Variation in fat, lactose and protein in human milk over 24 h and throughout the first year of lactation

Leon R. Mitoulas1*, Jacqueline C. Kent1, David B. Cox1, Robyn A. Owens2, Jillian L. Sherriff3 and Peter E. Hartmann1

Departments of 1Biochemistry and 2Computer Science, The University of Western Australia, 35 Stirling Highway, Crawley, WA 6009, Australia
3School of Public Health, Curtin University, GPO Box U1987, Perth, WA 6845, Australia

(Received 30 July 2001 – Revised 2 January 2002 – Accepted 30 January 2002)

Fat in human milk is extremely variable and can represent up to 50 % of infant energy intake. To accurately determine milk composition and infant intake at 1 (n 17), 2 (n 17), 4 (n 17), 6 (n 15), 9 (n 6) and 12 (n 5) months of lactation, samples of fore- and hind-milk were collected from each breast at each feed over 24 h periods from an initial group of seventeen women. The content of fat in milk varied over 24 h, with a mean CV of 47·6 (SE 2·1) % (n 76) and 46·7 (SE 1·7) % (n 76) for left and right breasts respectively. The 24 h amounts of fat, lactose and protein in milk differed between women (P=0·0001), but were consistent between left and right breasts. Daily milk production differed between breasts (P=0·0001) and women (P=0·0001). Accordingly, amounts of fat (P=0·0008), lactose (P=0·0385) and protein (P=0·0173) delivered to the infant over 24 h also differed between breasts and women (P=0·0001). The energy content of milk and the amount of energy delivered to the infant over 24 h were the same between breasts, but differed between women (P=0·0001). The growth rate of a group of only six infants in the present study was not related to either the concentrations or amounts of fat, lactose, protein and energy in milk over the first 6 months of life. These results show the individuality of milk composition and suggest that only a rigorous sampling routine that takes into account all levels of variation will allow the accurate determination of infant intake of fat, lactose, protein and energy.

Milk fat: Infant intake: Sampling routine: Human lactation

Of the major digestible energy components (fat, lactose and protein) in human milk, fat is the most variable. Woolridge (1995) listed several factors that either individually or in concert could account for the variability in fat content of human milk. Major factors included the amount of milk removed at both the last and current breast-feed, the length of the interval between breast-feeds, and the fat content at the end of the last breast-feed. Daly et al. (1993a) showed that approximately 70 % of the variation in fat content of breast milk was due to the extent of fullness of the breast (see Cox et al. 1996), essentially incorporating all the predictors proposed by Woolridge (1995) and expressing them as one term. Furthermore, the discovery of local (autocrine) control systems for milk synthesis (Henderson & Peaker, 1984) and possibly milk-fat synthesis (Heesom et al. 1992), combined with differences in milk production and storage capacity between breasts within mothers (Daly et al. 1993a), could result in different rates of milk and fat synthesis between breasts. These factors make it difficult to design a sampling protocol, suitable for all women, that will provide a true indication of energy density and intake of breast milk without affecting the natural routine of the demand-fed infant (Prentice & Prentice, 1988; Lucas & Davies, 1990; Prentice et al. 1996).

We have used a sampling protocol similar to that of Hartmann et al. (1986) that takes into account changes in fat content of milk during a feed, differences between breasts, changes over the course of the day and ensures minimum interference with infant feeding behaviour. We have determined the volume of milk removed together with the fat content, lactose and protein concentrations and the calculated energy content for milk from each breast at each feed over a 24 h period at 1, 2, 4, 6, 9 and 12 months of lactation for women who were breast-feeding. These data were used to determine mean 24 h concentrations.

* Corresponding author: Leon R. Mitoulas, fax +61 8 9380 1148, email Leon.Mitoulas@uwa.edu.au
in milk and amounts of each component delivered to the infant from breast milk from 1 to 12 months after birth.

