Promotion of intragranular co-aggregation with LH by enhancement of secretogranin II storage resulted in increased intracellular granule storage in gonadotrophs of GnRH-deprived male mice

J. L. Crawford*, J. R. McNeilly, L. Nicol and A. S. McNeilly†

MRC Human Reproductive Sciences Unit, Centre for Reproductive Biology, The University of Edinburgh Chancellor’s Building, 49 Little France Crescent, Old Dalkeith Road, Edinburgh EH16 4SB, UK

Intracellular associations indicate that granins may play a role in the regulatory mechanisms involved in differential secretion of gonadotrophins. The effect of GnRH on mRNA expression, storage and secretory patterns of granins and gonadotrophins was investigated in male mice. GnRH antiserum (G/A) was injected into mice in the treatment group (n = 15) at 12 h intervals for 2 days and a subset (n = 9) was killed. Buserelin (G/A + B) was administered to the remaining mice (n = 6), which were killed 2 h later; control mice (n = 6) were killed at the onset of the study. LHβ mRNA content was lower in G/A and G/A + B mice compared with controls, whereas plasma LH concentrations were higher in G/A + B mice. FSHβ mRNA content did not change, whereas plasma FSH concentrations were lower in G/A mice compared with controls, and higher in G/A + B mice compared with both G/A and control mice. Secretogranin II (SgII) and CgA mRNA contents were not different between experimental groups. There were more granules per gonadotroph in G/A mice, and considerably fewer after Buserelin treatment. Immunogold labelling of gonadotrophs revealed the presence of LH+/SgII+ and LH+/SgII− granules, and negligible numbers of LH+/SgII+ granules. Both the numbers of LH+/SgII+ granules and overall granule antigenicity for SgII were higher in G/A mice compared with controls and G/A+B mice. In contrast, there were fewer LH+/SgII− granules per gonadotroph in G/A mice compared with controls. In conclusion, absence of GnRH input to the pituitary gland resulted in preferential storage of SgII and subsequently increased intragranular co-aggregation with LH. Administration of Buserelin to G/A mice resulted in the apparent release of LH+/SgII+ granules that was reflected by an increase in plasma LH concentrations, indicating that these granules were in the regulated secretory pathway. In contrast, secretion of LH+/SgII− granules did not appear to be influenced by the actions of Buserelin and, therefore, may have been destined for constitutive release, possibly to maintain basal plasma LH concentrations.

Introduction

Granins are acidic glycoproteins that possess aggregative (Gerdes et al., 1989; Gorr et al., 1989) and calcium-binding (Cozzi and Zanini, 1986; Reiffen and Gratzl, 1986) properties, and are closely associated with gonadotrophins at the subcellular level (Watanabe et al., 1991, 1993; Jeziorowski et al., 1997). Hence, granins may be involved in the intracellular regulation of differential secretion of gonadotrophins in the specialized anterior pituitary cell, the gonadotroph. The regulation of pulsatile secretion of LH and of tonic secretion of FSH is complex and involves interactions between GnRH, activin, inhibin and ovarian-derived steroids (Clarke and Cummins, 1982; Levine et al., 1985; Farnworth et al., 1988; Farnworth, 1995). Specific inhibition of GnRH influence throughout administration of a GnRH agonist for 6 weeks in ewes completely abolished both pulsatile LH release and responsiveness of LH and FSH to GnRH stimulation, and also reduced both LHβ and FSHβ mRNA contents (to 5% and 30% of luteal control values, respectively), and pituitary content of LH and FSH (to 3% and 30% of control values, respectively). The significant reduction in β-subunit gene expression of both gonadotrophins to basal values was associated with only a 50% reduction in plasma concentrations of LH and FSH (McNeilly et al., 1991), demonstrating that LH and FSH are secreted constitutively, as well as via the regulated secretory pathway. It is likely that both constitutively and regulatory released granules are contained within the same gonadotroph. In sheep, Crawford et al. (2000) showed that 24 h after an exogenously induced preovulatory LH surge, 100% of gonadotrophs observed were devoid of properly formed

*Present address: Reproductive Biology Group, AgResearch, Wallaceville Animal Research Centre, PO Box 40063, Upper Hutt, New Zealand

