Newborn calf intestinal absorption of immunoglobulins extracted from colostrum

J.-F. GRONGNET, Elisabeth GRONGNET-PINCHON, D. LEVIEUX (*), M. PIOT (**), J. LAREYNIE

Summary. In the newborn calf, colostrum immunoglobulin absorption was compared with absorption of immunoglobulins extracted from colostrum. Reduced absorption of the latter resulted in plasma immunoglobulin levels unable to protect the animals against infections. Adding complete milk powder to the immunoglobulin solution slightly improved the level of absorption. Postprandial plasma alkaline phosphatases levels exhibited changes differing with the nature of the meals. This reinforces the idea that intestinal function was probably disturbed by immunoglobulin solutions. Moreover, reduced appetite for immunoglobulin solution was evident by the second meal, when most of the animals previously subjected to this diet suffered from a variety of problems (scour, asthenia), possibly caused by harmful effects of immunoglobulin solution on their digestive tract.

Introduction.

It is well known that copious ingestion of colostrum prevents sanitary problems in newborn ruminants and piglets (Dardillat, Trillat and Larvor, 1978). For colostrum to be effective, it must contain a sufficient amount of immunoglobulins (Stott and Fellah, 1983; Tshibangu et al., 1982) that are ingested soon after birth (Edwards et al., 1982; Stott et al., 1979) by lambs and calves which are able to absorb them. Very wide variations have been found between individual levels of plasma immunoglobulins after the ingestion of apparently similar quantities of pooled colostrum. These results led to studies on factors which might act on immunoglobulin absorption. Some authors considered the importance of environmental factors by studying:

- prepartum maternal feeding (Blecha et al., 1981; Grongnet, 1981; Halliday et al., 1978; Khalaf et al., 1979; Loh et al., 1971; Olson et al., 1981);
- ambient temperature (Blecha and Kelley, 1981; Cabello and Levieux, 1980; Kelley, Blecha and Regnier, 1982; Olson, Papasian and Ritter, 1980; Rafai et al., 1981; Stott et al., 1976; Stott, 1980);
- season of calving (Edwards, Broom and Collis, 1982; Frerking and Aekens, 1978; Gay, McGuire and Parish, 1983; Gonzalez, Villouta and Ferrando, 1976;
Selman, McEwan and Fisher, 1970; Smith, O’Neil and Simmons, 1967; Tshibangu et al., 1982;
- presence of the dam and methods of feeding colostrum (Fallon, 1978; Joly, 1981; Lomba et al., 1978; McCoy et al., 1970; Selman et al., 1971a, b; Smith, O’Neil and Simmons, 1967; Stott et al., 1979; Stott, 1980);
- size of the litter (Halliday, 1978; Logan and Irwin, 1977);

Other authors studied factors more closely related to the physiological state of newborn animals:
- prematurity and length of gestation (Cabello and Levieux, 1981a, b; George et al., 1979);
- adrenal hormones (Boyd and Hogg, 1981; Cabello and Levieux, 1980; Daniels, Hardy and Malinowska, 1973; Daniels et al., 1973; Gillette and Filkins, 1966; Halliday, 1959; Husband, Brandon and Lascelles, 1973; Johnston and Oxender, 1979; Morris and Morris, 1976; Patt and Eberhart, 1976; Rafai et al., 1981; Stott et al., 1976);
- thyroid hormones (Boyd and Hogg, 1981; Cabello and Levieux, 1981a, b; Cabello et al., 1983; Pethes, Frenyo and Rudas, 1982);
- sex (Halliday and Williams, 1979; Norman, Hohenboken and Kelley, 1981);

It should be noted that the results concerning most of these factors were often conflicting.

Such discrepancies also exist between results dealing with the influence of the physical-chemical features of the excipient mixed with the immunoglobulins. Their importance is far from negligible. Balfour and Comline (1962) noted a dramatic fall in absorption when immunoglobulins were administered in a saline solution, even if the solution was as well balanced ionically as colostrum. Luckily, a low molecular-weight protein, extracted from colostrum by these authors, restored absorption to a satisfactory level when added to their solution. Hardy (1969) established that adding potassium isobutyrate to a saline solution of immunoglobulins or polyvinylpyrrolidone increased intestinal absorption considerably; Baumwart et al., (1977), who added the same substance to colostrum, did not confirm this.

