Autocrine inhibition of milk secretion in the lactating tammar wallaby (*Macropus eugenii*)

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ABSTRACT

The lactating tammar wallaby progressively alters the rate of secretion and composition of its milk to provide appropriate nutrition for the developing offspring, whose needs are signalled by changes in the pattern and efficiency of its sucking. Tammars are also capable of asynchronous concurrent lactation, when the mother provides a dilute milk for a newborn young permanently attached to the teat (phase 2A of lactation), and a concentrated milk from an adjacent mammary gland for a young-at-heel (phase 3). The relationship between suckling behaviour and milk secretion, and the ability of adjacent glands to function independently, suggests that milk secretion is controlled locally, within each mammary gland, by a mechanism sensitive to frequency and completeness of milk removal. To determine if tammar milk contains a factor able to control milk secretion, milk fractions have been screened in tissue and cell culture bioassays. A 6–30 kDa fraction of phase 3 whey was found to inhibit milk constituent synthesis and secretion *in vitro*, and inhibitory activity was associated with two discrete fractions obtained by anion exchange chromatography, which contained protein bands migrating anomalously at 66 kDa and 63 kDa in SDS-PAGE. These bands were recognised in Western blotting by antiserum raised against a bovine autocrine inhibitor of milk secretion. By the same criteria, milk secreted in phase 2B of tammar lactation, when milk secretion is low and suckling intermittent but less vigorous than phase 3, also contained a feedback inhibitor of milk secretion. The results indicate that, as in dairy animals, marsupial milk secretion is under local control through feedback inhibition by a milk protein, and raise the possibility that autocrine feedback may influence the transition from phases of low milk secretion (phase 2A, 2B) to a high rate in the final third phase of lactation.

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INTRODUCTION

Marsupials have adopted a reproductive strategy based on a short gestation with the birth of an immature young. This is followed by extensive growth and physiological development of the dependent young during a comparatively long period of lactation (Tyndale-Biscoe & Janssens 1988). The need to provide appropriate nutrition for this development requires that marsupials such as the tammar wallaby (*Macropus eugenii*) progressively alter their rate of milk secretion (Dove & Cork 1989), and the composition and concentration of all the major milk constituents (Green 1984, Green & Merchant 1988, Nicholas 1988a) during the course of a lactation.

Lactation in the tammar is divided into four distinct phases which correlate with changes in the secretion of specific milk proteins and changes in the suckling patterns and development of the pouch young (Nicholas *et al.* 1997). Phase 1 comprises mammary gland development during pregnancy and lactogenesis, which occurs at parturition. Phase 2A comprises the first 100 days of lactation when the young is in the pouch and permanently attached to the teat, and phase 2B describes the subsequent 100 days when the young remains in the pouch but relinquishes the teat and sucks less frequently. Phase 2 milk is dilute, with elevated levels of carbohydrate and low levels of fat and protein (Nicholas 1988a). The transition to phase 3 of lactation is characterised by an increase in the rate of milk synthesis, and the secretion of a concentrated milk high in protein and fat and low in carbohydrate. The young grows faster in phase 3 and leaves the pouch, begins to eat herbage, but
continues to suck the mammary gland. Induction of phase 3 lactation is accompanied by characteristic changes in milk protein gene expression. Late lactation protein (LLP)-A (LLP-A) is secreted from approximately day 160 of lactation and is maximal during phase 3, whereas LLP-B is expressed exclusively in phase 3 (Bird *et al.* 1994, Nicholas *et al.* 1994, 1997). The 18 kDa early lactation protein is expressed only during phase 2A (Nicholas *et al.* 1997), whereas whey acidic protein is expressed predominantly in phase 2B (Nicholas *et al.* 1995, 1997). In contrast, α-lactalbumin, α-casein, β-lactoglobulin and β-1,4 galactosyltransferase are present throughout phase 2B and phase 3 of lactation (Messer & Elliot 1987, Bird *et al.* 1994, Nicholas *et al.* 1994).