†Correspondence Email: a.mcneilly@hrsu.mrc.ac.uk


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granules, which was consistent with an absence in pulsatile LH secretion, indicating that all gonadotrophs contained some stored LH that was released only on extracellular stimulation. However, basal LH concentrations in these animals remained unaltered, confirming that at least a small proportion of LH was being released constitutively from some or all gonadotrophs (Crawford et al., 2000). In addition, three types of granule were identified in gonado-
trophs of the rat, and granule populations shifted in males after stimulation by either castration or GnRH treatment in males (Watanabe et al., 1991, 1993, 1998a), as well as in females during the oestrous cycle (Watanabe et al., 1998b). Small LH- and SgII-positive granules, and large FSH- and CgA-positive granules, were located in gonadotrophs of male and female rats. After stimulation in male rats and at impending ovulation in female rats, a third type of intermediate-sized granule appeared, which contained an electron-dense LH- and SgII-positive core, and an electron-lucent CgA-positive outer region.

The mechanism through which granins may act as regulatory agents in differential gonadotrophin sorting, packaging and trafficking is unknown, although the potential role of a highly conserved N-terminal disulphide-bonded loop positioned on chromogranin A (CgA) and particularly chromogranin B (CgB) in the sorting of regulated proteins has received attention. Prevention of disulphide bond formation of CgB in a rat neuroendocrine cell line PC12 resulted in mis-sorting of CgB to the constitutive pathway (Chanat et al., 1993, 1994; Krömer et al., 1998). Alternatively, the addition of the disulphide-bonded loop region of CgB on to a constitutively secreted protein was sufficient to redirect the fusion protein into a regulated pathway (Glombik et al., 1999), apparently mediated through binding to a sorting receptor within the trans-Golgi network, such as the membrane-associated carboxypeptidase E (Cool et al., 1997). This theory was later contested by several findings including a recently discovered alternative role of the disulphide loop in homodimerization in CgA (Thiele and Huttner, 1998), the absence of the first cysteine residue at the N-terminal region failing to prevent anchorage to vesicle membrane-coupled columns (Yoo and Kang, 1997), and a lack of correlation between redox state of the loop in situ and entry of CgB into the regulated secretory pathway (Arvan and Castle, 1998). Moreover, it has recently been shown that CgA contains a binding site for inositol (1,4,5)-triphosphate (IP3) receptor (Yoo, 2000), a receptor that also functions as a Ca2+-channel within secretory granule membranes (Yoo, 1994; Yoo et al., 2000).

Despite a very small number of studies on the intracellu-
lar associations between granins and gonadotrophins in the rat (Watanabe et al., 1991, 1993, 1998a,b), the benefits of studying the relationship between granins and gonado-
trophins in mice are considerable in view of the wide range of transgenic mice models available. Therefore, the aim of the present study was to define the mechanisms through which granins may facilitate synthesis, and storage or secretion of gonadotrophins from gonadotroph cells in male mice. Transcriptional, storage and secretory relationships between granins and gonadotrophins in male mice deprived of GnRH input through administration of GnRH antibody and then subsequently stimulated with GnRH agonist were investigated. Shifts in granule populations in relation to morphology and antigenicity for gonadotrophins and granins were investigated and correlated with alterations in their transcription and secretion to determine the intracellular processing mechanisms.

**Materials and Methods**

**Animals and experimental design**

Wild-type (WT; n = 21) male mice were used in this experiment. Six mice were killed at the onset of the study to act as controls. Four injections (i.p.) of GnRH antiserum (0.1 ml G/A; Clarke et al., 1978; Fraser et al., 1981; McNeilly et al., 1984) were administered at 12 h intervals to 15 mice; nine of these G/A-treated mice were killed after the final injection. Synthetic GnRH was administered by i.p. injection (50 ng Buserelin; Sigma Chemical Co., St Louis, MO; G/A + B group) to the other six G/A-treated mice, which were killed 2 h later. A blood sample was collected immediately before death from all animals and the pituitary glands were extracted immediately after death.