**Methods**

**Subjects**

Healthy mothers and infants were recruited through the Nursing Mothers’ Association of Australia, Western Australian Branch, or private health care centres. All mothers supplied written informed consent to participate in the study, which was approved by the Human Research Ethics Committee, The University of Western Australia. Subject details have been previously reported (Cox et al. 1996, 1999). Briefly, mothers were between 18 and 35 years of age, twelve were multiparous (five mothers having had two children, seven mothers having had three children) and five primiparous, with a mean weight of 68.69 (SE 2.06; range 55.3–84.4) kg (n 17). All infants were born at term (except one, born at 31 weeks) and were exclusively breast-fed on demand for at least 4 months, with complementary solid foods being introduced between 4 and 6 months of age. All mothers maintained their own breast-feeding patterns throughout study periods. All study periods were within 1 week of the indicated month of lactation.

**Milk sampling**

Milk samples (≤ 1 ml) were collected before and after each feed from each breast by either manual breast pump (Kaneson Expression and Feeding Bottle; Yanase Waitch KK, Osaka, Japan) or hand expression into 5 ml polypropylene vials (Disposable Products Pty Ltd, Adelaide, Australia). Samples were initially stored in a household freezer for a maximum of 24 h and then transported on ice to the laboratory where they were stored at −20°C until analysed.

**Biochemical analyses**

**Milk fat.** The content of fat in fore- and hind-milk samples was determined using the modified colorimetric spectrophotometric method of Stern & Shapiro (1953), as previously described (Cox et al. 1996). Briefly, 2.5 μl portions of the milk samples (warmed to 37°C) and standards (triolein, 0–200 mM) were added to redistilled ethanol (600 μl) and mixed for 10 s. Hydroxylamine hydrochloride (2 μl; 100 μl) and NaOH (3.5 μl; 100 μl) were then added to each sample and the samples mixed and left to stand at room temperature for 20 min. Each sample was acidified by the addition of HCl (4 μl; 100 μl) and colour production achieved by the addition of a FeCl3–TCA solution (7.5 g TCA in 10 ml 0.37 M-FeCl3–0.1 M-HCl; 100 μl). The tube contents were mixed and 250 μl from each tube was pipetted into duplicate wells on a ninety-six-well microtitre plate. Absorbance of each well was determined at 540 nm using a plate spectrophotometer (Titertek Multiscan MCC/340; Flow Laboratories, McLean, VA, USA). The detection limit of this assay was 0.45 (SE 0.41) g/l (n 17). The assay CV was 8.1 % (n 13).

**Milk lactose.** The concentration of lactose in fore- and hind-milk samples was determined using the modified method of Kuhn & Lowenstein (1967), as described by Arthur et al. (1989). Briefly, defatted milk samples and lactose standards (0–300 mM) were diluted 1 in 150 with distilled deionised water. Duplicate portions of diluted samples and standards (5 μl) were pipetted into wells on a flat-bottom ninety-six-well microtitre plate and reagent 1 (8 U B-galactosidase/ml, 0.1 M-MgCl2 in 0.1 M-potassium phosphate buffer, pH 7.2; 50 μl) was added to each well and the plate mixed and incubated at 37°C for 60 min. Following this step, reagent 2 (9.6 U glucose oxidase/ml, 2.5 U peroxidase/ml, 500 μg 2,2-azino-di-(3-ethyl-benzthiazolin-sulfonate)-6-sulfonate/ml in 0.1 M-potassium phosphate buffer, pH 7.2; 200 μl) was added to each well and the absorbance measured at 405 nm on a plate spectrophotometer at 5 min intervals until a peak absorbance was reached at approximately 45 min. The recovery of a known amount of lactose added to milk samples was 102 (SE 1) % (n 17). The detection limit of this assay was 16.4 (SE 0.4) g/l (n 35) and the inter assay CV was 5.6 % (n 35).

**Milk protein.** The concentration of protein in fore- and hind-milk samples was determined using a commercial protein assay kit (Bio-Rad Laboratories, Richmond, CA, USA). The assay procedure was a modification of that of Atwood & Hartmann (1992), in that samples were diluted 1 in 30 with double-deionised water. To overcome the problems inherent in the choice of a milk standard the protein concentration of an aliquot of mature breast milk was determined by the Kjeldahl procedure (Hambraeus et al. 1978), as described by Atwood & Hartmann (1992). The remaining sample was then diluted with double deionised water to provide a range of standards (0–1 g/l). Briefly, defatted milk samples were diluted 1 in 30 with double-deionised water and pipetted (5 μl) in duplicate, with standards, onto a ninety-six-well microtitre plate. To each well was added 250 μl Bio-Rad protein assay reagent (diluted 1 in 5 with distilled deionised water and filtered through Whatman no. 1 paper) and the plate mixed for 1–2 min and then allowed to stand for 5 min. Absorbance was then measured at 620 nm using a plate spectrophotometer. The recovery of a known amount of protein added to milk samples was 99.96 (SE 1.03) % (n 17). The detection limit of this assay was 0.033 (SE 0.002) g/l (n 53) and the inter assay CV was 4.72 % (n 35).