Tammar, and other macropodid marsupials, are capable of asynchronous concurrent lactation (ACL), whereby the mother provides a dilute milk for a newborn young continually attached to the teat of one mammary gland (phase 2A) whilst supplying a concentrated milk from another mammary gland (phase 3) for a young-at-heel which has vacated the pouch (Nicholas 1988b, Nicholas *et al.* 1994, 1995). The ability of adjacent glands to maintain ACL demonstrates that they are functioning independently, despite being within the same systemic hormonal milieu. Independent control of milk secretion rate in each mammary gland also occurs in other mammals. In ruminants, frequency of milking regulates the rate of milk secretion by a local, i.e. intramammary mechanism (Peaker 1995), through feedback inhibition of milk secretion by a protein named FIL (feedback inhibitor of lactation; Wilde *et al.* 1995a). FIL is synthesised by the mammary epithelial cell and, once secreted in milk, acts on the same cells to inhibit constitutive secretion in a concentration-dependent and reversible manner (Rennison *et al.* 1993, Wilde *et al.* 1995b). Thus, the endocrine regulation of lactation is modulated by an autocrine mechanism which acts to match milk supply with demand. Local control of milk secretion by milk removal is not peculiar to ruminants; in human lactation, milk secretion rate in each breast is regulated according to completeness of breast emptying at the previous feed (Daly *et al.* 1993). This too may occur through feedback inhibition by a milk protein since screening of human milk constituents in mammary tissue cultures has identified a milk protein fraction which inhibits milk constituent synthesis (Prentice *et al.* 1989). In the tammar, induction of phase 3 lactation is accompanied by a change to less frequent but more vigorous (and presumably more efficient) sucking by the young. Therefore, one possible explanation for the difference between adjacent mammary glands in the tammar is that gland development and the increase in milk secretion is led by the offspring through changes in the pattern and completeness of milk removal. To investigate this possibility, we have screened tammar milk in mammary tissue and cell cultures, to determine if it contains a factor able to exert feedback inhibition on milk secretion.

**MATERIALS AND METHODS**

**Materials**

Culture media were obtained from Northumbria Biologicals (Cramlington, UK) or Life Technologies (Uxbridge, UK). Collagenase (Worthington type III, 150 U/mg) was from Lorne Laboratories (Reading, Berks, UK). Hormones, antibiotics, calf thymus DNA and general laboratory chemicals were from Sigma Chemical Company, Poole, Dorset, UK. [U-14C] glucose was from Amersham International, Little Chalfont, Bucks, UK. L-[4,5-3H]leucine was from ICN Biomedicals Ltd, High Wycombe, UK. Ultrafiltration membranes were from Amicon Ltd, Stonehouse, UK.

**Animals**

Tammar wallabies were bred and kept in open grassed yards at the Victorian Institute of Animal Science, Attwood, Victoria, Australia. The animals had free access to water and supplements of lucerne, hay and oats. The stage of lactation was determined from known birth dates or by measurement of the head length of pouch young (Poole *et al.* 1991). Mid-pregnant New Zealand White and non-pregnant female Dutch rabbits were from Hi-Line, Lymm, Cheshire, UK. Mice were Tuck’s no. 1 strain (H Tuck & Son, Battlebridge, UK).

**Milk sampling and fractionation**

Milk samples were collected from 13 tammars milked at 2–3 weekly intervals over a 6 month period encompassing phase 3 of lactation. In addition, 20 animals were sampled on one occasion only during phase 2B of lactation (day 150–170 of lactation). The animals were caught in the field, separated from their young for 4 h and milked under short-term general anaesthesia (mehexitone sodium (Breital); Eli Lilly, West Ryde, Australia) after intramuscular injection of 0.1 U of oxytocin (Heriot Agvet Pty Ltd, Rowville, Australia). Milk samples were immediately frozen at −20 °C and shipped to the UK on dry ice. The average volume of milk collected in phase 2B was 0.5 ml.
Bioassay of milk fractions

Ultrafiltered whey and anion exchange fractions thereof were tested for their ability to inhibit casein and lactose synthesis in mammary tissue explants prepared from mid-pregnant rabbits. Groups of 30 explants were cultured in Medium 199 containing bovine insulin (5 µg/ml), hydrocortisone (0·1 µg/ml) and ovine prolactin (1 µg/ml) for 2 h in the presence or absence of anion exchange fractions (5–30 µg protein/ml; three replicates per experiment). The culture was terminated by centrifugation of the cell suspension (500 g, 1 min), and the cell pellet and supernatant were frozen and stored at −20 °C. Radiolabel incorporated in cellular and secreted protein was assayed by precipitation with trichloroacetic acid (final concentration 10%, w/v). Cell lysates were prepared by sonication (Kontes KT50 cell disrupter, Uxbridge, UK; setting 20, 15 s) in 0·1 M NaH2PO4 pH 7·4 containing 2 M NaCl and assayed for DNA content (Labarca & Paigen 1980). Synthetic and secretory activity were expressed per unit DNA.
Protein analysis