Each pituitary gland was weighed and divided transversely into two evenly sized pieces, each containing a similar area of anterior pituitary. One half of the pituitary gland from each animal was randomly allocated for storage in liquid nitrogen for molecular biological analysis. The other half was further divided evenly in an anteroposterior direction and randomly allocated for either processing for confocal microscopy or transmission electron microscopy (TEM).

**RNA extraction and real-time PCR**

LHβ, FSHβ, SgII and CgA mRNA contents were deter-
mined using a quantitative real-time PCR method. Frozen pituitaries were transferred directly to RNazol B (AMS Biotechnology, Witney) and total RNA was extracted using the method recommended by the manufacturer. An aliquot of 800 ng in 8 μl from each total RNA sample was treated with DNase I from DNA-free® kit (Ambion, Inc., Austin, TX) following the manufacturer’s instructions to remove any residual DNA. A portion (300 ng; 40 ng μl–1) of DNase-
treated total RNA from each sample was reverse-transcribed using reagents including oligo d(T)18 provided with Taqman reverse transcription reagents kit (PE Biosystems, Warrington) in accordance with the manufacturer’s instructions. A programme of 10 min at 25°C, 30 min at 48°C and 5 min at 95°C was performed on a PTC-100 programmable thermal controller (GRi; Felsted, Dunmow).

Primers and probes were designed using the computer package ‘Primer Express’ (PE Biosystems). Primers and FAM-labelled probes for mLHβ (forward primer: TGTCTTAGCATGGTGTCGGATCT; reverse primer: AAGGCTACAG-
GAAAAGGACTATG; probe: CGGCTGTTTTGCTCTGTG, mFSHB (forward primer: GAGAGCAATCTGCTGCTCCATA; reverse primer: GCAGAAAAGGACATCTCTCCCT; probe: CTTGGAATTGACACACATCACCCTGAGTGA), mSgI (forward primer: CAAAGACACATCTCTCTG; reverse primer: AGGGAATTTGGCATTAAAGC; probe: TCCTGAAAAGTCTGGAGTACCCCTCAAACCA) and mCgA (forward primer: CCGACTGACCATCATCTTTCTG; reverse primer: CCGCTGACCATCATTCTTCTG; probe: CCACTTCCATGCAGCTACAAACCGA) were also developed by PE Biosystems. Ribosomal 18S primers and the VIC-labelled probe were supplied in a Taqman ribosomal RNA control reagents kit (PE Biosystems).

A reaction mix for real-time PCR was prepared by combining 1 x Taqman universal PCR master mix, specific forward (300 nmol l–1) and reverse (300 nmol l–1) primers, and specific Taqman probe (200 nmol l–1) for the mRNA of interest, ribosomal 18S forward (50 nmol l–1) and reverse (50 nmol l–1) primers and ribosomal 18S probe (200 nmol l–1). Samples were prepared in duplicate by placing aliquots of 48.88 µl reaction mix and 3.12 µl cDNA sample into autoclaved RNase-free 0.25 ml Eppendorf tubes and mixing well. Two 25 µl aliquots from each tube were transferred to adjacent wells in a 96-well real-time PCR plate. Optical caps were fixed on to the plates and the reaction was run on ABI Prism 7700 (PE Biosystems) using standard conditions. Controls included samples that underwent reverse transcription PCR with the exclusion of Multiscribe reverse transcriptase to check the effectiveness of DNase treatment, and reactions that omitted addition of template. Before analysis, serial dilutions of a sample (1:1 to 1:64) to ensure normalization of the amount of RNA used were made, and the line of best fit was plotted against log (input RNA). Quantification of bCgA (INCSTAR Corporation, Stillwater, MN).

Gonadotrophin radioimmunoassays

Reagents were supplied by the NIADDK and concentrations of plasma LH and FSH were measured by radioimmunoassay; all samples were included in duplicate in the same assay. The reference preparation used was rat LH-RP-1 and rat FSH-RP-3, and the sensitivity of the assays was 0.2 and 1.2 ng ml–1 for LH and FSH, respectively. The intra-assay coefficients of variation were < 10% (McNeilly et al., 1996, 2000).