**Milk energy.** The energy content for each feed was calculated using the conversion factors (Garza et al. 1985) of 38.7, 16.5 and 23.7 kJ/g for fat, lactose and protein respectively.

**Determination of 24 h milk production**

Milk yield was determined for each breast by test weighing the mother, as described by Arthur et al. (1987). Test weighing was carried out at each mother’s home over a 24–28 h period using an electronic Sauter balance (weighing platform, Model EC 240; evaluator unit with data output printer, Model ED 3300; FSE Scientific, Perth, WA, Australia). Briefly, mothers weighed themselves before and after each feed from each breast. To account for the insensitive water loss, which occurred during feeding,
mothers were instructed to reweigh themselves 20 min after the end of each feeding session. The rate of water loss for this 20 min period was then used to calculate insensitive water loss during the feeding period.

**Determination of 24 h nutrient and energy intake**

Fore- and hind-milk concentrations were averaged to provide a concentration of fat, lactose, protein and energy for each feed. The volume of the feed was then used to determine the amount taken by the infant and the total energy provided at each feed. The sum of the amount of each nutrient and the energy provided for all feeds over the study period (24–28 h) and the total volume delivered over the study period were then used to determine an average concentration. This concentration and the corrected 24 h volume (Arthur et al. 1987) were then used to determine the amount delivered in 24 h to the infant.

**Infant growth rates**

Infant growth rates were determined as previously described (Kent et al. 1999). Briefly, subjects provided records of birth weights and weight gains up to 6 months of age, as measured by midwives attending the birth and Community Health Nurses respectively.

**Statistical analysis**

Seventeen mothers initially provided data for milk production and fat content whereas lactose, protein and energy data were initially obtained for only nine of the seventeen mothers (Table 1). For all metabolites the number of mothers decreased as the study progressed, due to the cessation of the collection of samples from eleven mothers after 6 months of lactation. Storage capacity and average feed volumes were determined for only six mothers up to 6 months of lactation, the maximum period of exclusive demand breast-feeding. In all cases left and right breasts were treated separately, therefore n, unless otherwise stated, represents the number of individual breasts sampled.

All longitudinal analyses were performed using The SAS System for Windows v6.12 (SAS Institute Inc., Cary, NC, USA) with the general linear means (PROC GLM) procedure. Student’s paired t tests and other statistics were performed using Statview™ SE+Graphics (Abacus Concepts Inc., Berkeley, CA, USA). All values are reported as means with their standard errors, unless otherwise stated.

**Results**

**Variation of fat, lactose and protein in human milk over 24 h**

The fat content of hind-milk was significantly higher than that of fore-milk (P<0.05), but there were no significant differences for either lactose or protein. The mean CV (%) in fore- and hind-milk samples collected from left and right breasts for all mothers was 47·6 (SE 2·1) and 46-7 (SE 1-7) for fat (n 76), 9-86 (SE 1-66) and 8-37 (SE 1-21) for lactose (n 46), and 11-9 (SE 1-2) and 12-3 (SE 1-1) for protein (n 46) respectively.

The fat content of fore- and hind-milk samples obtained from an irregular feeding pattern (mean feed volume 60 (SD 35) ml) from breasts with larger storage capacities (an example is shown in Fig. 1(A), storage capacity 271 ml) varied more than the fat content of fore- and hind-milk samples for more regular feeding patterns (mean feed volume 73 (SE 30) ml) from breasts with smaller storage capacities (an example is shown in Fig. 1(B), storage capacity 124 ml). The standard deviation of the mean fat content of each feed over a 24 h period, as a measure of variability, was negatively correlated with the mean feed volume when expressed as a percentage of the storage capacity (Fig. 2; P<0.05, r = −0.319, n 44).