Among other products studied, only poly-L-arginine had a beneficial effect (Smith, Witty and Brown, 1968). All the others such as duodenal fluid (James and Polan, 1978), histamine (Patt et al., 1972), cow saliva (Balbierz et al., 1976), L-methionine and L-leucine (Smith and Pierce, 1967), large quantities of glucose (Lecce, 1966) or lactose (Werhahn, Klobasa and Butler, 1981) were ineffective or detrimental to absorption. It should be noted that a clear comparative interpretation of the results is doubtful, given the widely different experimental conditions used.

Moreover, if any colostral component exerts a positive and direct action upon immunoglobulin absorption (Balfour and Comline, 1962), the very strong antitrypsin activity of colostrum seems to affect it, but in an indirect way, by preventing
proteolysis. Once again, the results are very debatable (Baintner, 1973a, b; Chamberlain, Perry and Jones, 1965; Jensen and Pedersen, 1982).

Among the factors affecting immunoglobulin absorption by the newborn calf, lamb and piglet intestine, those concerning the other constituents of the solution are of great importance today because cheap, orally dispensable, immunoglobulins would be produced industrially in the near future. Their efficiency will, of course, depend on how well they can be absorbed.

The aim of the present work was to compare the absorption of immunoglobulins extracted from colostrum with that of immunoglobulins remaining in the colostrum.

Material and methods.

Animals. — Three groups (A, B, C) of newborn male and female Holstein Friesian calves, born from the same dairy herd, were used. Group A (40 calves) was born during the winter of 1979-1980, group B (32 calves) during the winter of 1980-1981, and group C (23 calves) during the winter of 1981-1982. All were born spontaneously at term and without dystocia. Calving was often slightly accelerated by a mild traction exerted on the fore limbs of the calf after it appeared in the vulva of the dam. No pharmacodynamic agents were used to facilitate parturition.

Diets. — Twenty-eight hours after birth and twice a day thereafter, the calves were fed with a standard milk replacer. Previous to this, they were fed four times, at exactly 4, 10, 16 and 22 h after birth. At those times, group A calves were fed on colostrum from the first milking of their dams. When several parturitions occurred at very short intervals, all the resulting colostra were mixed before feeding in order to homogenize the immunoglobulin levels as much as possible since there was no pooled colostrum already prepared; 25 g/kg body weight (BW) were given each time to each calf.

The four meals of group B animals consisted of an immunoglobulin solution (ISB); its basic composition is shown in table 1. This solution, using colostrum

<table>
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<th>TABLE 1</th>
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<td><strong>Elementary composition of immunoglobulin solutions. Comparison with colostrum.</strong></td>
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<tr>
<td><strong>Dry matter (%)</strong></td>
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<tr>
<td>Immunoglobulin solution ISB</td>
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<tr>
<td>Immunoglobulin solution ISC</td>
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<tr>
<td>Colostra (Group A, all dams ; mean ± SD)</td>
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<tr>
<td>Reference pooled colostra</td>
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<td>2 ( 8 cows)</td>
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* ITU/g : inhibited trypsin unit ; ** ND : not determined.
obtained at the first milking, frozen and thawed, was prepared by the following successive steps: — dilution (x 2); — removal of fat by centrifugation; — dilution (x 3); — casein coagulation at pH 4.6 by addition of HCl, 6N; — removal of the insoluble by centrifugation; neutralization by NaOH; — ultrafiltration, concentration and washing on Romicon, MP 100 (R) membrane (nominal separation level: 100 000 daltons) which retained the immunoglobulins in the retentate and removed α-lactalbumin and β-lactoglobulin in the ultrafiltrate.

At the end of washing, the retentate was frozen in units of one liter. After each birth, it was thawed by warming in a mild water bath at not more than 40-50 °C. Considering the level of immunoglobulins G (IgG) in the retentate (83.6 g/l) and the average level of IgG in the colostra used for group A (98.6 ± 23.6 g/l), the delivery of the retentate at the rate of 29 g/kg BW allowed group B calves to receive as much immunoglobulin as group A calves (2.5 g/kg BW).