Confocal microscopy

The antisera used included a monoclonal antiserum against bLHβ (518B7; J. F. Roser, Department of Animal Science, University of California, CA; Matteri et al., 1987) and rabbit-raised antiserum against hFSHB (M91; S. Lynch, Birmingham), bSgI (Pel-Freez Biologicals, Rogers, AR) and bCgA (INCAST Corporation, Stillwater, MN).

Pituitary pieces were fixed in Bouin’s fixative for 6–8 h before transfer to 70% (v/v) ethanol for storage until processing. Tissue was rehydrated with 50% (v/v) ethanol, 25% (v/v) ethanol and 0.01 mol PBS l–1 (PBS tablets; Sigma Chemical Co.) for 1 h each at room temperature, and then incubated overnight in permeabilization/blocking buffer (P/B buffer; 1% (w/v) BSA, 10% normal goat serum (NGS) and 0.003% (v/v) Triton-X100) at 4°C. Mixes of 5 µg ml–1 LH antibody, with either 1:200 FSH, 1:500 SgI or 1:1000 CgA antibodies, were made up in P/B buffer. Each piece of tissue was divided into three and a different antibody mix was added to each triplicate sample and incubated at 4°C for 72 h. The tissue pieces were washed twice for 1 h and then once for 2 h in 0.01 mol PBS l–1 at 4°C. A mix of secondary antibodies raised in rabbit (1:20 anti-mouse whole molecule IgG TRITC and 1:20 anti-rabbit whole molecule IgG FITC; Sigma Chemical Co.) were diluted in P/B buffer and added to each sample, and then incubated overnight at 4°C. The tissue was finally washed three times for 1 h and left overnight in 0.01 mol PBS l–1 at 4°C. Pituitary pieces were placed on washed glass slides (Chance Propper Ltd, Agar Scientific Ltd, Stanstead) inside adhesive Easi-Seal squares (Hybaid Ltd, Ashford) and mounted in Citifluor AF1 (Glycerol/PBS solution; Citifluor Ltd, London) before being encased with coverslips (Chance Propper Ltd) and sealed with nail polish.

Transmission electron microscopy

Pituitary tissue pieces were fixed and processed using a modification of a method described in Berrymann and Rodewald (1990) that reports high specific staining in pancreatic and jejunal tissue in the rat without sacrificing cell ultrastructure. Modifications included the fixative (2% (w/v) paraformaldehyde, 0.1% (v/v) glutaraldehyde (25% EM grade; Agar Scientific Ltd), 0.2% (w/v) picric acid and 0.5 mmol calcium chloride l–1 in 0.1 mol phosphate buffer l–1, pH 7.4), embedding resin (Unicryl resin) and embedding capsules (gelatin capsules, size 00; Agar Scientific Ltd).

Ultrathin sections (80 nm) were cut with a 4 mm ultra-diamond Diatome knife (45°; Leica UK Ltd, Milton Keynes) using a Reichert Jung Ultracut ultramicrotome (Wild Leitz Ltd, Milton Keynes) and mounted on hexagonal 200 mesh gold grids (Agar Scientific Ltd). Gonadotrophs were identified by an immunogold labelling method, using antisera described in the confocal microscopy methods section (dilutions were 5 µg ml–1 for LH, 1:200 for FSH, 1:1000 for SgI and 1:500 for CgA). Grids were blocked with 5% (v/v) NGS in Tris histochemical buffer (THB, pH 8.2; BDH Laboratory Supplies) and were then washed in THB. After 2 h incubation in LH antisera, the sections were washed in THB and incubated for a further 2 h in goat anti-mouse (1:50, 5 nm diameter gold particles for monoclonal antisera; British Biocell International Ltd) IgG–gold particle conjugates. Sections were washed in THB and then in ddH2O. This process was repeated for double immunogold labelling using FSH, SgI or CgA primary antibody, and then goat anti-rabbit (1:50, 10 or 15 nm diameter gold particles for polyclonal antisera; British Biocell International Ltd).
IgG–gold particle conjugates secondary antibody. After the final wash step, sections were fixed in 4% (v/v) glutaraldehyde. Sections were first stained in 2% (w/v) osmium tetroxide (Agar Scientific Ltd) for 1 h and then triple stained in Reynolds lead citrate (10 min; 0.08 mol lead nitrate l⁻¹, 0.12 mol trisodium citrate l⁻¹ and 0.16 mol sodium hydroxide l⁻¹), 2% (w/v) uranyl acetate (1 h) and again in Reynolds lead citrate (10 min), washing thoroughly in ddH₂O between incubations. Sections were viewed under a Philips CM 120 Biotwin TEM.