The composition of tammar whey fractions was analysed by SDS-PAGE in 10% gels under reducing conditions (Laemmli 1970). Gels were stained with Coomassie Blue or blotted onto PVDF membrane (Millipore) using standard methods (Towbin et al. 1979). Membranes were blocked for a minimum of 1 h at 4 °C with 4% (w/v) gelatin in PBS containing 0·1% (v/v) Tween 20 (PBST) and incubated for 2 h at room temperature with polyclonal rabbit antisera to tammar α-lactalbumin (1:500, Maher & Nicholas 1987), tammar LLP-A (1 µg IgG/ml; Nicholas et al. 1994), ovine β-lactoglobulin (1:500; Osborne et al. 1995) or bovine FIL (1:500; Wilde et al. 1995a). Blots were washed three times in PBST, and developed by sequential incubation with goat anti-(rabbit Ig)–alkaline phosphatase conjugate and bromochloroindolyl phosphate/nitro blue tetrazolium substrate (Sigma) according to the manufacturer’s instructions. When only a small amount of protein was available, as with phase 2B milk, fractions were resolved in high density Phastgels and silver stained using the Phast silverstain kit (Pharmacia).

ELISA

Whey fractions obtained by anion exchange chromatography were diluted to 1 µg/ml in coating buffer (15 mM Na2CO3, 35 mM NaHCO3, 3 mM NaN3, pH 9·6), and used to coat microtitre plates (Dynatech, Billingshurst, UK) by incubation at 4 °C for 16 h. The wells were blocked with 0·5% (w/v) gelatin in the same buffer, washed with PBST then incubated at room temperature for 2 h with 100 µl PBST containing polyclonal antisera against tammar α-lactalbumin (1:500 dilution), tammar LLP-A (1 µg IgG/ml) or ovine β-lactoglobulin (1:500 dilution). After repeated washing, binding was determined by incubation with anti-(rabbit Ig)–horseradish peroxidase conjugate (Scottish Antibody Production Unit, Carluke, UK) and O-phenylenediamine in citrate phosphate buffer (0·1 M NaH2PO4, 2H2O, 50 mM citric acid, pH 5·5) containing 0·012% H2O2. The absorbance was measured at 492 nm using a Labsystems MCC platereader (Helsinki, Finland).

Statistical analysis

Results are presented as means ± S.E.M. for n ≥ 3. Statistical significance was determined by general linear mean analysis or Student’s paired t-test.

RESULTS

Tammar milk fractions were initially screened for an inhibitor of milk secretion in rabbit mammary explants cultures, a system used previously to identify an autocrine regulator of milk secretion in goat’s milk (Wilde et al. 1995a). When tested at equal protein concentration, inhibition of lactose synthesis was observed with phase 2B but not phase 3 whey (Table 1). Preparation of 6–30 kDa whey fractions, and exclusion of a greater proportion of phase 3 tammar whey constituents of >30 kDa had no effect (Table 1). Based on these initial observations, the 6–30 kDa whey constituents from phase 3 milk were resolved by anion exchange chromatography (Fig. 1), and a
A series of fractions covering the entire salt gradient were tested for their ability to inhibit protein secretion in suspension cultures of mouse mammary acini. Bioassay in acini culture measured essentially the same response as tissue explant culture, i.e. the ability of milk fractions to inhibit synthesis and secretion of key milk constituents, and comparison of key milk fractions in each system confirmed that they produced similar effects (results not shown).

Bioassay of fractions from three individual animals showed inhibitory activity in acini consistently associated with fractions in the range 0–73 mM salt (Fig. 2). Other fractions constituting the majority of the whey proteins had no significant effect. Similar results were obtained with two samples of phase 3 milk pooled from several animals (results not shown).

SDS-PAGE of the inhibitory fractions of phase 3 whey showed that each contained a mixture of proteins, with several protein bands common to two or more fractions (Fig. 3). To obtain fractions containing discrete populations of proteins, proteins eluting at 0–73 mM salt were collected according to their A_280 elution profile. With this strategy, inhibition of protein secretion in acini cultures was associated with fractions designated 1 and 4 (Fig. 4), which eluted at 35–45 mM and 60–73 mM NaCl respectively (Fig. 1).

