not shown).

Lactating mare PRL profiles are characterized by long discrete episodic bursts, defined by increasing PRL concentrations due to more or less superimposed or partially overlapping secretory pulses. These are separated by relatively quiescent periods when PRL concentrations returned to basal levels. In mare #101, concentrations ranged from 1 to 23 ng/ml with up to 5-fold changes detected in 75 min. Mean 24 h PRL concentrations decreased from 5.27 to 4.18 ng/ml from week 1 to week 6 of lactation.
Fig 7. Plasma PRL (A) and GH (B) in response to an intravenous injection of 2.34 mg TRH in a lactating mare (□), a non-lactating mare (■) a stallion (♦) and an injection of physiological saline in a lactating mare (♦). Blood samples were collected at 15 min intervals from -60 to 180 min relative to injection.

Fig 8. Plasma GH in response to an intravenous injection of 670 μg of hGRF$_{1-29}$ or saline administered to 3 Dutch warm-blooded mares. Blood samples were collected at -180, -150, -120, -90, -60, -30, 0, 5, 10, 15, 30, 45, 60, 90, 120, 150, 180 and 240 min relative to the administration of hGRF$_{1-29}$ at noon.
Fig 9. Basal circulating levels of PRL (▲) and GH (△) in blood samples taken at 15 min intervals over a 24-h period in the lactating mares, #101 (profile A) #102 (profile B), and #103 (profile C). The profiles represent the arithmetic mean hormone concentrations of duplicate determinations at that time point. The dark rectangles (on top) represent periods of darkness.
In mare #102 PRL concentrations ranged from 1 to 46 ng/ml with up to 9-fold changes detected in 75 min. Mean 24 h PRL decreased from 10.89 to 6.48 ng/ml from weeks 1 to 8 of lactation. Two episodic bursts were detected in this mare with both light and dark peaks indicating the presence of a biphasic secretory pattern (fig 9). No major elevations in PRL could be detected in #104, the non-lactating mare or in #105 the stallion 24-h PRL profiles; the overall 24-h mean PRL concentrations were 3.18 and 3.29 ng/ml respectively.

DISCUSSION

Highly sensitive, specific RIAs for eGH and ePRL have been developed employing highly purified and well-characterised preparations of hormone. These assays have been employed in the assay of closely timed plasma samples over 24 h in both lactating and non-lactating animals. Specific patterns of secretion for each hormone were described.

To our knowledge, this is the first report of a homologous RIA for eGH. Previous studies have relied on heterologous assays, employing antiserum to pGH and pGH as the label (Thompson et al, 1992). The validation data presented here show that the sensitivity, precision and accuracy of the RIA is adequate for the investigation of changes in GH concentrations. Cross-reactivity studies of purified preparations of several mammalian GHs and PRLs as well as equine gonadotrophins in the RIA have demonstrated that the assay was highly specific for eGH. In the case of ePRL, its cross-reactivity was low suggesting very little cross-contamination of the ePRL standard with eGH. No interference by plasma components was observed in the RIA as demonstrated by the parallelism of the inhibition curves of plasma PRL and crude PRL pituitary extracts, and the good recoveries of exogenous ePRL added to plasma samples.

Similar yields and size estimates for eGH have been obtained by previous investigators (Saxena and Henneman, 1966; Hartree et al, 1968; Conde et al, 1973). Equine GH showed full cross-reactivity in the bGH RRA employing rabbit liver and labelled bGH but had a low potency when the receptor preparation was from mare liver with the same label. We were unable to demonstrate specific binding of labelled eGH to either mare or rabbit liver receptors. Equine GH was equipotent with pGH in the Snell dwarf mouse bioassay and in a homologous porcine GH RIA confirming the purity of eGH and its immunological identity with porcine GH.