Every second gonadotroph cell in every second hexagonal grid slot was selected until ten immunogold-labelled gonadotroph cells were identified from 80 nm thick sections of pituitary tissue from five animals in each group. Within each gonadotroph, two areas of cytoplasm (containing ten granules each) on opposite sides of the nucleus were randomly selected and numbers of LH⁺ve/SgII⁺ve, LH⁺ve/SgII⁻ve and LH⁻ve/SgII⁻ve granules, as well as number of gold particles per granule, were counted while being viewed on the TEM. The total number of granules per gonadotroph was calculated by counting all granules within these selected cells.

Statistical analysis

All mass plasma and mRNA values, as well as gold particle and granule counts, are reported as mean ± SEM. Data were analysed using one-way ANOVA and where a significant (P < 0.05) interaction was calculated, a post-hoc Fisher’s protected least significant difference test was performed, using Statview statistical computer package (version 4.02; SAS Institute Inc., San Francisco, CA).

Results

Pituitary masses, plasma LH and FSH concentrations, and gramin and gonadotrophin mRNA contents

There were no significant differences in mean pituitary gland wet mass between experimental groups (data not shown). Mean plasma LH concentrations were significantly higher (P < 0.0001) in G/A + B mice compared with controls and G/A mice (Fig. 1a). Mean concentrations of plasma FSH were lower (P < 0.001 and P < 0.0001, respectively) in G/A mice compared with both control and G/A+B mice, but were higher (P < 0.05 and P < 0.0001) after Buserelin treatment compared with both controls and G/A mice (Fig. 1b). Mean LHβ mRNA contents were lower (P < 0.05) in both treatment groups compared with control animals (Fig. 1c). There were no significant differences in mRNA expression for FSHβ (Fig. 1d), SgII (Fig. 1e) or CgA (Fig. 1f).

Ultrastructural observations

Gonadotrophs had a lighter appearance than other types of pituitary cell after an osmium tetroxide and triple staining protocol (Fig. 2a). A dichotomy of granular staining was observed within gonadotrophs of control mice: granules were either completely electron-lucent or contained an electron-dense core surrounded by an electron-lucent outer region (‘halo of space’; Fig. 2b). Most granules observed in gonadotrophs from G/A mice were of the latter type (Fig. 2c), whereas entirely electron-lucent granules were prevalent in G/A + B mice (Fig. 2d).

Total numbers and antigenicity of granules

There were more (P < 0.01) granules per gonadotroph (80 nm thick section) in G/A mice compared with controls, but after Buserelin treatment fewer granules were observed compared with both G/A (P < 0.0005) and control (P < 0.05) mice (Fig. 3a).

There were fewer (P = 0.005) LH⁺ve/SgII⁻ve and more (P < 0.0005) LH⁺ve/SgII⁺ve granules in G/A mice compared with controls (Fig. 3b). After Buserelin treatment, the number of LH⁺ve/SgII⁺ve granules was lower (P < 0.0001) compared with G/A mice and similar to values of control animals, whereas numbers of LH⁺ve/SgII⁻ve granules had not
The amount of LH in relation to SgII in individual granules was determined by counting the gold particles bound to these antigens within granules to assess the association between LH and SgII during formation of granules and their individual partitioning and storage. This enabled the LH$^{+ve}$/SgII$^{+ve}$ granules to be divided into groups dependent on the proportion of LH and SgII present. The numbers of LH$^{+ve}$/SgII$^{+ve}$ granules in which LH-bound gold particles consisted of 26–50% and 51–75% of total gold particles (sum of LH- and SgII-bound gold particles) present per granule was higher ($P < 0.05$ and $P < 0.0005$, respectively) in G/A mice compared with controls (Fig. 3c). After Buserelin administration, the numbers of these granules had decreased ($P < 0.05$ and $P < 0.0001$, respectively), again to values comparable to those observed in control mice (Fig. 3c). In contrast, the numbers of LH$^{+ve}$/SgII$^{+ve}$ granules

![Image of transmission electron micrographs](image-url)
in which LH-bound gold particles consisted of 0–25% and 76–99% of total gold particles present per granule were not different between experimental groups.