The protein composition of inhibitory fractions 1 and 4 was assessed by SDS-PAGE and comparison with the known molecular weights of tammar milk proteins (Nicholas et al. 1997), by immunoblotting and by non-competitive ELISA using specific antibodies. Resolution of fraction 1 by SDS-PAGE identified prominent bands migrating at 66 and 63 kDa (Fig. 5A), and in non-competitive ELISA the fraction showed weak reactions with antisera against β-lactoglobulin (18 kDa) and with LLP-A (24 kDa), a major whey protein in phase 3 tammar milk (Nicholas et al. 1987) (Fig. 6). This antiserum probably also recognises the B isoform of LLP (Nicholas et al. 1987). Fraction 4 showed similar 66 and 63 kDa bands on SDS-PAGE and a strong band of 18 kDa (Fig. 5A) which, based on ELISA...
Inhibition of mammary secretion by fractions 1 and 4 is, however, unlikely to be due to LLP or β-lactoglobulin. LLP-A and LLP-B were found predominantly in fraction 5 (Fig. 6), which was not inhibitory in the bioassay (Fig. 4). Likewise, the β-lactoglobulin content of fraction 3 was similar to that of fraction 4 and in excess of that in fraction 1 (Fig. 6), but fraction 3 did not inhibit protein secretion in mammary acini cultures (Fig. 4). On the other hand, the 63 kDa band common to fractions 1 and 4 was recognised in Western blotting by a polyclonal antiserum raised against a bovine FIL (Wilde et al. 1995a) (Fig. 5B). Anti-FIL antiserum showed negligible reaction with fractions 2 and 3, which were inactive in cell culture bioassay (Fig. 4). The presence of high molecular weight bands in fractions 1 and 4 was unexpected, since the material was derived from ultrafiltered whey, and contrasted with the fractions' elution profiles during gel filtration chromatography. Gel filtration of inhibitory fractions 1 and 4 confirmed the absence of molecules >30 kDa, and showed only one protein of 5-7 kDa common to the two fractions (results not shown).

Phase 3 and phase 2B tammar milk differ in protein composition (Nicholas et al. 1997) but, as with phase 3 milk, whole whey from phase 2B milk and a 6–30 kDa fraction thereof inhibited casein and lactose synthesis in rabbit explant cultures (Table 1). The small volumes of milk obtainable at this stage of lactation (~0.5 ml per animal) and the low protein content of phase 2B whey, reflected in the anion exchange elution profile of the 6–30 kDa fraction (Fig. 7A), precluded extensive bioassay of anion exchange fractions. However, using pooled material from ten animals, acinar protein synthesis and secretion were inhibited by an anion exchange fraction eluting at 35–80 mM NaCl, similar to the elution position of phase 3 fractions 1 and 4 (Fig. 7B). Western blotting of phase 2B whey fractions with anti-FIL antiserum also indicated the presence of an inhibitory protein in phase 2B milk. SDS-PAGE showed the presence of a 66 kDa component in protein eluting at 55–80 mM NaCl during anion exchange chromatography, and the study, is likely to be β-lactoglobulin (Fig. 6). Inhibition of mammary secretion by fractions 1 and 4 is, however, unlikely to be due to LLP or β-lactoglobulin. LLP-A and LLP-B were found predominantly in fraction 5 (Fig. 6), which was not inhibitory in the bioassay (Fig. 4). Likewise, the β-lactoglobulin content of fraction 3 was similar to that of fraction 4 and in excess of that in fraction 1 (Fig. 6), but fraction 3 did not inhibit protein secretion in mammary acini cultures (Fig. 4). On the other hand, the 63 kDa band common to fractions 1 and 4 was recognised in Western blotting by a polyclonal antiserum raised against a bovine FIL (Wilde et al. 1995a) (Fig. 5B). Anti-FIL antiserum showed negligible reaction with fractions 2 and 3, which were inactive in cell culture bioassay (Fig. 4). The presence of high molecular weight bands in fractions 1 and 4 was unexpected, since the material was derived from ultrafiltered whey, and contrasted with the fractions' elution profiles during gel filtration chromatography. Gel filtration of inhibitory fractions 1 and 4 confirmed the absence of molecules >30 kDa, and showed only one protein of 5-7 kDa common to the two fractions (results not shown).