Examination showed that the retentate was pathogen-free and of satisfactory bacteriological quality.

Group C was divided into three subgroups: 8 calves (subgroup C1) received pooled colostrum obtained at the first milking, according to group A conditions; 7 calves (subgroup C2) received an immunoglobulin solution (ISC, table 1) prepared as ISB; 8 calves (subgroup C3) were treated as subgroup C2, except that 160 g of milk powder were added per kg of immunoglobulin solution. This restored the dry matter percentage to nearly that of usual first milking colostrum, obtained by biochemical components enclosed in the colostrum, for most of them.

The alimentary levels assigned to these three subgroups were calculated so that the quantity of IgG ingested per kg of body weight was the same as that administered to groups A and B.

The four experimental meals were given to all the calves in a bucket fitted with a teat at the bottom. In case of mild inappetence, manual pressure was applied to the teat to facilitate ingestion. In case of acute inappetence, oesophageal intubation was practised.

Sampling and analysis. — Blood was taken by puncture of the jugular vein of every calf at the very moment of birth and at 4, 6 (7 h for group C), 10, 16, 22 and 28 h afterwards.

Similar sampling was carried out again 2, 3 and 4 days after birth and at the end of the first and second postnatal weeks. The level of IgG in blood plasma was determined by single radial immunodiffusion according to Mancini, Carbonata and Heremans (1965) and the level of alkaline phosphatase according to Bessey, Lowry and Brock (1946) using Boehringer-Mannheim (R) reagents. Cortisol was titrated in groups A and B according to Bosc and Fèvre. (1977) on blood samples collected at 0, 6 and 22 h and 3 days after birth.

As for the immunoglobulin solutions (ISB and ISC, table 1), IgG levels were also measured by the method cited above in all the colostra used. Antitryptic activity was measured according to Valdebouze et al. (1980) in the ISB and ISC solu-
tions and in two pooled colostra not related to this experiment but used as a good reference because of the large number of individual colostra mixed.

All the bull calves in group C were put into digestibility crates as soon as half an hour after birth, and their faeces were collected for 72 h. The faeces of all calves were mixed, lyophilized, and the amount of IgG was determined in order to calculate the apparent digestibility (AD) of these proteins. 72 h was considered to be long enough to permit complete transit of the last, immunoglobulin-rich meal.

Results.

Vitality, appetite and general health of the calves. — The vitality of all calves was good during the four hours before the first meal. The various treatments did not apparently impair it, except for a female calf (N° 209) belonging to group C2. This animal, exhibiting excellent vitality at birth, avidly ingested the first meal of immunoglobulin solution. She then refused to stand up, and died seventeen hours after birth. Postmortem examination revealed haemorrhagic enteritis of the jejunum.

A transient diarrhoea affected most of the animal of groups B and C2. The diarrhoea began at ten hours after birth and stopped approximately one day later. Its precocious and transient nature led us to exclude an infectious origin and, on the contrary, to suspect an alimentary one.

Colostrum meals were characterized by very few refusals (fig. 1), but this was not true of the immunoglobulin solutions. Although the first meal was ingested rather well (25 % refusals for group B, 0 % for group C2), refusal rate reached a maximum at the second meal (84 and 88 % for groups B and C2, respectively). The refusal rate pattern of group C3 was quite similar to that of group C2, although lower.

![Graph showing percentage of refusals of the first meals after birth in dairy calves fed colostrum or immunoglobulin solutions.](image)

FIG. 1. — Percentage of refusals of the first meals after birth in dairy calves fed colostrum or immunoglobulin solutions.

The satisfactory levels of ingestion recorded at the first meal of immunoglobulin solution seem to demonstrate that the appetency of the immunoglobulin solutions was not the basic cause of the numerous refusals noted later. On the
contrary, the first meal might have provoked digestive troubles, resulting in a general loss of appetite. On the other hand, this loss was corrected rather quickly since the refusal rate returned to a more reasonable level at the third and fourth meals.