Overall granule antigenicity (defined as the total number of gold particles per granule) for SgII was higher (*P* < 0.005) in G/A mice compared with controls, but after Buserelin treatment antigenicity was lower (*P* < 0.005) compared with G/A mice and similar to values observed in controls (Fig. 3d). There was no difference in granule antigenicity for LH between experimental groups (Fig. 3d).

In sections immunogold-labelled for LH (5 nm gold particle) and SgII (15 nm gold particle), the labels for both proteins were located throughout the matrix of the granule (Fig. 4a). On the rare occasion that FSH (15 nm) labelling was observed, the gold particles appeared to be located near the periphery of the granule (Fig. 4b), although labelling for both FSH and CgA was sparse, indicating that the TEM processing conditions were too harsh for the preservation of these proteins. Therefore, these results were excluded from the study.

Confocal microscopy observations

Pituitary sections were immunofluorescently bi-labelled for LH and FSH, LH and SgII, and LH and CgA, and were viewed using a confocal microscope. In those sections labelled for LH and FSH, mono-hormonal (for both gonadotrophins) and bi-hormonal gonadotrophs were identified in animals from all groups (Fig. 5a–f). Although the predominant populations were bi-hormonal, there were no differences in the proportions of these populations between groups (data not shown). A similar pattern was observed for sections labelled for LH and SgII (Fig. 6a–f); proportions of mono-labelled and bi-labelled cells were not different between experimental groups (data not shown). Sections labelled for LH and CgA also contained both mono-labelled (for both antigens) and bi-labelled cells (Fig. 7a–f). Punctate labelling of CgA was observed in nearly all CgA-positive cells (Fig. 7e,f). Gonadotrophs that were positive for both LH and CgA were prevalent in all experimental groups, although the only cell populations that changed were those labelled for CgA only, with a higher proportion (*P* < 0.05) in G/A + B mice compared with control and G/A mice (data not shown).

Discussion

This study investigated the relationship between transcription, storage and secretory patterns of granins and gonadotrophins in mice, and has shown for the first time that SgII does indeed play a role in facilitating storage and secretion of LH. Evidence that SgII is involved in the regulatory processing of LH strengthens the hypothesis that granins are involved in the regulation of differential secretion of gonadotrophins. This finding paves the way for future investigations in this area, and hypogonadal and transgenic mice will provide ideal physiological model systems to interpret these regulatory mechanisms.

Storage of LH within gonadotrophs is necessary for its rapid and concomitant pulsatile release under the action of
Co-aggregation of intragranular SgII with LH

Fig. 4. Transmission electron micrographs of areas of cytoplasm within immunogold-labelled 80 nm sectioned gonadotrophs that contained granules positive for (a) LH (5 nm) and secretogranin II (SgII; 15 nm), and (b) LH (5 nm) and FSH (15 nm) in male mice. Note that gold particles bound to LH and SgII were located throughout the matrix of the granules, whereas most of the gold particles bound to FSH are situated near the periphery of the granules (arrows). Scale bars represent 250 nm.

Fig. 5. Localization of LH (red) and FSH (green) immunofluorescence by confocal microscopy at (a,b,c) low and (d,e,f) high magnification in thick pituitary sections containing gonadotrophs from (a,d) control male mice, or from male mice treated for 2 days with either (b,e) GnRH antiserum (G/A) or (c,f) G/A followed by a bolus of Buserelin (G/A + B). The yellow staining represents areas of dual LH–FSH immunofluorescent labelling within gonadotrophs. Note that LH only, FSH only and co-localized LH–FSH-labelled gonadotrophs were present in animals from all experimental groups. Scale bars represent (a,c,e) 50 μm and (b,d,f) 10 μm.
GnRH and, presumably, is directed at the level of exocytosis. Total intracellular granule storage increased after GnRH antiserum administration in the present study, which is perhaps not surprising considering that probably negligible amounts of stored LH were released over the 48 h treatment period due to the absence of exocytotic-inducing stimulation. The type of granule that appeared to be stored within this 48 h treatment period was LH+/SgII+ granules containing moderately even numbers of both antigens (25–75% LH-bound and 75–25% SgII-bound gold particles). In contrast, there were fewer LH+/SgII- granules in G/A mice compared with controls, indicating that this type of granule was continually released during this time.