**Variation between left and right breasts**

Milk production (Fig. 3) was found to differ significantly between left and right breasts at 1, 2, 4 and 12 months of lactation (P<0.05). Overall, the mean 24 h milk production for the left breast was 356 (SD 129) ml and for the right breast was 443 (SD 141) ml.

There were no differences in either the contents of fat and energy, or concentrations of lactose and protein between milk from left and right breasts from 1 to 12 months of lactation (Fig. 4). Although there were some significant differences between the left and right breasts in the amounts of fat, lactose and protein delivered to the infant, there were no significant differences between left and right breasts in the amount of energy delivered to the infant at any stage of lactation (Fig. 4).

Milk production of left and right breasts was consistently different (P<0.05) for five of the mothers over the first 6 months of lactation. For these mothers the breast that produced more milk was termed the ‘preferred’ breast (Fig. 5). There was no difference in the fat content of milk between preferred and non-preferred breasts. Nevertheless, the amount of fat delivered to the infant was greater (P=0·005) from the preferred breast.

**Variation over the first year of lactation**

The composition and volume of milk removed from each breast over the first 12 months of lactation is shown in Table 2. The mean volume of milk produced per breast from 1 to 12 months of lactation was 399 (SE 11) ml/
24 h. Although there was no significant difference in milk production from 1 to 6 months of lactation, milk production at 12 months was significantly less than that at 6 months of lactation ($P < 0.05$; Table 2).

The fat content of milk and the amount of fat delivered in 24 h per breast over the first 12 months of lactation were $37.4$ (SE 0.6) g/l and $14.8$ (SE 0.5) g respectively. The fat content of milk differed greatly between women ($P < 0.0001$), with some mothers producing milk with a fat content of $50$ g/l, while in others it was $35$ g/l at all stages of lactation (Fig. 6). Mean fat content showed marked differences over the 12 months of lactation ($P < 0.0001$), decreasing from $39.9$ (SE 1.4) g/l at 1 month to $35.2$ (SE 1.4) g/l at 2 months and then increasing again to $40.7$ (SE 1.7) g/l at 9 months. In contrast, the mean amount of fat delivered to the infant did not change with stage of lactation.

The mean concentration of lactose and protein in milk from 1 to 12 months of lactation was $61.4$ (SE 0.6) g/l and $9.2$ (SE 0.2) g/l respectively (Table 2), but there were significant differences ($P < 0.0001$) between women. The concentration of lactose in milk did not change with stage of lactation, whereas the concentration of protein decreased from $10.5$ (SE 0.4) g/l at 1 month to $8.0$ (SE 0.4) g/l at 6 months, and then remained steady. The amount of lactose and protein delivered to the infant (Table 2) differed between women ($P < 0.0001$) and declined with stage of lactation ($P < 0.0253$).

The energy content and amount delivered to the infant was $2.65$ (SE 0.04) kJ/ml and $1007$ (SE 39) kJ per breast respectively. Both the energy content and amount delivered to the infant differed between women ($P < 0.0001$). However, only the energy content differed with stage of lactation ($P = 0.0001$), decreasing from $2.7$ (SE 0.1) kJ/ml at 1 month to $2.5$ (SE 0.1) kJ/ml at 2 months and then increasing to $2.8$ (SE 0.1) kJ/ml at 9 months.
Growth rates for six infants were 20.4 (SE 2.0) g/d from birth for the first 6 months of life. No significant relationships were found between growth rate of the infants and either the amount taken in by the infant or the mean milk concentrations of fat, lactose, protein and energy for the first 6 months of lactation. Energy intake (kJ/kg body weight) from breast milk for four of the infants decreased significantly (P<0.0006) from 1 month (456 (SE 64)) to 6 months of lactation (268 (SE 33)).