The death of calf 209 and the excessively high refusal rates strengthen the notion that the immunoglobulin solutions were harmful to the digestive tract of the calves. The diarrhoeic episode reinforces this presumption, even if Selman, McEwan and Fisher (1970) noted a similar phenomenon in calves well fed with colostrum.

**Immunoglobulins, alkaline phosphatase and cortisol in blood plasma; antitrypsic activity in colostrum and immunoglobulin solutions.** — In all the groups, maximal plasma levels of immunoglobulins were found at 28 hours after birth (fig. 2), but the levels reached were very different: 32.8 ± 1.9 g/l in group A and 25.4 ± 3.5 in group C1. Groups which received the immunoglobulin solutions showed very low levels: 6.6 ± 0.5 in group B and 8.6 ± 1.3 in group C2. Group C3, which received an immunoglobulin solution supplemented with milk powder, exhibited intermediate levels which were significantly higher than those of groups B and C2 between 16 and 96 h after birth. However, these levels were very different from those of groups A and C1.

![Graph showing changes in plasma IgG levels](image)

**FIG. 2.** — Changes in plasma IgG levels in neonatal dairy calves fed colostrum or immunoglobulin solutions.

Immunoglobulin apparent digestibility was 0.93 ± 0.03 for the colostrum diet, 0.91 ± 0.02 for the immunoglobulin solution alone and 0.85 ± 0.06 for the immunoglobulin solution mixed with milk powder.

Antitrypsic activity is shown in table 1. The levels of this activity in relation to the immunoglobulin solutions were close to the values obtained with the two colostrum pools used as a reference.
The first drink of colostrum started an abrupt rise in plasma alkaline phosphatase levels, followed by a decline some hours after (fig. 3). The patterns of the curves seemed to be specific to the diet: the rise was rapid and high in calves fed the immunoglobulin solutions, and the drop sudden and low. In calves fed colostrum, the curve was less sharp, resulting in levels significantly lower than those of groups B and C₂ + C₃ at the onset of the experiment (7 and 10 h), and exceeding them afterwards (16, 22 and 28 h).

In groups A and B, cortisolemia was very high at birth (fig. 4) and decreased throughout the experiment. The group B levels, slightly more elevated at birth, became significantly higher than those of group A later.
Discussion.

The groups fed colostrum exhibited a maximal plasma IgG level near to the highest values in the literature (Gay, McGuire and Parish, 1983; Stott et al., 1979). This was in agreement with the good conditions in which the meals were delivered and the ability of the calves to absorb immunoglobulins. Consequently, the very low levels recorded in the calves fed the immunoglobulin solutions must be regarded as resulting from the composition of this solution. This ineffectiveness is probably the outcome of extracting the low molecular-weight protein liable to intestinal immunoglobulin absorption or of removing the precursor of this molecule (Balfour and Comline, 1962). In our experiment, the improvement seen when milk powder was added to the solution would suggest that these products were also present in the milk but in lower quantities.

The immunoglobulin solutions used here also had a low osmotic pressure owing to their equally low concentrations of lactose and minerals (table 1). However, this feature must not be considered as the prime cause of malabsorption since Balfour and Comline (1962) observed the same phenomenon with a well-equilibrated solution. However, the fact that the osmotic pressure was too low would be at the origin of the harmful action of the immunoglobulin solutions on the calf digestive tract.

This unfortunate property seems to be established by the clinical observations cited above and confirmed by the changes in cortisolemia which increased rapidly in the animals fed the immunoglobulin solutions. Considering this last result some caution must be taken, since feeding such solutions, very poor of energy level could be considered as underfeeding. It has been noted in humans that fasting increases cortisolemia (Palmblad et al., 1977) and in the newborn calf itself that a delay in feeding colostrum postpones the decrease in cortisolemia, which is very high at birth (Lamotte and Eberhart, 1976; Nightengale and Stott, 1981).

The fate of the immunoglobulins which did not appear in the blood circulation of the calves fed the immunoglobulin solutions is unknown. They were not found in the faeces in a complete state because their apparent digestibility was high and did not differ from one diet to another. It must be remembered that the immunoglobulin levels were determined by an immunological method which would be ineffective if the molecule was fractionated.