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Journal of Molecular Endocrinology (1998) 21, 169–177

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principally in fraction c at 65–70 mM (Fig. 8A). This band was recognised specifically by anti-FIL antiserum (Fig. 8B). The presence of LLP-A in fraction c, and also in the principal phase 2B peak, designated e, was shown by non-competitive ELISA, and reflects the late stage of phase 2B lactation necessary to obtain sufficient milk for analysis (results not shown).

DISCUSSION

The change in suckling pattern during the transition from phase 2B to phase 3 of lactation (Tyndale-Biscoe & Janssens 1988) suggests that, as in other species, milk secretion rate in the tammar wallaby may be dependent on frequency and completeness of milk removal. In this study, tissue and cell
culture bioassay showed that phase 3 tammar milk contains proteins able to inhibit the secretion of milk constituents. Both proteins cross-reacted with antisera raised against a bovine autocrine inhibitor of lactation (Addey et al. 1995, Wilde et al. 1995a), suggesting that they may perform a similar function in the tammar, i.e. marsupial milk secretion may be under autocrine control through feedback inhibition by a milk constituent, as it is in dairy animals (Wilde et al. 1995a,b). The structure of the inhibitory proteins in tammar milk has yet to be determined. However, it was notable that they, like the FIL proteins isolated from goat’s and cow’s milk, behaved anomalously in SDS-PAGE.

Although previously ultrafiltered through a 30 kDa cut-off membrane, the FIL-related proteins in tammar milk migrated in SDS-PAGE with apparent molecular masses of 66 and 63 kDa. The caprine FIL also migrates at ~66 kDa in SDS-PAGE, but passes through the 30 kDa ultrafiltration membrane, and elutes on gel filtration with a molecular mass of 7.4 kDa (Wilde et al. 1995a). Similarly, gel filtration of inhibitory fractions 1 and 4 has confirmed the absence of molecules of >30 kDa and shown only one protein of 5.7 kDa common to the two fractions (results not shown). Thus, the presence of high molecular weight bands in ultrafiltered tammar phase 3 whey may be explained by the characteristically anomalous behaviour of molecules which share an ability to inhibit secretion in mammary culture bioassays.

The inhibitory proteins in phase 3 tammar milk may provide a mechanism for matching the mother’s milk supply with the changing demand of the developing young as its growth accelerates, and its diet becomes increasingly more herbivorous (Tyndale-Biscoe & Janssens 1988). That is, feedback inhibition may act to regulate milk secretion rate according to frequency or completeness of milk removal from the mammary gland (Peaker & Wilde 1996). Control of milk secretion rate may be signalled by changes in FIL’s concentration in milk. Thus, addition of purified caprine FIL to milk stored in one goat mammary gland decreased milk secretion rate temporarily, and in a concentration-dependent manner (Wilde et al. 1995a). Conversely, immunological neutralisation of FIL in goat’s milk elicited a temporary increase in milk secretion rate (Wilde et al. 1996). Alternatively, or in addition, autocrine feedback may be modulated by changing the activity of a receptor-mediated signal transduction pathway (Wilde et al. 1997a). Whichever is the case, the intracellular effect of autocrine feedback is to block constitutive secretion in the mammary epithelial cells (Rennison et al. 1993). The inhibitory activity of phase 2B tammar milk fractions in tissue and cell culture bioassays suggests that feedback inhibition may also control milk secretion during the period when the young is intermittently attached to the teat. Inhibition was, as in phase 3, associated with 6–30 kDa whey proteins, and Western blotting showed that the activity was ascribable to the presence of a FIL-like protein in milk. Tammar milk proteins, like milk composition itself, are regulated asynchronously throughout lactation. The expression and concentration of some milk proteins change significantly, whereas others are expressed at relatively constant levels (Messer & Elliot 1987, Bird et al. 1994, Nicholas et al. 1994, 1995, 1997). If the expression or milk concentration of tammar FIL is amongst those which changes during the transition to phase 3, it could be an important determinant of the increase in milk secretion and, by analogy with other species (Peaker & Wilde 1996, Wilde et al. 1997b), the developmental adaptations which act to sustain it. Bioassay suggested that, when compared at equal concentrations, the whey protein fraction of phase 2B milk was indeed more inhibitory than that derived from phase 3 milk. On this basis, and taking into account the concentration of whey proteins in phase 2B and phase 3 milk (15-7 and 30-9 mg/ml respectively), inhibition compared on a milk volume basis is calculated to be >10-fold higher in phase 2B milk. Thus, the transition to phase 3 of lactation may be promoted by a decrease in FIL-mediated autocrine feedback on milk secretion.

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