**Infant growth**

Fig. 4. Milk fat content (A), amount of fat delivered over 24 h (B), milk lactose concentration (C), amount of lactose delivered over 24 h (D), milk protein concentration (E), amount of protein delivered over 24 h (F), milk energy content (G) and amount of energy delivered over 24 h (H) for left (○—○) and right (●—●) breasts from 1 month to 12 months of lactation. Differences at individual time points between left and right breasts are indicated, *P<0.05, **P<0.01, ***P<0.001. Values are means with their standard errors represented by vertical bars; n at each time point is shown in Table 1. For details of subjects and procedures, see p. 30.
A number of protocols have been employed to measure the variation in the fat content of milk both during a breast-feed and between breasts. These procedures range from the removal of milk from the non-feeding breast at
Milk composition and infant intake

Fig. 6. Daily fat content of milk from all mothers at 1, 2, 4, 6, 9 and 12 months of lactation. Values are mean (—) and standard deviations (...) and ranges, represented by vertical bars. For details of subjects and procedures, see p. 30.

alternate breast-feeds over the course of the day (Butte et al. 1984; Garza & Butte, 1986; Nommersen et al. 1991) to the collection of either random or timed milk samples (Hall, 1979; Lauber & Reinhardt, 1979; Bitman et al. 1983; Allen et al. 1991). We used a sampling protocol similar to that of Hartmann et al. (1986) that involved the collection of fore- and hind-milk samples and the measurement of the production of milk from each breast at each feed over a period of 24 h at 1, 2, 4, 6, 9 and 12 months of lactation to account for the variation in fat and to accurately determine the concentrations of fat, lactose, protein and energy in human milk from an initial group of seventeen mothers. Whilst the total number of mothers recruited may seem low it must be stated that this demanding sampling protocol involved the collection and multiple analysis of over 2000 milk samples, as well as requiring mother participation for a minimum of 6 months and a maximum of 12 months.

The maximum amount of milk that can be stored in the breast is available to the infant, storage capacity (Daly et al. 1993b; Kent et al. 1999), and the infant’s appetite indirectly influenced the content of fat in milk. For the breast with a large storage capacity (≥200 ml) the volume of each feed over the course of the day can vary greatly, with the infant rarely draining the breast at any one feed (Fig. 1(A)). This factor allows for less regularity in timing and volume of milk removed for each feed, and results in increased variation in the content of fat in fore- and hind-milk (Fig. 1(A)). Thus, any one feed is unlikely to approximate the daily average. On the other hand, for a breast with a small storage capacity (≤150 ml) the feed volume more closely approximates the storage capacity, and the variation between the fat content of fore- and hind-milk will be low (Fig. 1(B)) and is more likely to be representative of the daily average. Consequently, the collection of milk samples either randomly or at particular times of the day will be unrepresentative for women with larger storage capacities because of the variability exhibited both in the fat content of fore- and hind-milk and in the intake of milk at each breast-feed.

The significant difference in milk production between breasts (P<0.0001; Fig. 3) is in agreement with previous findings (Morrison, 1952; Hytten, 1954; Cox et al. 1996). The daily mean content of fat and energy together with the concentrations of lactose and protein in milk were found to be the same between breasts within women over the course of the study (Fig. 4(A, G, C and E respectively)). Furthermore, when left and right breasts were reclassified as preferred and non-preferred, based on the consistent differences (P<0.05) in milk production (Fig. 5(A)), no differences between breasts were found in the fat content of milk (Fig. 5(B)). This similarity in the daily mean composition of milk between breasts (either left or right, or preferred or non-preferred) within women validates those methods that remove milk from the non-feeding breast at alternate breast-feeds over the course of the day for the determination of milk composition. In addition, the finding that there was no significant difference between the mean 24 h content of fat in milk in each breast may reflect an additional level of control on milk synthesis in women. Apart from the endocrine and autocrine control mechanisms of milk synthesis and production (Hartmann et al. 1998) there may also exist a ‘metabolic’ level of control based on the homeorhetic model proposed by Bauman & Currie (1980).

The changes observed for all measured components over the first 12 months of lactation (Table 2) were similar to those reported previously. Milk production was constant for the first 6 months (Dewey & Lonnerdal, 1983; Hartmann et al. 1995; Cox et al. 1996), after which there was a steady decline (Neville et al. 1991). The fat content of milk decreased between 1 and 4 months (Butte et al. 1984), before increasing by 12 months of lactation (Ferris & Jensen, 1984; Allen et al. 1991). The concentration of protein in milk decreased by 6 months of lactation (Hytten, 1954; Prentice et al. 1981; Butte et al. 1984; Allen et al. 1991) and then remained constant (Neville et al. 1991), whereas the concentration of lactose remained constant throughout the first year of lactation (Hartmann & Kulski, 1978). The decrease in the energy content at 2 months and the subsequent increase by 9 months can be attributed to the changes in fat content of the milk (Fig. 5).