Some proteolysis could have occurred (Kumano et al., 1976), but to a similar extent in the diets since the antitrypsic activity of the immunoglobulin solutions and the reference colostra were the same. Thus, the main reason for the wide differences between the plasma immunoglobulin levels could not be determined.

The rapid disappearance of immunoglobulins from the blood compartment due to a severe proteinuria was also unexpected. Such a phenomenon was already observed in the newborn calf by McDougall (1965), but it is unlikely that its intensity would depend so much on the type of diet.

There is no alternative but to admit with Matte et al. (1982) that a great part of the immunoglobulins seemed to be lost or, at least, was hardly perceptible by the means used here. However, after careful examination of the curves, a hypo-
thesis can be proposed in relation to alkaline phosphatases (fig. 3). Healy (1975), in association later with Dinsdale (Healy and Dinsdale, 1979), was the first to report that a rise in plasma alkaline phosphatase levels always occurred at the time the immunoglobulins appeared in the blood circulation. This enzyme would be a part of the wall of subcellular vesicles containing immunoglobulins and released into the lymph circulation by enterocytes (Dinsdale and Healy, 1982). The enzyme is also present in colostrum (Linden and Maraval, 1979), but at a rate quite unable to account for the very high plasma concentrations after colostrum ingestion. This is contrary to gamma-glutamyl transferase, an enzyme appearing at the same moment and whose colostral origin has been determined by Braun et al. (1982).

Here, the origin of alkaline phosphatase is clearly endogenous and its abundance in plasma evidences a release of vesicles containing immunoglobulins. Therefore in groups B and C2 + C3, after an extremely sharp rise, the levels suddenly dropped to significantly lower levels than those of the groups fed colostrum. This leads us to postulate a premature termination of transport, resulting in very low levels of plasma immunoglobulin. It should be noted, however, that cessation of transport does not necessarily mean cessation of absorption since, in the piglet, Baintner (1973a, b) measured the time-lapse between these two phenomena, an interval associated with the sequestration of immunoglobulins in the intestinal wall.

Conclusions. Colostrum is a very complex biochemical medium. It contains immunoglobulins required for the sanitary protection of the newborn animals. Besides these, colostrum also contains other factors, as yet poorly defined, which allow the immunoglobulins to be absorbed and transported. It is of interest to identify these factors because, if not, it will be impossible to carry out the project of feeding newborn ruminants with immunoglobulins industrially extracted from colostrum, blood serum and especially from lactoserum when maternal colostrum is lacking. It is clear that without these unknown factors, immunoglobulins are unable to pass into the plasma of newborn animals.

Résumé. Absorption des immunoglobulines par le veau nouveau-né.

L'exploitation industrielle des surplus de colostrum bovin étant envisageable à court terme, on a étudié, chez le veau nouveau-né, l'absorption intestinale des immunoglobulines extraites du colostrum par ultrafiltration. Quatre-vingt quinze animaux issus du même troupeau laitier lors de trois campagnes de vêlage successives ont été répartis en lots suivant les règles expérimentales usuelles. Trois régimes ont été comparés : colostrum de première traite, solution saline d'immunoglobulines extraites du colostrum par ultrafiltration et solution saline d'immunoglobulines extraites du colostrum par ultrafiltration additionnée de poudre de lait entier. Les repas ont été administrés très exactement quatre, dix, seize et vingt-deux heures après la naissance. Les quantités des différents régimes ont été ajustées de façon à ce que les quantités moyennes d'immunoglobulines reçues par les animaux fussent identiques. Des prélèvements sanguins réalisés suivant une cinétique précise, s'étendant de la naissance à vingt-huit heures de vie aérienne ont montré que les immunoglobulines con-
tenues dans la solution saline étaient absorbées dans une très faible mesure. On a pu augmenter significativement cette absorption par l’adjonction de poudre de lait entier à la solution. Le dosage dans le plasma des phosphatases alcalines, issues pour la plupart de l’intestin, à ce stade de la vie, a montré que l’évolution de leur teneur était très dépendante du régime. Ceci pourrait attester de différences dans l’évolution du fonctionnement de l’intestin qui seraient provoquées par le régime mais pour lesquelles les auteurs ne sont pas en mesure, actuellement, de proposer une explication.

References