These changes in granule populations in G/A mice compared with controls were associated with reduced LH mRNA expression and constant plasma LH concentrations. This latter result may be due to a combination of existing low concentrations of circulating LH in control mice and the possible inability to detect even significant changes in circulating plasma LH concentrations from one sample taken before death due to the pulsatile nature of LH release. However, administration of this GnRH antiserum in sheep in the follicular stage of the oestrous cycle also did not affect basal concentrations of LH, but promptly abolished its pulsatile secretion (McNeilly et al., 1984). Therefore, it is possible that in the present study, GnRH antibody abolished pulsatile LH secretion, while basal concentrations remained unaffected, but the inability to collect blood samples intensively from mice makes it impossible to comment on pulsatile hormone release in this species. Disparity between β-subunit mRNA expression and plasma concentrations of LH is well known and is facilitated through the ability of gonadotrophs to store LH (Turgeon et al., 1996). Moreover, changes in stability of LHβ mRNA transcript, through alterations in 3′-poly(A) tail length, potentially provide additional means of post-transcriptional regulation of LH synthesis and are influenced by treatment with GnRH (Weiss et al., 1992), progesterone (Wu and Miller, 1991) and oestradiol (Park et al., 1996), and occur during the follicular phase of the cycle in sheep (Crawford and McNeilly, in press).

Integration of these results not only implies that LH+/SgII+ granules were released constitutively to maintain basal LH concentrations in the present study, but also that SgII may have been required for increased packaging and storage of LH, as well as perhaps contributing to the exocytotic responsiveness of granules to GnRH stimulation. Alternatively, SgII secretion may be entirely regulated and, therefore, when exocytosis is blocked, SgII is also trapped into increased storage.
storage. Even though transcription rate for LHβ had decreased in mice treated with GnRH antiserum, total granular antigenicity for LH did not change, indicating that the rate of granule packaging may have slowed to allow intragranular collection of LH equivalent to pretreatment amounts. This was substantiated by increased SgII content within granules despite constant SgII mRNA content, and further indicates active aggregation of LH and SgII under these conditions.

The significant increase in plasma LH concentrations in G/A + B mice in the present study verified the ability of Buserelin to bypass the inhibitory actions of GnRH antibody (Clarke et al., 1978) and indicated that the bulk of LH released in these mice may have been derived from the intracellular stores of gonadotrophs. Indeed, it appeared that the predominant type of granule released after Buserelin treatment was LH+/SgII+ve granules (containing moderately even numbers of both antigen), whereas numbers of LH+/SgII-ve granules remained unaltered compared with those in G/A mice. This finding supports the hypotheses that LH+/SgII+ve granules were released in a regulated fashion in response to GnRH, that LH+/SgII+ve granules were released constitutively to maintain basal concentrations of LH and that SgII plays an active role in regulated secretion of LH. If this is true, the reduction in numbers of LH+/SgII-ve granules evident after GnRH antiserum treatment in these mice compared with that in controls indicates that formation of these granules was curbed by a reduced rate of LHβ transcription. In other words, these granules may accumulate within the cytoplasm as a net result of varying LHβ mRNA content and steady exocytotic rates, which may be under autocrine control and not due to regulated storage.

The marked attenuation in plasma FSH concentrations after administration of GnRH antiserum, in conjunction with unaltered FSHβ mRNA content, indicates that a proportion of FSH is released via the regulated secretory pathway. This contention was confirmed by the action of Buserelin, which resulted in increased plasma FSH concentrations, presumably from release of intracellular FSH stores. Unfortunately, as a result of the sensitivity of FSH and CgA antigenicity to TEM processing methods, very little of either of these antigens was detected within granules in the present study. Therefore, due to the limited information obtained on FSH and CgA, no conclusions can be drawn with respect to the relationship between granin and FSH storage and synthesis.