The minimum requirements for the estimation of the energy intake of the breast-fed infant are the measurement of milk production over a 24 h period together with the accurate determination of the average composition of the breast milk consumed by the infant over the same period (Hartmann et al. 1998). The mean total 24 h milk production from both breasts was 798 (SD 232) ml (Table 2), which is consistent with previous reports for milk production in women (Butte et al. 1984; Dewey et al. 1986; Hartmann et al. 1995; Cox et al. 1996). The mean fat, lactose and protein contents (g/l) of breast milk (37.4 (SE 0·6), 61·4 (SE 0·6), 9·16 (SE 0·19) respectively; Table 2) was determined by averaging the fore- and hind-milk
concentrations of fat, lactose and protein for each breastfeed from each breast over the 24 h period and weighting the values for the amount of milk consumed from each feed from each breast at 1, 2, 4, 6, 9 and 12 months of lactation (Picciano, 1984). Although these values are similar to those reported previously (Jensen et al., 1995), there was considerable variation about the means (CV 21.2, 9.3 and 19.9 % respectively). Thus, the concentrations of fat, lactose, protein, and energy, as well as milk production, differed significantly between women ($P=0.0001$), resulting in infants of the same age receiving different daily intakes. These results highlight the differences in milk composition between women and emphasise the inadequacy of using population averages of milk composition to determine either the intakes of individual breast-fed infants or the dietary requirements for lactation of individual mothers (Hartmann et al., 1995).

Growth rates over the first 6 months of life for the six infants monitored were not related to either the concentrations or amounts of fat, lactose, protein and energy in milk. Indeed, the only factor to have an effect on growth rate was milk intake (Kent et al., 1999). These results, from a small group of infants, are supported by the findings of a larger study (Butte et al., 1984), and together they highlight the importance of initially addressing milk intake by the infant and the possible mismanagement of breast-feeding, rather than questioning milk quality or composition, during the clinical treatment of slow-weight-gain infants (Lawrence & Lawrence, 1999).

Previous reports have shown energy intakes of formula-fed infants to be greater than those of exclusively breast-fed infants (Garza & Butte, 1990; Heiring et al., 1993). In the current study the increase in body weight between 1 and 6 months of age was obtained for four of the infants. For these breast-fed infants energy intake (kJ/kg body weight) at 1 month of age was not different from what they would have received if they had been fed formula, as directed by the manufacturer. However, at 6 months of age the energy intake from breast milk had decreased significantly ($P<0.05$), whereas that for formula did not. These results, obtained with a different sampling protocol, corroborate those of Heinig et al. (1993) and show, as expected, that each infant received less energy (kJ/kg body weight) as lactation progressed (Dewey & Lonnerdal, 1983). When combined with the findings of Kramer (1981) and von Kries et al. (1999) that breast-feeding has a protective effect against childhood obesity, these results, albeit from a small group of only four subjects, add to the dispute over the current recommendations for the energy intake for formula-fed infants, and further support the establishment of new dietary guidelines based on the energy intakes of breast-fed infants.

In conclusion, from our studies using an initial group of seventeen women we have found that milk composition does not differ between either left and right breasts or between preferred and non-preferred breasts. It is due to this factor that the results on milk composition obtained by the current study (using a rigorous sampling routine) are similar to those of previous studies, indicating that it is possible to determine population averages by a variety of sampling schedules. However, the finding that the daily variation in milk fat content together with average milk composition and production differ significantly between women shows that all other sampling schedules cannot provide an accurate indication of the intake of fat, lactose, protein and energy of the individual infant and must, therefore, be interpreted cautiously.

Acknowledgements

The authors wish to thank the mothers and babies who participated in this study, together with the Nursing Mothers’ Association of Australia. We would also like to thank Dijana Mihic and Tracey Williams for technical assistance and Dr Lyle Gurrin for statistical advice. This study was supported by the Grains Research and Development Corporation of Australia, Meadow Lea Foods Ltd, the Lotteries Commission of Western Australia, the Australian Research Council and the National Health and Medical Research Council of Australia.

References


Dewey KG, Finley DA, Strode MA & Lonnerdal B (1986) Relationship of maternal age to breast milk volume and composition. In Human Lactation 2: Maternal and Environmental...