Similar to the binding pattern seen with eGH and bGH in mare liver (ie higher affinity of bGH), although ePRL could bind to mare and rabbit mammary gland and liver, it competed much less effectively than oPRL. This is similar to the results of Jerry
et al (1991) who found that specific binding of oPRL to porcine mammary membranes was greater than specific binding of the homologous pPRL. It is possible that oPRL binds to sites in the equine mammary gland in addition to the authentic lactogenic receptor or that it has a higher affinity for the equine lactogenic receptor. It has been suggested that homologous PRLs are unstable and may be uniquely cleaved at the plasma membrane and released as 16 and 8 kDa fragments (Clapp, 1987). It is also possible that chemical damage to ePRL, but not oPRL, during iodination could explain this discrepancy in binding. Parke and Forsyth (1975) have also demonstrated that preparations of canine and ePRL with high pigeon crop sac stimulating activity, were less than 1 iu/mg in terms of the ovine PRL standard in the rabbit liver RRA. The high potency of ePRL in the pigeon crop sac bioassay compares well to the values obtained by other investigators (Chen et al, 1979; Li and Chung, 1983) and its low potency in the mouse mammary gland bioassay emphasizes the need to examine the action of hormones using homologous systems. The purity of our ePRL standard was confirmed in a previously established homologous ePRL RIA (Worthy et al, 1986) and significant cross-reactivity was obtained with a homologous porcine PRL RIA, which conforms well with the close structural homology between these hormones (Lehrman et al, 1988).

The lack of response of plasma GH to hGRF(1-29) was somewhat surprising, but it is known that several factors can affect an animal's ability to respond. Dubreuil et al (1987) found that the GH response to GRF in the swine was affected by age, sex and the timing of the GRF infusion relative to a natural pulse of GH with females responding better than males. Thompson et al (1992) have reported a response to porcine GRF in 8 out of 9 mares administered the hormone. The results of the TRH stimulation test showed that it is unlikely that the horse responds to this hormone by rising plasma GH, confirming the results of Thompson et al (1992). However elevations in PRL concentrations after TRH administration were detected, supporting the observations of previous investigators (Thompson and Nett, 1984; Thompson et al, 1986a; Johnson, 1987). The results also suggest that the PRL response to TRH in the mare may become refractory as in the lactating mare a large peak of PRL was detected prior to TRH injection and the rise in PRL post-TRH administration was not observed until 1 h afterwards.

Peak lactation in the mare occurs at around 8 weeks and weaning in natural herds occurs 35–40 weeks after birth (Duncan et al, 1984). The 24-h profiles shown are from mares in early to peak lactation. Closely timed blood samples over 24-h periods showed 2 quite distinct patterns of secretion for PRL and GH in the lactating mare. Large 'unimodal pulses' of GH and 'episodic bursts' of PRL were the typical patterns of secretion in the lactating mare. The pattern of secretion of GH is similar to that described in lactating pigs and cows (Buttle, 1987a; Bines et al, 1983). In lactating cows, no consistent pattern of GH secretion was found at any stage of lactation but very high short-lived peaks of GH occurred at irregular intervals in high yielding cows at peak lactation. In the dry period and in low yielding cows these peaks were smaller and less frequent, compared with the smaller GH elevations observed in the non-lactating mare. Buttle (1987a) has ascribed the pattern of GH secretion in lactating pigs to weight loss and negative energy balance. It is noteworthy that many of the GH pulses recorded in the mare coincided with the onset or occurred during the dark period. In man the onset of typical slow-wave sleep stimulates an increase in the blood level of GH (Honda et al, 1969) and this release is inhibited by the occurrence of paradoxical sleep.
No consistent pattern of PRL secretion was detected between all the mares but each mare had her own characteristic pattern. In mare #101 one episodic burst occurred in the afternoon. Roser et al (1987) have also shown evidence of a midday diurnal surge in PRL in mares. In mare #102 a biphasic pattern was observed with both light and dark peaks. Similarly Lincoln (1979) observed both a day- and a night-time peak of PRL in the ram, and a similar diurnal rhythm has been reported with a nadir around noon and a peak before midnight in the ewe (Wallace et al, 1988). The 3 lactating mares were freely lactating, ie foals remained with the mares, and all animals were fed ad libitum ensuring undisturbed patterns of hormone secretion. When Wiest and Thompson (1987) separated dams from foals and reunited them some time later they found that PRL was released in some but not all mares. However the continuous sucking pattern throughout certain time intervals typical of the foal may result in the release of relatively small quantities of the hormone throughout the period of stimulation, as occurs in the lactating rat (Grosvenor and Whitworth, 1974) making elevations in hormone concentrations due to the effects of sucking difficult to observe. Moreover, when sucking episodes occur frequently there is insufficient time between them for PRL levels to fall to low levels and so basal concentrations of PRL remain elevated.

In conclusion, we have developed new homologous RIAs for eGH and ePRL using highly purified and well-characterised preparations of hormones. The RIA for eGH has enabled initial studies of the physiology of eGH and it will enable us to carry out more extensive studies to define more clearly the physiological role of eGH in the horse.

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