In the few granules in which specific FSH label was observed, FSH appeared to be situated preferentially near the periphery of the granule, although the granular
membrane was not easily definable. In contrast, strong and sensitive labelling of LH and SgII revealed their location throughout the matrix of the granule. This partitioning has also been reported in rat gonadotrophs (Watanabe et al., 1993, 1998b). Indeed, compartmentalization of stored CgA was thought to be responsible for the punctate labelling of CgA observed by confocal microscopy in the present study. Recent studies have identified IP3 receptors on membranes of secretory granules, as well as IP3 receptor-binding sites on CgA (for a review, see Yoo, 2000), and, therefore, CgA may serve as a gate-keeper in the sorting of gonadotrophins (Arvan and Castle, 1998) and facilitate granule membrane–plasma membrane interactions at the event of exocytosis. The increase in CgA-only cells observed by confocal microscopy in G/A + B mice in the present study indicates that CgA may not be released after Buserelin administration, but may be recycled together with the granule membrane. Although early immunocytochemical studies showed localization of the three major granins to within gonadotrophs and a few corticotrophs, whereas most thyrotrophs contained CgA and a few contained SgII (O’Connor et al., 1983; Cohn et al., 1984; Rosa et al., 1985; Rundle et al., 1986; Hearn; 1987), GnRH receptors are only present on gonadotrophs in the anterior pituitary gland.

The characteristic morphology of regulated secretory cells is the accumulation of secretory granules (Burgess and Kelly, 1987). The pathway of formation of nascent storage granule begins with a protrusion of the trans-Golgi network to create a condensing vacuole (Palade, 1975; Novikoff et al., 1977) that transforms into an immature granule capable of stimulus-dependent secretion (Arvan et al., 1991; Tooze et al., 1991) only on detachment from the trans-Golgi network (Tooze and Tooze, 1986). Maturation of an immature granule to a mature secretory granule involves an apparent reduction in volume and surface area (Sesso et al., 1980), as well as an increase in dry mass concentration (Wong et al., 1991) that results in an extremely stable electron-dense core (Tooze and Tooze, 1986), which may be separated from the membrane by a ‘halo of space’ (Burgess and Kelly, 1987). It was interesting to note the dichotomy of granule morphology observed in the present study. Equal proportions of entirely electron-lucent granules (presumably immature granules) and granules that contained an electron-dense core surrounded by an electron-dense outer region (presumably mature granules) were observed in control mice. However, these ‘presumptive mature’ granules were the prevalent type of granule in G/A mice, probably due to increased storage time caused by the block on stimulated exocytosis. Conversely, presumptive immature granules were the predominant type of granule in G/A + B mice, indicating that it was presumptive ‘mature’ granules that were preferentially released after exocytotic stimulation. Despite immature granules being capable of stimulus-dependent secretion (Arvan et al., 1991; Tooze et al., 1991), it is possible that mature granules have a higher affinity for granule membrane–plasma membrane fusion that is required for exocytosis.

In conclusion, removal of GnRH input reduced LHβ mRNA content but did not affect basal plasma LH concentrations. In combination with increased storage of LH+/ve/SgII-ve granules, decreased storage of LH+/ve/SgII-ve granules and overall augmentation of intragranular anti-gercity of SgII indicate that SgII and LH actively co-aggregate to promote intracellular granule storage in the absence of GnRH input. In addition, LH+/ve/SgII-ve granules appeared to have been destined for constitutive release and may have functioned to maintain basal LH concentrations, whereas LH+/ve/SgII-ve granules appeared to have been sorted into the regulated pathway and were under the direct influence of GnRH. The absence of GnRH action on the pituitary did not alter FSHβ mRNA content but resulted in attenuated plasma FSH concentrations, indicating that a proportion of FSH had been sorted into storage at this time. Subsequent stimulation by Buserelin caused a marked release of intracellular LH+/ve/SgII-ve granule stores, reflected as a detectable increase in both plasma LH and FSH concentrations, whereas LH+/ve/SgII-ve granules were not released.

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